A TREATISE ON SYSTEMS
VOLUME I

ANALYTICAL DESCRIPTION OF
HUMAN-INFORMATION NETWORKS

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This Treatise on Systems is both a work for studying and for reference. It is composed of two volumes: the first, a restoration of an earlier book written in 2002, and the second a continuation consisting of new perspectives. My focus for almost 25 years has been the question: how can we ask the right questions about systems, without having to learn every lesson the hard way?

I wrote this first volume hurriedly in the university summer vacation of 2002, as a text for a new course in analytical thinking, as part of an experimental Oslo University College Master of Science programme in Network and System Administration. It bore the title Analytical Network and System Administration: Managing Human-Computer Systems, and was intended as a synthesis of all the methods one could use to analyze systems of humans interacting with computers from a mathematical logical perspective, drawn from any and every discipline. My aim was to present systems from the perspective of analysis of resources, scaling, and maintenance, for a future of ‘pervasive computing’, in which computers would be literally everywhere\(^1\). This was the time of the Xerox PARC vision, which would later be rebranded as ‘The Internet of Things’. I was searching for ways to describe pragmatic systems using mathematical methods, feeling a deep dissatisfaction with the recipe approaches of OSI, ITIL, and the academic literature that made no attempt to address any issues that they had not already experienced. Although great engineers can come up with analytical approaches on their own, given plenty of time and experience, I believed that good engineering should not be the preserve to brilliant individuals. With a little training in how to think about problems, analysis is available to everyone.

Many of the methods I picked could be drawn from my own research into systems and their scaling behaviours. In spite of a valiant and well-meaning effort, looking back, some of the approaches (e.g. game theory) seem as naive and idealistic as the corresponding ideas did in their native fields of economics and biology. I was clutching at straws, asking what could be done to go beyond the ad hoc methods of the day? Others have implicitly entered modern datacentre design (queueing theory) without any help from me. Unfortunately, for many years datacentre engineering disappeared from public view into the private sector: Amazon, Google, and later others sucked up

\(^{1}\)See the story of this in my book In Search of Certainty [Bur13a]
talented engineers and created a divide between those pushing the envelope, and those left behind.

So why revive this now? Isn’t it too late? I don’t think so. Today, we have passed the age of the treatise, yet (call me old fashioned) I still believe in the virtue of coherent thought, and most engineers have still not made the transition to treating system engineering as a science. Recently, I was honoured to write the introduction to Google’s Site Reliability Engineering handbook[BJPM16], which makes a perfect companion volume to this theoretical compendium. Most engineers turn their noses up at theory, but Google’s work show how that is mistaken. Most will read the non-technical account of Site Reliability Engineering, in the modern way (as a branded concept), and find it better than a book on theoretical thinking. But, I believe that serious individuals will still be grateful for a cultural context for these ideas. If this treatise helps only one other person than me, it will have been worth it. And the wonders of modern print on demand make its availability viable.

The university degree, which motivated this older volume, was not to be a course in computer science and distributed algorithms (which would not have been comprehensible to students who applied for engineering, and would have given little insight into how actual systems work in practice). Nor could I claim that anyone was actually using the approaches in the world of industry (except, of course, myself out of sheer dogged insistence that science is important). Ultimately, the attempt was ridiculed by the self-appointed guardians of system administration, with the exception of a few maverick enthusiasts2.

The single semester course, for me, was a great success. It accomplished what I set out to do beyond my wildest expectations. Students, who professed to never having learned or been interested in mathematics, became some of the highest achievers, reaching more than an ability to solve equations by rote: they came to understand what the equations meant, how they were approximations to reality, and what the possible answers meant. Later, when I left the university, the course was dropped in favour of some standard mathematical methods, without specific application to systems, and the book became priced beyond the reach of sub-financiers by the publisher.

More interestingly, this was exactly the time when the industry was undergoing a revolution thanks to the explosion of web services. In the their secretive way Google was building some of the largest systems on the planet. They were later followed by Facebook and by others. Many of these companies used CFEngine and its principles of scaling to build their early datacentres. Typical computer systems grew from being of the order of 100 computers to 10,000 and then 100,000, and beyond. As the years went by, some of the engineers involved told me that they secretly admired the original book, and the work it was based on. Having tried more traditional programming, many recreated CFEngine-like technologies and architectures due to scaling issues. Moreover, many authors began to write that the knowledge they applied to understand the system was more like

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2 This pattern seems to follow me constantly, but over time the works have been rediscovered and appreciated at least in part.
physics than computer science. This began with the server administrators, database designers, then the network administrators. The acknowledgement of physics as the science of scale was in line with my expectations. Most computer scientists never confront the ideas of scale or dynamical stability, because they are focused on microscopic mechanisms of intent and possibility. I later wrote a popular science account of the work *In Search of Certainty* to try to reach a wider audience without use of mathematics.

One outcome of this hurriedly prepared book was that it focused my attention on a huge hole in theory, concerning the role of policy: i.e. the choices and decisions that are not based on rational methods, but which are simply chosen because they are what people intend to do. Systems are, after all, designed with a purpose in mind. Physics is designed to not acknowledge such subjectivities. This was an important link between the intentional design of computer science and the unintended emergent outcomes of actual systems. Following up on this point led directly to Promise Theory[BB14].

In this second edition (in addition to bringing the book back from the edge of price reality to where readers can actually afford it), I have chosen not to alter the book significantly. However, I have attempted to update it somewhat, peppering the text with hints about where the theory led, as well as examples from the present, without trying to alter its historical naivety, poised as it was on the brink of modern datacentre design.

Some fifteen years later, we have learned much in practice, often in ways that could have been shortened by a little theoretical know-how, while the ideas presented in this first volume are still only just entering mainstream, second-hand through technologies of those few individuals who applied them, with or without my help. I register a growing interest in science amongst IT operations engineers, nonetheless, and a curious interest in Promise Theory for its attempt to bridge what seem like disparate and contradictory worlds of humanity and technology. There is no other analytical language to describe the phenomena they confront in the daily lives. Most authors still rely on rhetoric and explicit demonstration to argue their cases. I am humbly pleased that these early writings seem to have played some tiny role in bringing science to bear through a minority of willing minds.

Today, the potential for theoretical modelling is even more exciting. Human-computer systems have spread beyond the datacentre into our daily lives. The idea of ‘semantic spacetime’[Bur14, Bur15, Bur16b] takes Promise Theory to new levels of scale, to view deeply embedded functional systems at high density for the coming age of pervasive embedded computing. Today, we also have actual data about scaling to compare to theory, as well as the qualitative experiences described in Google’s own story of system design[BJPM16], and these accounts vindicate the use of analysis quite well. I hope the restoration of this introductory volume along with the second volume, which looks forward, may inspire a few individuals to continue to delve more into analytical methods, to better understand the human-information systems that we are creating move the planet forward.
FOREWORD TO FIRST EDITION

It is my great honor to introduce a landmark book in the field of network and system administration. For the first time, in one place, one can study the components of network and system administration as an evolving and emerging discipline and science, rather than as a set of recipes, practices or principles. This book represents the step from ‘mastery of the practice’ and ‘scientific understanding’, a step very similar to that between historical alchemy and chemistry.

As recently as ten years ago, many people considered ‘network and system administration’ to comprise remembering and following complex recipes for building and maintaining systems and networks. The complexity of many of these recipes and the difficulty of explaining them to nonpractitioners in simple and understandable terms—encouraged practitioners to treat system administration as an ‘art’ or ‘guild craft’ into which practitioners are initiated through apprenticeship.

Current master practitioners of network and system administration are perhaps best compared with historical master alchemists at the dawn of chemistry as a science. In contrast to the distorted popular image of alchemy as seeking riches through transmutation of base metals, historical research portrays alchemists as master practitioners of the subtle art of combining chemicals towards particular results or ends. Practitioners of alchemy often possessed both precise technique and highly developed observational skills. Likewise, current master practitioners of network and system administration craft highly reliable networks from a mix of precise practice, observational skills and the intuition that comes from careful observation of network behaviour over long time periods. But both alchemists and master practitioners lack the common language that makes it easy to exchange valuable information with others: the language of science.

Alas, the alchemy by which we have so far managed our networks is no longer sufficient. When networks were simple in structure, it was possible to maintain them through the use of relatively straightforward recipes, procedures and practices. In the post-Internet World, the administrator is now faced with managing and controlling networks that can dynamically adapt to changing conditions and requirements quickly and, perhaps, even unpredictably. These adaptive networks can exhibit ‘emergent properties’ that are not predictable in advance. In concert with adapting networks to serve human needs, future administrators must adapt themselves to the task of
management by developing an ongoing, perpetually evolving, and shared understanding.

In the past, it was reasonable to consider a computer network as a collection of cooperating machines functioning in isolation. Adaptive networks cannot be analysed in this fashion; their human components must also be considered. Modern networks are not communities of machines, but rather communities of humans inextricably linked by machines: what the author calls ‘Cooperating ecologies’ of users and machines. The behaviour of humans must be considered along with the behaviour of the network for making conclusions about network performance and suitability.

These pressures force me to an inescapable conclusion. System administrators cannot continue to be alchemist-practitioners. They must instead develop the language of science and evolve from members of a profession to researchers within a shared scientific discipline. This book shows the way.

Though we live thousands of miles apart, the author and I are ‘kindred spirits’—forged by many of the same experiences, challenges and insights. In the late 1980s and early 1990s, both of us were faculty, managing our own computer networks for teaching and research. Neither of us had access to the contemporary guilds of system administration (or each other), and had to learn how to administer networks the hard way-by reading the documentation and creating our own recipes for success. Both of us realized (completely independently) that there were simple concepts behind the recipes that, once discovered, make the recipes easy to remember, reconstruct and understand. Concurrently and independently, both of us set out to create software tools that would avoid repeated manual configuration.

Although we were trained in radically differing academic traditions (the author from physics and myself from mathematics and computer science), our administrative tools, developed completely in isolation from one another, had very similar capabilities and even accomplished tasks using the same methods. The most striking similarity was that both tools were based upon the same ‘principles’. For the first time, it very much looked like we had found an invariant principle in the art of system and network administration: the ‘principle of convergence’. As people would say in the North Carolina backwoods near where I grew up, ‘if it ain’t broke, don’t fix it’.

The road from alchemy to discipline has many steps. In the author’s previous book, Principles of Network and System Administration, he takes the first step from practice (‘what to do’) to principles (‘why to do it’). Recipes are not created equal; some are better than others. Many times the difference between good and poor recipes can be expressed in terms of easily understood principles. Good recipes can then be constructed top-down, starting at the principles. Practitioners have approached the same problem bottom-up, working to turn their tested and proven recipes into sets of ‘best practices’ that are guaranteed to work well for a particular site or application. Recently, many practitioners have begun to outline the ‘principles’ underlying their practices. There is remarkable similarity between the results of these two seemingly opposing processes, and the author’s ‘principles’, and the practitioners’ ‘best practices’ are now quickly meeting on a common
middle ground of principles.

In this book, for the first time, the author identifies principles of scientific practice and observation that anyone can use to become proficient ‘analysts’ of network and system administration practices. This will not make one a better practitioner, but rather will allow one to discuss and evaluate the practice with others in a clear and concise manner. The reader will not find any recipes in this book. The reader will not find principles of practice. Rather, the book explains the principles behind the science and chemistry of cooking, so that one can efficiently derive one’s own efficient and effective recipes for future networks.

Proficient system administrators have always been capable of this kind of alchemy, but have found it challenging to teach the skill to others. This book unlocks the full power of the scientific method to allow sharing of analyses, so that future administrators can look beyond recipe, to shared understanding and discipline. In this way, now-isolated practitioners can form a shared scientific community and discipline whose knowledge is greater than the sum of its parts.

Looking at the table of contents, one will be very surprised to note that the traditional disciplines of ‘computer science’ and ‘computer engineering’-long considered the inseparable partners of system administration-are not the basis of the new science. Rather, experimental physics has proven to be the Rosetta Stone that unlocks the mysteries of complex systems. To understand why, we must examine the fundamental differences in economics between the disciplines of computer science and engineering and the disciplines of network and system administration.

Traditional computer science and engineering (and, particularly, the sciences involved in building the systems that system administrators manage) are based upon either an operational or axiomatic semantic model of computing. Both models express ‘what a program does’ in an ideal computing environment. Software developers build complex systems in layers, where each subsequent layer presumes the correct function of layers upon which it is built. Program correctness at a given layer is a mathematical property based upon axioms that describe the behaviour of underlying layers. Fully understanding a very complex system requires understanding of each layer and its interdependencies and assumptions in dealing with other layers.

System administrators have a differing view of the systems they manage compared to that of the developers who designed the systems. It is not economically feasible to teach the deep knowledge and mathematical understanding necessary to craft and debug software and systems to large populations of human system administrators. System administrators must instead base their actions upon a high-level set of initial experimental hypotheses called the ‘system documentation’. The documentation consists of hypotheses to be tested, not axioms to be trusted. As administrators learn how to manage a system, they refine their understanding top-down, by direct observation and ongoing evaluation of hypotheses.

Turning system and network administration into a discipline requires one to learn some skills, previously considered far removed from the practice. Evaluating hypotheses requires a
rudimentary knowledge of statistics and the experimental method. These hypotheses are built not upon operational or axiomatic semantic models of computing, but upon specialized high-level mathematical models that describe behaviour of a complex system. With this machinery in hand, several advanced methods of analysis-prevalent in experimental physics and other scientific disciplines—are applied to the problem of understanding management of complex systems.

Proficient system administrators are already skilled experimental scientists; they just do not acknowledge this fact and cannot effectively communicate their findings to others. This book takes a major step towards understanding the profession of system and network administration as a science rather than as an art. While this step is difficult to take, it is both rewarding and necessary for those pioneers who will manage the next generation of networks and services. Please read on, and seek to understand the true nature of networking—as a process that involves connecting humans, not just computers.

Alva Couch Tufts University, USA, 2003
This is a research document and a textbook for graduate students and researchers in the field of networking and system administration. It offers a theoretical perspective on human-computer systems and their administration. The book assumes a basic competence in mathematical methods, common to undergraduates courses. Readers looking for a less theoretical introduction to the subject may wish to consult [Bur00b].

I have striven to write a short book, treating topics briefly rather than succumbing to the temptation to write an encyclopædia that few will read or be able to lift. I have not attempted to survey the literature or provide any historical context to the development of these ideas (see [ABC01]). I hope this makes the book accessible to the intelligent lay reader who does not possess an extensive literacy in the field, and would be confused by such distractions. The more advanced reader should find sufficient threads to follow to add depth to the material. In my experience, too much attention to detail merely results in forgetting why one is studying something at all. In this case, we are trying to formulate a descriptive language for systems.

A theoretical synthesis of system administration plays two roles: it provides a descriptive framework for systems that should be available to other areas of computer science, and it proffers an analytical framework for dealing with the complexities of interacting components. The field of system administration meets an unusual challenge in computer science: that of approximation. Modern computing systems are too complicated to be understood in exact terms.

In the flagship theory of physics, quantum electrodynamics, one builds everything out of two simple principles:

1. Different things can exist at different places and times.
2. For every effect, there must be a cause.

The beauty of this construction is its lack of assumptions and the richness of the results. In this text, I have tried to synthesize something like this for human-computer systems. In order to finish the book, and keep it short and readable I have had to compromise on many things. I hope that the
result nevertheless contributes in some way to a broader scientific understanding of the field and will inspire students to further serious study of this important subject.

Some of this work is based on research performed with my collaborators Geoff Canright, Frode Sandnes and Trond Reitan. I have benefitted greatly from discussions with them and others. I am especially grateful for the interest and support of other researchers, most notably Alva Couch for understanding my own contributions when no one else did. Finally I would like to thank several for reading draft versions of the manuscript and commenting: Paul Anderson, Lars Kristiansen, Tore Jonassen, Anil Somayaji, Jan Bergstra.

MB June 2002
CHAPTER 1

INTRODUCTION

Technology: the science of the mechanical and industrial arts.
[Gk. tekhne art and logos speech].

– Odhams dictionary of the English language

1.1 SYSTEMS AND THEIR MANAGEMENT?

The management of systems, called variously management theory or system administration is about the design, running and maintenance of human-information systems. Human-information systems are ‘communities’ of people and machines that collaborate actively to execute a common task. Examples of human-information systems include business enterprises, service institutions and any extensive machinery that is operated by, or interacts with human beings. The human players in a human-information system are often called the users and the machines are referred to as hosts, but this suggests an asymmetry of roles which is not always the case.

System administration is primarily about the technological side of a system: the architecture, construction and optimization of the collaborating parts, but it also occasionally touches on softer factors such as user assistance (help desks), ethical considerations in deploying a system, and the larger implications of its design for others who come into contact with it. System administration deals first and foremost with the system as a whole, treating the individual components as black boxes, to be opened only when it is possible or practical to do so. It does not conventionally consider the design of user-tools such as third-party computer programs, nor does it attempt to design enhancements to the available software, though it does often discuss meta-tools and improvised software systems which can be used to monitor, adjust or even govern the system. This
omission is mainly because user-software is acquired beyond the control of a system administrator; it is written by third parties, and is not open to local modification. Thus users’ tools and software are treated as ‘given quantities’ or ‘boundary conditions’.

For historical reasons, the study of system administration has fallen into two camps: those who speak of network management and discuss its problems in terms of software design for the management of black box devices by humans (e.g. using SNMP), and those who speak of system administration and concern themselves with practical strategies of machine and software configuration at all levels, including automation, human-information issues and ethical considerations. These two viewpoints are complementary, but too often ignore one another. This book considers human-information systems in general, and refers to specific technologies only by example. It is therefore as much about purely human administrative systems as it is about computers.

1.2 What is a System?

A system is most often an organized effort to fulfill a goal, or at least carry out some predictable behaviour. The concept is of the broadest possible generality. A system could be a mechanical device, a computer, an office of workers, a network of humans and machines, a series of forms and procedures (a bureaucracy) etc. Systems involve themes, such as collaboration and communication between different actors, the use of structure to represent information or to promote efficiency, and the laws of cause and effect. Within any mechanism, specialization of the parts is required to build significant innovation; it is only through strategy of divide and conquer that significant problems can be solved. This implies that each division requires a special solution.

A computer system is usually understood to mean a system composed primarily of computers, using computers or supporting computers. A human-information system includes the role of humans, such as in a business enterprise where computers are widely used. The principles and theories concerning systems come from a wide range of fields of study. They are synthesized here in a form and language which is suitable for scholars of science and engineering.

1.3 What is Administration?

The word administration covers a variety of meanings in common parlance. The American Administration is the government of the United States, i.e. a political leadership. A university administration is a bureaucracy and economic resource department, which works on behalf of a board of governors to implement university policy and to manage its resources. The administrative department of a company is generally the part which handles economic procedures and payment transactions. In human-information system administration the definition is broadened to include all of the organizational aspects and also engineering issues, such as system fault diagnosis. In
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this regard, it is like the medical profession, which combines checking, management and repair of bodily functions. The main issues are:

- System design and rationalization
- Resource management
- Fault finding

In order to achieve these goals, it requires

- Procedure
- Team work
- Ethical practices
- Appreciation of security

Administration comprises two aspects: technical solutions and arbitrary policies. A technical solution is required to achieve goals and sub-goals, so that a problem can be broken down into manageable pieces. Policy is required to make the system, as far as possible, predictable: it pre-decides the answers to questions on issues which cannot be derived from within the system itself. Policy is therefore an arbitrary choice, perhaps guided by a goal or principle.

The arbitrary aspect of policy cannot be disregarded from the administration of a system, since it sets the boundary conditions under which the system will operate, and supplies answers to questions which cannot be determined purely on the grounds of efficiency. This is especially important where humans are involved: human welfare, permissions, responsibilities and ethical issues are all parts of policy. Modelling these intangible qualities, formally, presents some challenges and requires the creative use of abstraction.

The administration of a system is an administration of temporal and resource development. The administration of a network of localized systems (a so-called distributed system) contains all of the above, and additionally the administration of the location of and communication between the system’s parts. Administration is thus a flow of activity, information about resources, policy making, record keeping, diagnosis, and repair.

1.4 STUDYING SYSTEMS

There are many issues to be studied in system administration. Some issues are of a technical nature, while others are of a human nature. System administration confronts the human-machine interaction as few other branches of computer science do. Here are some examples:
• **System design** (e.g. how to get humans and machines to do a particular job as efficiently as possible. What works? What does not work? How does one know?)

• **Reliability studies** (e.g. failure rate of hardware/software, evaluation of policies and strategies)

• **Determining and evaluating methods for ensuring system integrity** (e.g. automation, cooperation between humans, formalization of policy, contingency planning etc.)

• **Observations which reveal aspects of system behaviour which are difficult to predict** (e.g. strange phenomena, periodic cycles).

• **Issues of strategy and planning.**

Usually system administrators do not decide the purpose of a system, they are regarded as supporting personnel. As we shall see, this view is somewhat flawed from the viewpoint of system design, however. It does not always make sense to separate the human and computer components in a system; as we move farther into the information age, the fates of both become more deeply intertwined.

To date, little theory has been applied to the problems of system administration. In a subject which is complex, like system administration, it is easy to fall back on qualitative claims. This is dangerous, however, since one is easily fooled by qualitative descriptions. Analysis proceeds as a dialogue between theory and by experiment. We need theory to interpret results of observations and we need observations to back up theory. Any conclusions must be a consistent mixture of the two. At the same time, one must not believe that it is sensible to demand hard-nosed Popper-like falsification of claims in such a complex environment. Any numbers which we can measure, and any models we can make must be considered valuable, provided they actually have a sensible interpretation.

**Human-computer interaction**

The established field of human-computer interaction (HCI) has grown up, in computer science, around the need for reliable interfaces in critical software scenarios (see for instance [She96, Zad73]). For example, in the military, real danger could come of an ill-designed user interface on a nuclear submarine; or in a power plant, a poorly designed system could set off an explosion or result in blackouts.

One can extend the notion of the human-computer interaction to think less as a programmer and more as a physicist. The task of physics is to understand and describe what happens when different parts of nature interact. The interaction between fickle humans and rigid machinery leads to many unexpected phenomena, some of which might be predicted by a more detailed functional
understanding of this interaction. This does not merely involve human attitudes and habits; it is a problem of systemic complexity — something that physics has its own methods to describe. Many of the problems surrounding computer security enter into the equation through the human-computer interaction. Of all the parts of a system, humans bend most easily: they are often both the weakest link and the most adaptable tools in a solution, but there is more to the human-computer interaction than psychology and button pushing. The issue reaches out to the very principles of science: what are relevant time scales for the interactions and for the effects to manifest? What are the sources of predictability and unpredictability? Where is the system immune to this interaction, and where is the interaction very strong? These are not questions that a computer science analysis alone can answer; there are physics questions behind these issues. Thus, in reading this book, you should not be misled into thinking that physics is merely about electrons, heat and motion: it is broad methodology for ‘understanding phenomena’, no matter where they occur, or how they are described. What computer science lacks from its attachment to technology, it must regain by appealing to the physics of systems.

POLICY

The idea policy plays a central role in the administration of systems, whether they are dominated by human or technological concerns.

**Definition 1** (Policy - heuristic). A policy is a description of what is intended and desirable about a system. It includes a set of ad hoc choices, goals, compromises, schedules, definitions and limitations about the system. Where humans are involved, compromises often include psychological considerations, and welfare issues.

A policy provides a frame of reference in which a system is understood to operate. It injects a relativistic aspect into the science of systems: we cannot expect to find absolute answers, when different systems play by different rules and have different expectations. A theory of systems must therefore take into account policy as a basic axiom. Much effort is expended in the chapters that follow to find a tenable definition of policy.

STABILITY AND INSTABILITY

It is in the nature of almost all systems to change with time. The human and machine parts of a system change, both in response to one another, and in response to a larger environment. The system is usually a predictable, known quantity; the environment is, by definition, an unknown quantity. Such changes tend to move the system in one or two directions: either the system falls into disarray or it stagnates. The meaning of these provocative terms is different for the human and the machine parts:
• Systems will fall into a stable repetition of behaviour (a limit cycle) or reach some equilibrium at which point further change cannot occur without external intervention.

• Systems will eventually invalidate their assumptions and fail to fulfil their purpose.

Ideally a machine will perform, repetitively, the same job over and over again, because that is the function of mechanisms: stagnation is good for machines. For humans, on the other hand, this is usually regarded as a bad thing, since humans are valued for their creativity and adaptability. For a system mechanism to fall into disarray is a bad thing.

The relationship between a system and its environment is often crucial in determining which of the above is the case. The inclusion of human behaviour in systems must be modelled carefully, since humans are not deterministic in the same way that machines (automata) can be. Humans must therefore be considered as being part system and part environment. Finally, policy itself must be our guide as to what is desirable change.

SECURITY

Security is a property of systems, that has come to the forefront of our attention in recent times. How shall we include it in a theory of system administration?

Definition 2 (Security). Security concerns the possible ways in which a system’s integrity might be compromised, causing it to fail in its intended purpose. In other words, a breach of security is a failure of a system to meet its specifications.

Security refers to ‘intended purpose’, so it is immediately clear that it relates directly to policy and that it is a property of the entire system in general. Note also that, while we associate security with ‘attacks’ or ‘criminal activity’, natural disasters or other occurrences could be equally to blame for the external perturbations that break systems.

A loss of integrity can come from a variety of sources, e.g. an internal fault, an accident or a malicious attack on the system. Security is a property that requires the analysis of assumptions that underpin the system, since it is these areas which one tends to disregard and which can be exploited by attackers, or fail for diverse reasons. The system depends on its components in order to function. Security is thus about an analysis of dependencies. We can sum this up in a second definition:

Definition 3 (Secure system). A secure system is one in which every possible threat has been analyzed and where all the risks have been assessed and accepted as a matter of policy.
1.5 WHAT’S IN A THEORY?

When it was first published, I wrote that this book was not a finished theory, like the theory of relativity, or the theory of genetic replication. It was not to be understood the end of a story, but a beginning. In particular, system administration as applied to computers, was at the start of its scientific journey, not at its end. In the intervening years, there have been many changes in the information technology industry, and developments like the explosion of social media and cloud computing, implicitly using the ideas presented in the book. In most cases, the technological accomplishments were built without any credit due to the former edition. In other cases, companies like Facebook, LinkedIn, as well as Wall Street used the ideas through the proxy of the CFEngine software, which embodied the learning documented therein.

DRAMATIS PERSONAE

The players in systems and their administration are:

- Computers.
- Networks.
- System users or participants.
- Policy.
- Administrators.

The goal of this volume is to apply standard methods of rational scientific methodology to the description of systems, and to provide an overview of those methods in a way that cannot be found elsewhere. Thus, its aim is to form a bridge between mathematics and system engineering. We seek a clear and flexible language (rooted in mathematics) in which to write their script. It will deal with basic themes of:

- Time (when events occur or should occur).
- Location (where resources should be located).
- Value (how much the parts of a system contribute or are worth).
- Randomness and predictability (our ability to control or specify).

It must answer questions that are of interest to the management of systems. We can use two strategies:
• Type I (pure science) models that describe the behaviour of a system without attempting to interpret its value or usefulness. These are ‘vignettes’ that describe what we can observe and explain it in impartial terms. They provide a basic understanding of phenomena that leads to expertise about the system.

• Type II (applied science) models add interpretations of value and correctness (policy) to the description. They help up to make decisions by impressing a rational framework on the subjectivities of policy.

A SNAPSHOT OF REALITY

The system administrator rises and heads for the computer. Grabs coffee or cola and proceeds to catch up on E-mail. There are questions, bug-reports, automatic replies from scripted programs, spam and lengthy discussions from mailing lists.

The day proceeds to planning, fault finding, installing software, modifying system parameters to implement (often ad hoc) policy that enables the system to solve a problem for a user, or which makes the running smoother (more predictable) — see fig. 1.1. On top of all of this, the administrator must be thinking about what users are doing. After all, they are the ones who need the system and the ones who most often break it. How does ‘the system’ cope with them and their activities as they feed off it and feed back on it. They are, in every sense, a part of the system. How can their habits and skills be changed to make it all work more smoothly? This will require an appreciation of the social interactions of the system and how they, in turn, affect the structures of the logical networks and demands placed on the machines.

There are decisions to be made, but many of them seem too uncertain to be able to make a reliable judgement on the available evidence. Experimentation is required, and searching for advice from others. Unfortunately, you never know how reliable others’ opinions and assertions will be. It would be cool if there were a method for turning the creative energy into the optimal answer. There is ample opportunity and a wealth of tools to collect information, but how should that information be organized and interpreted? What is lacking is not software, but theoretical tools.

What view or philosophy could unify the different facets of system administration: design, economics, efficiency, verification, fault-finding, maintenance, security, and so on? Each of these issues is based on something more primitive or fundamental. Our task is therefore to use the power of abstraction to break down the familiar problems into simpler units that we can master and then reassemble into an approximation of reality. There is no unique point of view here (see next chapter).

Theory might lead to better tools but also to better procedures. If it is to be of any use, it must have predictive power as well as descriptive power. We have to end up with formulae and procedures that make criticism and re-evaluation easier and more effective. We must be able to
summarize simple ‘laws’ about system management (thumb-rules) that are not based only on vague experience, but have a theoretical explanation based on reasonable cause and effect.

How could such a thing be done? For instance: how might we measure how much work will be involved in a task?

- We would have to distinguish between the work we actually do and how much work is needed in principle (efficiency and optimization).

- We would look for a mathematical idea with the characteristics or properties of work. We find that we can map work into the idea of ‘information’ content in some cases (now we have something concrete to study).

- Information or work is a statistical concept: information that is transmitted often can be compressed on average — if we do something often, efficiencies can be improved through economies of scale.
By starting down the road of analysis, we gain many small insights that can be assembled into a deeper understanding. That is what this book attempts to do.

The system administrator wonders if he or she will ever become redundant, but there is no sign of that happening. The external conditions and requirements of users are changing too quickly for a system to adapt automatically, and policy has to be adjusted to new goals and crises. Humans are the only technology on the planet that can address that problem for the foreseeable future. Besides, the pursuit of pleasure is a human condition, and part of the enjoyment of the job is that creative and analytical pursuit.

The purpose of this book is to offer a framework in which to analyze and understand the phenomena of human-computer management. It is only with the help of theoretical models that we truly can obtain a deeper understanding of system behaviour.

STUDIES

The coming chapters describe a variety of languages for discussing systems, and present some methods and issues that are the basis of the author’s own work. Analysis is the scientific method in action, so this book is about analysis. It has many themes:

1. **Observe** — we must establish a factual basis for discussing systems.

2. **Deduce cause** — we establish probable causes of observed phenomena.

3. **Establish goals** — what do we want from this information?

4. **Diagnose ‘faults’** — what is a fault? It implies a value judgement, based on policy.

5. **Correct faults** — devise and apply strategies.

Again, these concepts are intimately connected with ‘policy’, i.e. a specification of right and wrong. In some sense, we need to know the ‘distance’ between what we would like to see and what we actually see.

This is all very abstract. In the day to day running of systems, few administrators think in such generalized, abstract terms — yet this is what this book asks you to do.

**Example 1** (A backup method). A basic duty of system administrators is to perform a backup of data and procedures: to ensure the integrity of the system under natural or unnatural threats. How shall we abstract this and turn it into a scientific enquiry?

We might begin by examining how data can be copied from one place to another. This adds a chain of questions: i) how can the copying be made efficient? ii) what does efficient mean? iii) how often do the data change, and in what way? What is the best strategy for making a copy: immediately after every change, once per day, once per hour? We can introduce a model for the
change, e.g. a mass of data that is more or less constant, with small random fluctuating changes to some files, driven by random user activity. This gives us something to test against reality. Now we need to know how users behave, and what they are likely to do. We then ask: what do these fluctuations look like over time? Can they be characterized, so that we can tune a copying algorithm to fit them? What is the best strategy for copying the files?

The chain of questions never stops: analysis is a process, not an answer.

Example 2 (Resource management). Planning a system’s resources, and deploying them so that the system functions optimally is another task for a system administrator. How can we measure, or even discuss the operation of a system to see how it is operating? Can important (centrally important) places be identified in the system, where extra resources are needed, or the system might be vulnerable to failure? How shall we model demand and load? Is the arrival of load (traffic) predictable or stochastic? How does this affect our ability to handle it? If one part of the system depends on another, what does this mean for the efficiency or reliability? How do we even start asking these questions analytically?

Example 3 (Pattern detection). Patterns of activity manifest themselves over time in systems. How do we measure the change, and what is the uncertainty in our measurement? What are their causes? How can they be described and modelled? If a system changes its pattern of behaviour, what does this mean? Is it a fault or a feature?

In computer security, intrusion detection systems often make use of this kind of idea, but how can the idea be described, quantified and generalized, hence evaluated?

Example 4 (Configuration management). The initial construction and implementation of a system, in terms of its basic building blocks is referred to as its configuration. It is a measure of the system’s state or condition. How should we measure this state? Is it a fixed pattern, or a statistical phenomenon? How quickly should it change? What might cause it to change unexpectedly? How big a change can occur before the system is damaged? Is it possible to guarantee that every configuration will be stable, perform its intended function, and be implementable according to the constraints of a policy?

In each of the examples above, an apparently straightforward issue generates a stream of questions that we would like to answer. Asking these questions is what science is about: answering them involves the language of mathematics and logic in concert with a scientific inquiry: science is about extracting the essential features from complex observable phenomena and modelling them in order to make predictions. It is based on observation and approximate verification. There is no “exact science” as we sometimes hear about in connection with physics or chemistry; it is always about suitably idealized approximations to the truth, or “uncertainty management”. Mathematics,
on the other hand, is not to be confused with science — it is about rewriting assumptions in
different ways; i.e. if one begins with a statement that is assumed true (an axiom) and manipulates
it according to the rules of mathematics, the resulting statement is also true by the same axioms. It
contains no more information than the assumptions on which it rests. Clearly mathematics is an
important language for expressing science.

1.6 HOW TO USE THE TEXT

Readers should not expect to understand or appreciate everything in this book in the short term.
Many subtle and deep lying connections are sewn in these pages that will take even the most
experienced reader some time to unravel. It is my hope that there are issues sketched out here
that will provide fodder for research for at least a decade, probably several. Many ideas about the
administration of systems are general and have been discussed many times in different contexts,
but not in the manner or context of system administration.

The text can be read in several ways. To gain a software-engineering perspective, one can
replace “the system” with “the software”. To gain a business management perspective, replace “the
system” with “the business”, or “the organization”. For human-computer administration, read “the
system” as “the network of computers and its users”.

The first part of the book is about observing and recording observations about systems, since we
aim to take a scientific approach to systems. Part 2 concerns abstracting and naming the concepts
of a system’s operation and administration in order to place them into a formal framework. In the
final part of the book, we discuss the physics of information systems, i.e. the problem of how to
model the time-development of all the resources in order to determine the effect of policy. This
reflects the cycle of development of a system:

- Observation.
- Design (change).
- Analysis.

1.7 SOME NOTATION USED

A few generic symbols and notations are used frequently in this book and might be unfamiliar.

The function \( q(t) \) is always used to represent a ‘signal’ or quality that is varying in the system,
i.e. a scalar function describing any value that changes in time. I have found it more useful to call
all such quantities by the same symbol, since they all have the same status.

\( q(x, t) \) is a function of time and a label \( x \) that normally represents a spatial position, such as a
memory location. In structured memory, composed of multiple objects with finite size the addresses
are multi-dimensional and we write $q(\vec{x}, t)$, where $\vec{x} = (x_1, \ldots, x_\ell)$ is an $\ell$-dimensional vector that specifies location within a structured system, e.g. $(6,3,8)$ meaning perhaps bit 6 of component 3 in object 8.

In describing averages, the notation $\langle \ldots \rangle$ is used for mean and expectation values, e.g. $\langle X \rangle$ would mean an average over values of $X$. In statistics literature, this is often written $E(X)$.

In a local averaging procedure, reduces a large set $X$ to a smaller set $x$ of compounded objects, thus does not result in a scalar value but a smaller set whose elements are identified by a new label. E.g. suppose we start with a set of 10 values $X$. We could find the mean of all values $\langle X \rangle_{10}$ giving a single value. now group them into 5 groups of 2. Now we average each pair and end up with 5 averaged values: $\langle X(x) \rangle_2$. This still has a label $x$ since it is a set of values, where $x = 1 \ldots 5$.

Applications and Further Study 1.

- Use these broad topics as a set of themes for categorizing the detailed treatments in forthcoming chapters.
CHAPTER 2

SCIENCE AND ITS METHODS

Science is culture,
Technology is art.

– Author’s slogan.

A central theme of this book is the application of scientific methodologies to the design, understanding and maintenance of human-computer systems. Ironically ‘Computer Science’ has often lacked classical scientific thinking in favour of reasoned assertion, since it has primarily been an agent for technology and mathematics. The art of observation has concerned mainly those who work with performance analysis.

While mathematics is about reasoning (it seeks to determine logical relationships between assumed truths), the main purpose of science is to interpret the world as we see it, by looking for suitably idealized descriptions of observed phenomena and quantifying their uncertainty. Science is best expressed with mathematics, but the two are independent. There are many philosophies about the meaning of science, but in this book we shall be pragmatical rather than encyclopædic in discussing these.

2.1 THE AIM OF SCIENCE

Let us define science in a form that motivates its discussion in relation to human-computer systems.

Principle 1 (Aim of science). The principal aim of science is to uncover the most likely explanation for observable phenomena.
Science is a procedure for making sure that we know what we are talking about, when discussing phenomena that occur around us. It is about managing our uncertainty. Science does not necessarily tell us what the correct explanation for a phenomenon is, but it provides us with tools for evaluating the likelihood that a given explanation is true, given certain experimental conditions. Thus, central to science is the act of observation.

Observation is useless without interpretation, so experiments need theories and models to support them. Moreover, there are many strategies for understanding observable phenomena: it is not necessary to have seen a phenomenon to be able to explain it, since we can often predict phenomena just by guesswork, or imagination. The supposed explanation can then be applied and tested once the phenomenon has actually been observed.

The day-to-day routine of science involves the following themes, in approximately this order:

**Observation of phenomena**

Normally we want to learn something about a system, e.g. find a pattern of behaviour so that we might predict how it will behave in the future, or evaluate a property so that we can make a choice or value judgement about it. This might be as simple as measuring a value, or it might involve plotting a set of values in a graph against a parameter such as time or memory.

**Example 5.** Performance analysts measure the rate at which a system can perform its task. They do this with the larger aim of making things faster or more efficient. Computer anomaly detectors, on the other hand, look for familiar patterns of behaviour so that unusual occurrences can be identified and examined more closely for their significance.

**Estimation of experimental error**

In observing the world, we must be cautious about the possibility of error in procedure and interpretation: if we intend to base decisions of observations, we need to know how certain we are of our basis. Poor data can mislead (garbage in; garbage out). Any method of observation admits the possibility of error in relation to one’s assumptions and methods.

- We make a mistake in measurement (either at random or repeatedly).
- The measuring apparatus might be unreliable.
- The assumptions of the experiment are violated (e.g. inconstant environmental conditions)

Although it is normal to refer to this as ‘experimental error’, a better phrase is *experimental uncertainty*. We must quantify the uncertainty in the experimental process itself, because this

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1This is how black holes were ‘discovered’ in astrophysics. It is now believed that there is unambiguous evidence for black holes.
Figure 2.1: A pattern of process behaviour. The solid curve is the measured expectation value of the behaviour for that time of week. The error bars indicate the standard deviation, which also has a periodic variation that follows the same pattern as the expectation value; i.e. both moments of the probability distribution of fluctuations has a daily and weekly period.
CHAPTER 2. SCIENCE AND ITS METHODS

contributes an estimation of how correct our speculations about the results are. Uncertainties are usually plotted as ‘error bars’ (see fig 2.1).

IDENTIFICATION OF RELATIONSHIPS

Once we know the main patterns of behaviour, we try to quantify them by writing down mathematical relationships. This leads to empirical relationships between variables, i.e. it tells us how many of the variables we are able to identify are independent, and how many are determined.

Example 6. It is known that the number of processes running on a college web server is approximately a periodic function (see fig. 2.1). Using these observations, we could try to write down a mathematical relationship to describe this. e.g.

$$f(t) = A + Be^{-\gamma(t-t_0)} \sin(\omega t),$$  

(2.1)

where \(t\) is time along the horizontal axis, and \(f(t)\) is the value on the vertical axis, for constants \(A, B, \omega, \gamma, t_0\).

In the example above, there are far too many parameters to make a meaningful fit. It is always possible to fit a curve to data with enough parameters (‘enough parameters to fit an elephant’ is a common phrase used to ridicule students); the question is how many are justified before an alternative explanation is warranted?

SPECULATION ABOUT MECHANISMS

Expressing observations in algebraic form gives us a clue about how many parameters are likely to lie behind the explanation of a phenomenon. Next we speculate about the plausible explanations that lead to the phenomena, and formulate a theory to explain the relationships. If our theory can predict the relationships and data we have provided, it is reasonable to call the speculation a theory.

CONFIRMATION OF SPECULATIONS

One must test a theory as fully as possible by comparing it to existing observations, and by pushing both theory and observation to try to predict something that we do not already know.

QUANTIFICATION OF UNCERTAINTY

In comparing theory and observation, there is much uncertainty. There is a basic uncertainty in the data we have collected, then there is a question of how accurately we expect a theory to reproduce those data.
Example 7. Suppose the formula above for fig. 2.1, in eqn. (2.1) can be made to reproduce the data to within twenty percent of the value on either side, i.e. the approximate form of the curve is right, but not perfect. Is this an acceptable description of the data? How close do we have to be to say that we are close enough? This ‘distance from truth’ is our uncertainty.

In a clear sense, science is about uncertainty management. Nearly all systems of interest (and every system involving humans) are very complex and it is impossible to describe them fully. Science’s principle strategy is therefore to simplify things to the point where it is possible to make some concrete characterizations about observations. We can only do this with a certain measure of uncertainty. To do the best job possible, we need to control those uncertainties. This is the subject of the next chapter.

2.2 Causality, Superposition and Dependency

In any dynamical system where several processes can coexist, there are two possible extremes:

- Every process is independent of every other. System resources change additively (linearly) in response to new processes.
- The addition of each new process affects the behaviour of the others in a non-additive (non-linear) fashion.

The first case is called superposition, i.e. that two processes can coexist without interfering. This is not true or possible in general, but it can be a useful viewpoint for approximating some system regimes. The latter case is more general and often occurs when a system reaches some limitation, or constraint on its behaviour, such as when there is contention over which process has the use of critical resources.

The principle of causality governs all systems at a fundamental level. It is simply stated:

**Principle 2 (Causality). Every change or effect happens in response to a cause, which precedes it.**

This principle sounds intuitive and even manifestly obvious, but the way in which cause and effect are related in a dynamical system is not always as clear as one might imagine. We would often like to be able to establish a causal connection between a change of a specific parameter and the resulting change in the system. This is a central skill in fault finding, for instance; however, such causal links are very difficult to determine in complex systems. This is one of the reasons why the administration of systems is hard.
2.3 CONTROVERSIES AND PHILOSOPHIES OF SCIENCE

Science and philosophy have long been related. Indeed, what we now call science was once ‘natural philosophy’, or pondering about the natural world. Those who practice science today tend to think little about its larger meaning, or even its methodology. Science has become an ‘industry’ — the high ideals that were afforded it in the seventeenth century have since been submerged in the practicalities of applying it to real problems.

Here are some assertions that have been made of science by philosophers ([Hor96]):

- “Science cannot determine the truth of an explanation, only its likelihood”.
- “Science can only determine the falsity of a theory, not whether it is true”.
- “We must distinguish between truth, which is objective and absolute, and certainty which is subjective”.

To the casual technologist, such assertions are likely to draw only scepticism as to the value of philosophy. However, those willing to reflect more deeply on the whole investigative enterprise will find many ideas in the philosophy of science that are both interesting and of practical importance. The difficulty in presenting the labours of careful thought in such a brief and summarial form is that it is easy to misrepresent the philosophers’ detailed arguments. No doubt they would be horrified by this summary if they were alive to read it.

One of the first modern philosophers of science was Sir Francis Bacon, of the sixteenth century. Bacon (who died of pneumonia after stuffing a chicken with ice to see if it would preserve its flesh — thus anticipating the deep-freeze) maintained that the task of science is to uncover a thing’s character, by noting the presence or absence of tell-tale qualities. Thus, to understand heat, for instance, we must examine a list of hot and cold things and discern what features are relevant and irrelevant to the production of heat; e.g. exposure to sunlight is relevant, but the width of an object is not. Next we would examine instances in which a phenomenon is present in varying degrees, noting what circumstances also vary. For example, to understand heat we must observe things at different temperatures and note what circumstances are present in varying degrees. Bacon recognized that we cannot examine an endless number of instances: at some point we must stop and survey the instances so far.

Especially in the seventeenth century, philosophy became intertwined with mathematics, or analytical thinking. The philosopher Descartes used geometry for his inspiration as to how best to conduct an impartial inquiry. John Locke, an understudy of Isaac Newton, hoped to draw inspiration from the phenomenal success of Newton’s laws of motion and the calculus, and derive an analytical way of addressing a ‘method of inquiry’ — what, today, we would call a ‘scientific

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2 At this point it would be natural to give a reference to a book in which a nice summary was presented. Alas, I have yet to find a clear exposition of the philosophy of science printed in English.
method’. His philosophy, now called *empiricism*, implies a reliance on experience as the source of ideas and knowledge.

Newton was a significant source of inspiration to philosophers because, for the first time, his work had made it possible to calculate the outcome of a hypothetical situation that no one had ever observed before, i.e. predict the future for idealized physical systems. During the Enlightenment, philosophers even came to believe that scientific inquiry could yield truths about human nature and thus that ethical principles might be best derived from such truths; this would therefore be a basis for a new order of society.

In the eighteenth century, others began to realize that this vision was flawed. David Hume discovered an important twist, namely that predictions about events that are not observed cannot be *proven* to be true or false, nor even to be probable, since observation alone cannot see into the future, and not not attempt to asses the *cause* of a phenomenon. He asserted that there are two sources of knowledge: analytical knowledge that is certain (provable assertions) but which cannot directly represent reality, and empirical knowledge or observations that are uncertain but which apply to the real world.

The empirical observation that releasing a stone causes it to fall to the ground is insufficient to prove, beyond doubt, that every stone will always fall to the ground in the future. This is a good example of how our limited experience shapes our view of the world. Before humans went into space, the assertion was always true; however, away from gravity, in the weightlessness of space, the observation becomes meaningless. Hume’s point is that we do not know what we don’t know, so we should not make unwarranted assumptions.

Although Hume’s ideas had an impact on philosophy, they were not generally accepted in science. Immanuel Kant and John Stuart Mill made attempts to solve some of Hume’s problems. Kant claimed to solve some of them by assuming that certain facts were to be regarded as axioms, i.e. articles of faith that were beyond doubt; i.e. that one should always set the stage by stating the conditions under which conclusions should be deemed “true”.

Kant supposed, moreover, that our perception of the world is important to how we understand it. In what sense are things real? How do we know that we are not imagining everything? Thus, how do we know that there are not many equally good explanations for everything we see? His central thesis was that the possibility of human knowledge presupposes the participation of the human mind. Instead of trying, by reason or experience, to make our concepts match the nature of objects, Kant held, we must allow the structure of our concepts shape our experience of objects.

Mill took a more pragmatic line of inquiry and argued that the truth of science is not absolute, but that its goals were noble; i.e. science is a self-correcting enterprise that does not need axiomatic foundations per se. If experience reveals a flaw in its generalities, that can be accommodated by a critical revision of theory. It would eventually deal with its own faults by a process of refinement.

*Epistemology* is a branch of philosophy that investigates the origins and nature, and the extent
of human knowledge. Although the effort to develop an adequate theory of knowledge is at least as old as Plato, epistemology has dominated Western philosophy only since the era of Descartes and Locke, largely as an extended dispute between rationalism and empiricism. Rationalism believes that some ideas or concepts are independent of experience and that some truth is known by reason alone (e.g. parallel lines never meet). Empiricism believes truth must be established by reference to experience alone.

Logical positivism is a twentieth-century philosophical movement that used a strict principle of verifiability to reject non-empirical statements of metaphysics, theology, and ethics. Under the influence of Hume and others, the logical positivists believed that the only meaningful statements were those reporting empirical observations. The tautologies of logic and mathematics could not add to these, but merely re-express them. It was thus a mixture of rationalism and empiricism.

The verifiability principle is the claim that the meaning of a proposition is no more than the set of observations that would determine its truth, i.e. that an empirical proposition is meaningful only if it either actually has been verified or could at least in principle be verified. Analytic statements (including mathematics) are non-empirical; their truth or falsity requires no verification. Verificationism was an important element in the philosophical program of logical positivism.

One of the most influential philosophers of science is Karl Popper. He is sometimes referred to as the most important philosopher of science since Francis Bacon. Karl Popper’s ideas have proven to be widely influential for their pragmatism and their belief in the rational. Popper rejected that knowledge is a social phenomenon — it is absolute. He supposed that we cannot be certain of what we see, but if we are sufficiently critical we can determine whether or not we are wrong, by deductive falsification, or a process of conjecture and refutation.

Popper believed that theories direct our observations. They are part of our innate desire to impose order and organization on the world, i.e. to systematize the phenomena we see, but we are easily fooled and therefore we need to constantly criticize and retest every assumption to see if we can falsify them. Hume said we can never prove them right, but Popper says that we can at least try to see if they are wrong.

Paul Feyerabend later argued that there is no such thing as an objective scientific method. He argued that what makes a theory true or false is entirely a property of the world-view of which that assertion is a part. This is relativism, i.e. objectivity is a myth. We are intrinsically locked into our own world view, perceiving everything through a particular filter, like a pair of sunglasses that only lets us see particular things.

We need only one flaw in an explanation to discount it; but we might need to confirm hundreds of facts and details to be sure about its validity i.e. “truth”. In the context of this book, science itself is a system that we shall use to examine others. We summarize with a pragmatic view of science:

| Principle 3 (Controlled environment) | Science provides an impartial method for investigating and describing phenomena within an idealized environment, under controlled conditions. |
CHAPTER 2. SCIENCE AND ITS METHODS

Science, we claim, is an investigative enterprise, whose aim is to characterize what is already there. Technology, on the other hand is a creative enterprise: it is about tool-building.

The relationship between science and technology is often presented as being problematical by technologists, but it is actually quite clear. If we do not truly understand how things work and behave, we cannot use those things to design tools and methods. In technology we immediately hit upon an important application of science, namely its role in making value judgements. A value judgement is a subjective judgement, e.g. one tool can be better than another, one system or method can be better than another — but how are such judgements made? Science cannot answer these questions, but it can assist in evaluating them, if the subjectivity can be defined clearly.

The situation is somewhat analogous to that faced by the seventeenth century philosophers who believed that ethics could be derived from scientific principles. Science cannot tell us whether a tool or a system is “good” or “bad”, because “good” and “bad” have no objective definitions. Science craves a discipline in making assertions about technology, and perhaps even guides us in making
improvements in the tools we make, by helping us to clarify our own thoughts by quantification of technologies.

2.5 HYPOTHESES

Although science sometimes springs from serendipitous discovery, its systematic content comes from testing existing ideas or theories and assertions. Scientific knowledge advances by undertaking a series of studies, in order to either verify or falsify a hypothesis. Sometimes these studies are theoretical, sometimes they are empirical and frequently they are a mixture of the two. Statistical reproducibility is an important criterion for any result, otherwise it is worthless, because it is uncertain. We might be able to get the same answer twice by accident, but only repeated verification can be trusted.

In system administration, software tools and human methods form the technologies that are used. Progress in understanding is made, with the assistance of the tools only if investigation leads to a greater predictive power or a more efficient solution to a problem.

- Scientific progress is the gradual refinement of the conceptual model that describes the phenomenon we are studying. In some cases, we are interested in modelling tools. Thus technology is closely related to science.

- Technological progress is the gradual creative refinement of the tools and methods referred to by the technology. In some cases, the goal is the technology itself, in other situations the technology is only an implement for assisting the investigation.

All problems are pieces of a larger puzzle. A complete scientific study begins with a motivation, followed by an appraisal of the problems, the construction of a theoretical model for understanding or solving the problems, and finally an evaluation or verification of the approach used and the results obtained. Recently much discussion has been directed towards finding suitable methods for evaluating technological innovations in computer science as well as to encouraging researchers to use them. Nowadays many computing systems are of comparable complexity to phenomena found in the natural world and our understanding of them is not always complete, in spite of the fact that they were designed to fulfil a specific task. In short technology might not be completely predictable, hence there is a need for experimental verification.

2.6 THE SCIENCE OF TECHNOLOGY

In technology the act of observation has two goals: i) to gather information about a problem in order to motivate the design and construction of a technology which solves it, and ii) to determine
CHAPTER 2. SCIENCE AND ITS METHODS

whether or not the resulting technology fulfils its design goals. If the latter is not fulfilled in a technological context, the system may be described as faulty, whereas in natural science there is no right or wrong. In between these two empirical book-marks lies a theoretical model which hopefully connects the two.

System administration is a mixture of science, technology and sociology. The users of computer systems are constantly changing the conditions for observations. If the conditions under which observations are made are not constant, then the data lose their meaning: the message we are trying to extract from the data is supplemented by several other messages which are difficult to separate from one another. Let us call the message we are trying to extract signal and the other messages which we are not interested in noise. Complex systems are often characterized by very noisy environments.

In most disciplines one would attempt to reduce or eliminate the noise in order to isolate the signal. However, in system administration, it would be no good to eliminate the users from an experiment, since it is they who cause most of the problems which one is trying to solve. In principle this kind of noise in data could be eliminated by statistical sampling over very long periods of time, but in the case of real computer systems this might not be possible since seasonal variations in patterns of use often lead to several qualitatively different types of behaviour which should not be mixed. The collection of reliable data might therefore take many years, even if one can agree on what constitutes a reasonable experiment. This is often impractical, given the pace of technological change in the field.

2.7 EVALUATING A SYSTEM - DEPENDENCIES

Evaluating a model of system administration is a little bit like evaluating the concept of a bridge. Clearly a bridge is a structure with many components each of which contributes to the whole. The bridge either fulfils its purpose in carrying traffic past obstacles or it does not. In evaluating the bridge, should one then consider the performance of each brick and wire individually? Should one consider the aesthetic qualities of the bridge? There might be many different designs each with slightly different goals. Can one bridge be deemed better than another on the basis of objective measurement? Perhaps only the bridge’s maintainer is in a position to gain a feeling for which bridge is the most successful, but the success criterion might be rather vague: a collection of small differences which make the perceptible performance of the bridge optimal, but with no measurably significant data to support the conclusion. These are the dilemmas of evaluating a complex technology.

The options we have for performing experimental studies are,

- Measurements.
- Simulations.
• User surveys. with all of the incumbent difficulties which these entail.

SIMPLICITY

Conceptual and practical simplicity are often deemed to be positive attributes of systems and procedures. This is because simple systems are easy to understand and their behaviours are easy to predict. We prefer that systems that perform a function do so predictably.

EVALUATION OF INDIVIDUAL MECHANISMS

For individual pieces of a system, it is sometimes possible to evaluate the efficiency and correctness of the components. Efficiency is a relative concept and, if used, it must be placed in a context. For example, efficiency of low level algorithms is conceptually irrelevant to the higher levels of a program, but it might be practically relevant. i.e. one must say what is meant by efficiency before quoting results. The correctness of the results yielded by a mechanism/algorithm can be measured in relation to its design specifications. Without a clear mapping of input/output the correctness of any result produced by a mechanism is a heuristic quality. Heuristics can only be evaluated by experienced users expressing their informed opinions.

2.8 ABUSES OF SCIENCE

Science is about constantly asking questions and verifying hypotheses to see if one’s world view holds up to scrutiny. However, the authority that science has won is not always been wielded in a benign way. History is replete with illegitimate ideas that have tried to hide behind the reputation of science, by embracing its terminology without embracing its forms.

Marketeers are constantly playing this game with us, inventing scientific sounding names for bells and whistles on their products, or claiming that they are ‘scientifically proven’ (an oxymoron). By quoting numbers, or talking about ‘ologies’ there are many uncritical forces in the world who manipulate our beliefs, assuming that most individuals will not be able to verify them or discount them\(^3\). In teaching a scientific method, we must be constantly aware of abuses of science.

\(^3\)Eugenics is one classic example where the words and concepts discovered by science were usurped for illegitimate means to claim that certain individuals were genetically superior to others. This was a classic misunderstanding of a scientific concept that was embraced without proper testing or understanding.
Applications and Further Study 2. The observation and analysis of systems involves these themes:

- Variables or measurables.
- Determinism or causality.
- Indeterministic, random or stochastic influences.
- Systems and their environments.
- Accounting and conservation.
CHAPTER 3

EXPERIMENT AND OBSERVATION

Trust, but verify!

–Russian Proverb

Collecting data to support an idea or hypothesis is central to the scientific method. We insist on the existence of evidence that can be examined and related analytically (by mathematics or other reasoning) to the phenomenon under consideration, because our trust in random observation or hearsay is only limited. The paraphrased proverb, “Trust but verify” is often cited in connection with system security, but it is equally pertinent here. In a sense, the scientific method is the security or quality assurance system for ‘truth’.

To study human-computer systems, we draw on analytical methods from the diverse branches of science, but our conclusions must be based on observed fact. Reliable observational evidence is most easily obtained where one can perform experiments to gather numerical data, then derive relationships and conclusions. Descriptive sciences do not always have this luxury and are forced to use a form of data collection that involves visual observation, classification or even by interview. This is less focused and therefore harder to use to support specific conclusions.

Example 8. A zoologist might find no problem in measuring the weight of animals, but might find it difficult to classify the colours of animals in order to relate this to their behaviour. When is red really brown? Fuzzy classifiers from day-to-day experience lead to difficulties for science — qualitative descriptions are prone to subjective interpretation.

Example 9. In human-computer systems, it is easy to measure numerical quantities such as rate of change of data, but qualitative features such as ‘lawfulness’ of users seem too vague to quantify.
Difficulties with qualitative characterizations can sometimes be eliminated by going to a lower level, or to a smaller scale of the system: e.g. the classification of animals might be done more precisely by looking at their DNA, and the lawfulness of a user might be measured by examining the policy conformance of each file and change made by the user.

3.1 DATA PLOTS AND TIME SERIES

In the observation of real systems, measurements are made and data are collected. If the data are collected at regular intervals, they are usually represented either as time-series, i.e. plots of a measured values versus the time at which the measurements were made, or as histograms that count the numbers of measurements that fall into certain domains (called classes) in the data values. Both types of diagram play important roles in understanding systems. In addition, various kinds of graphical representations are used to elucidate relationships between variables, such as plots of one variable against another, or log-log plots of the same variables that indicate power-law relationships.

Fig. 3.1 shows a typical series of measurements made from a computer system over the course of several weeks. By plotting all of the data on a against a timescale of a week, one sees a clear pattern in the data, but also a scatter in the values measured at each time. Error bars are drawn at each point where there are repeated measurement. These show the width of the standard deviation $\pm \sigma$ centred about the mean value. It is important to plot the scatter in data as a visual guide to the uncertainty in the claimed result (see section 3.4).

Since, at each each time, in fig. 3.1 there is a distribution of values, we can plot that distribution on a frequency plot like that in fig. 3.2. This is a kind of histogram in which the columns have been joined into an approximate curve. If the area under a frequency plot like this is normalized to unity, it represents a probability distribution for the measured values $P(q)$. The probability distribution for the measured values is important in gauging the stability of a system as well as in characterizing its fluctuation spectrum, as we shall see in chapter 8.

3.2 CONSTANCY OF ENVIRONMENT DURING MEASUREMENT

In science, our aim is to take small steps, by stripping away everything down to single cause-effect relationships, and then gradually putting things back together. Einstein is famous for having said that everything in should be made as simple as possible, but no simpler. By this, he meant that we should neither over-complicate nor over-simplify an explanation.

Most phenomena are governed by a number of parameters; e.g. suppose the rate of a computer is affected by three parameters:

$$R = R(c, m, s)$$  \hfill (3.1)
where $c$ is the CPU rate, $m$ is the amount of memory and $s$ is the speed of memory. If we want to discover just how $R$ depends on each of these, we must test each parameter individually, holding the others constant, else we might mix up the dependence on each parameter. Science ends up with neat formulae relating measurables, only because this isolation is possible. Such formulae describe the real world, but they do not really describe the ‘real environment’ because the environment is messy. Science therefore strives to ensure idealized environmental conditions for investigating phenomena, in order to take one thing at a time.

In the real world of human-computer systems, there are many variables and influences that affect a system, so we must strive to maintain constant conditions in all variables but the one we would like to test. This is rarely possible, and thus there is an inevitable uncertainty or experimental error in any experiment. An important task of science is to quantify this uncertainty.
CHAPTER 3. EXPERIMENT AND OBSERVATION

Principle 4. Scientific observation strives to isolate single cause-effect relationships, by striving to keep environmental conditions constant during measurement. The impossibility of completely constant external conditions makes it necessary to quantify the uncertainty in each measurement.

Figure 3.2: A frequency plot of the numbers of measurements of a given value.

Note that by isolating ‘single’ cause-effect relationships, we do not mean to imply that there is always a single variable that controls a process, only that each independent change can be identified with an independent parameter.

The way we do this for simple measurable values is relatively easy and is described in this chapter. However, not all situations are so easily quantifiable. Qualitative experiments, such as those of biology (e.g. classifying types of behaviour) also occur in the study of human-computer systems. If we do not actually begin with hard numbers, the estimate of uncertainty has to be made by finding a numerical scale, typically through a creative use of classification statistics; e.g. how many animals have exhibited behaviour $A$ and how many behaviour $B$? Or how far is behaviour $A$ from behaviour $B$ on some arbitrary scale, used only for comparison.

All scales are arbitrary in science (that is why we have many different units for weight, height,
frequency etc), what is important is how we relate these scales to observables.

### 3.3 Experimental Design

The cleverness of an experiment’s design can be crucial to its success in providing the right information. Our aim is to isolate a single channel of cause-effect at a time. We must ensure that the experimental observation does not interfere with the system we are measuring. Often an experiment yields unexpected obstacles which must be overcome. There can be a lot of work to answer even a simple question. (For examples from computer performance analysis, see [Jai91].)

**Example 10.** Suppose we wish to compare the behaviour of two programs for mirroring (copying) files, for backup. We notice that one program seems to complete its task very quickly, presenting a high load to the source and destination machines. The other takes much longer but presents almost no load. How shall we determine the reason?

We might begin by finding some data to copy. Data are composed of files of different sizes. Size might be important, so we shall be interested in how size affects the rate of copying, if at all. The first time we copy the files, every file must be transferred in full. On subsequent updates, only changes need to be copied. One program claims to copy only those bytes that are different; the other has to copy a whole file, even if only one byte has changed, so file size again becomes important.

We could investigate how the total time for copying is related to the total amount of data i) of all files, ii) of files that are copied. We might also be interested in what dependencies the programs have: do they use the Internet Protocol with TCP or UDP, IPv4 or IPv6? Does the computer kernel or operating system affect the performance of the two programs?

The stream of questions never ceases; we must decide when to stop. Which questions are we interested in, and when have they been sufficiently answered? This is a value judgement that requires experience and inquisitiveness from the investigator.

### 3.4 Stochastic (Random) Variables

Our inability to control, or even follow every variable in a system’s environment means that some of the changes appearing in the system seem random, or inexplicable.

**Definition 4 (Random process).** A random process is one in which there are too many unknowns to be able to trace the channels of cause and effect.

A stochastic or random variable is a variable whose value depends on the outcome of some underlying random process. The range of values of the variable is not at issue, but which particular
value the variable has at a given moment is random. We say that a stochastic variable \( X \) will have a certain value \( x \) with a probability \( P(x) \).

Usually, in an experiment a variable can be said to have a certain random component (sometimes called its ‘error’ from the historical prejudice that science is deterministic and the only source of randomness is the errors incurred by the experimental procedure) and an average stable value. We write this

\[
x = \langle x \rangle + \Delta x,
\]

where \( x \) is the actual value measured, \( \langle x \rangle \) is the mean or expectation value of all measurements (often written \( E(x) \) in statistical literature), and \( \Delta x \) is the deviation from the mean. The mean value changes much more slowly than \( \Delta x \). For example:

- Choices made by large numbers of users are not predictable, except on average.
- Measurements collected over long periods of time are subject to a variety of fluctuating conditions.

Measurements can often appear to give random results, because we do not know all of the underlying mechanisms in a system. We say that such systems are non-deterministic or that there are hidden variables that prevent us from knowing all the details. If a variable has a fixed value, and we measure it often enough and for long enough, the random components will often fall into a stable distribution, by virtue of the central limit theorem (see for instance ref. [GS01]). The best known example of a stable distribution is the Gaussian type of distribution.

### 3.5 Actual Values or Characteristic Values

There is a subtle distinction in measurement between an observable that has an actual ‘true’ value and one that can only be characterized by a typical value.

For example, it is generally assumed that the rest mass of the electron has a ‘true’ value that never changes. Yet when we measure it, we get many different answers. The conclusion must be that the different values result from errors in the measurement procedure. In a different example, we can measure the size of a whale and we get many different answers. Here there is not ‘true’ or ‘standard’ whale and the best we can do is to measure a typical or expected value of the size.

In human-computer systems, there are few if any measurements of the first type, because almost all values are affected by some kind of variation. For example, room temperature can alter the maximum transmission rate of a cable. We must therefore be careful about what we claim to be constant, and what is the reason for the experimental variation in the results. Part of the art in science is in the interpretation of results, within the constraints of cause and effect.
CHAPTER 3. EXPERIMENT AND OBSERVATION

3.6 OBSERVATIONAL ERRORS

All measurements involve certain errors. One might be tempted to believe that, where computers are involved, there would be no error in collecting data, but this is false. Errors are not only a human failing, they occur because of unpredictability in the measurement process, and we have already established throughout this book that computers systems can be unpredictable. We are thus forced to make estimates of the extent to which our measurements can be in error. This is a difficult matter, but approximate statistical methods are well known in the natural sciences, methods which become increasingly accurate with the amount of data in an experimental sample.

The ability to estimate and treat errors should not be viewed as an excuse for constructing a poor experiment. Errors can only be minimized by design. There are several distinct types of error in the process of observation.

The simplest type of error is called random error. Random errors are usually small deviations from the ‘true value’ of a measurement which occur by accident, by unforeseen jitter in the system, or some other influence. By their nature, we are usually ignorant of the cause of random errors, otherwise it might be possible to eliminate them. The important point about random errors is that they are distributed evenly about the mean value of the observation. Indeed, it is usually assumed that they are distributed with an approximately normal or Gaussian profile about the mean. This means that there are as many positive as negative deviations and thus random errors can be averaged out by taking the mean of the observations.

It is tempting to believe that computers would not be susceptible to random errors. After all, computers do not make mistakes. However this is an erroneous belief. The measurer is not the only source of random errors. A better way of expressing this is to say that random errors are a measure of the unpredictability of the measuring process. Computer systems are also unpredictable, since they are constantly influences by outside agents such as users and network requests.

The second type of error is a personal error. This is an error which a particular experimenter adds to the data unwittingly. There are many instances of this kind of error in the history of science. In a computer controlled measurement process, this corresponds to any particular bias introduced through the use of specific software, or through the interpretation of the measurements.

The final and most insidious type of error is the systematic error. This is an error which runs throughout all of the data. It is a systematic shift in the true value of the data, in one direction, and thus it cannot be eliminated by averaging. A systematic error leads also to an error in the mean value of the measurement. The sources of systematic error are often difficult to find, since they are often a result of misunderstandings, or of the specific behaviour of the measuring apparatus.

In order to measure the CPU usage of a computer system, for instance, we have to start a new program which collects that information, but that program inevitably uses the CPU also and therefore changes the conditions of the measurement. These issues are well known in the physical sciences and are captured in principles such as Heisenberg’s Uncertainty Principle, Schrödinger’s...
cat and the use of infinite idealized heat baths in thermodynamics. We can formulate our own verbal expression of this for computer systems:

**Principle 5** (Uncertainty). *The act of measuring a given quantity in a system with finite resources, always changes the conditions under which the measurement is made, i.e. the act of measurement changes the system.*

**Example 11.** *In order to measure the pressure of a bicycle-tyre, we have to release some of the pressure. If we continue to measure the pressure, the tyre will eventually be flat.*

**Example 12.** *In measuring the load on a computer system, most monitoring systems start a process which consumes a measurably significant amount of memory, and which forces the system to process kernel requests that steal resources from the existing system. If the goal is observational consistency, for diagnostic clarity, monitoring should be present in a predictable and non-intrusive way from at all times evenly, not spawned upon request, even if the latter might minimize load.*

The larger the available resources of the system, compared to the resources required to make the measurement, the smaller the effect on the measurement will be.

### 3.7 The Mean and Standard Deviation

In the theory of errors, we use the ideas above to define two quantities for a set of data: the mean and the standard deviation. Contrary to what one sometimes reads, these quantities are not necessarily tied to the normal distribution: they are just expressions of scale that can be used to characterize data sets. They are also called the first and second moments of the data.

The situation is now as follows: we have made a number \( N \) of observations of values \( v_1, v_2, v_3, \ldots, v_N \), which have a certain randomness and we are trying to find out a characteristic value \( v \) for the measurement. Assuming that there are no systematic errors, i.e. assuming that all of the deviations have independent random causes, we define the value \( \langle v \rangle \) to be the arithmetic mean of the data:

\[
\langle v \rangle = \frac{v_1 + v_2 + \cdots + v_N}{N} = \frac{1}{N} \sum_{i=1}^{N} v_i. \tag{3.3}
\]

Next we treat the deviations of the actual measurements as our guesses for the error in the measurements:

\[
\Delta g_1 = \langle v \rangle - v_1 \\
\Delta g_2 = \langle v \rangle - v_2 \\
\vdots \\
\Delta g_N = \langle v \rangle - v_N
\]
and define the *standard deviation* of the data by

\[ \sigma = \sqrt{\frac{1}{N} \sum_{i=0}^{N} \Delta g_i^2}. \]  

(3.4)

This is clearly a measure of the scatter in the data due to random influences. \( \sigma \) is the root mean square (RMS) of the assumed errors. These definitions are a way of interpreting measurements, from the assumption that one really is measuring the true value, affected by random interference.

**Definition 5** (Gaussian signal power). *A random signal that is distributed according to a Gaussian distribution has a characteristic amplitude \( \sigma \), and thus a squared amplitude of \( \sigma^2 \). Since the squared amplitude of a signal is associated with the power (in Watts) of a physical signal, the variance is often assumed to measure power.*

An example of the use of standard deviation can be seen in the error bars of the figures in this chapter. Whenever one quotes an average value, the number of data and the standard deviation should also be quoted in order to give meaning to the value. In system administration, one is interested in the average values of any system metric which fluctuates with time.

## 3.8 Probability Distributions and Measurement

When ever we repeat a measurement and obtain different results, a distribution of different answers is formed. The spread of results needs to be interpreted. There are two possible explanations for a range of values:

- The quantity being measured does not have a fixed value.
- The measurement procedure is imperfect and incurs a range of values due to error or uncertainty.

Often both of these are the case. In order to give any meaning to a measurement, we have to repeat the measurement a number of times and show that we obtain approximately the same answer each time. In any complex system, in which there are many things going on which are beyond our control (read: just about anywhere in the real world), we will never obtain exactly the same answer twice. Instead we will get a variety of different answers which we can plot as a graph: on the \( x \)-axis, we plot the actual measured value and on the \( y \)-axis we plot the number of times we obtained that measurement divided by a normalizing factor, such as the total number of measurements. by drawing a curve through the points, we obtain an idealized picture which shows the probability of measuring the different values.
Figure 3.3: The scatter is an estimate of the width of the populated regions of the probability distribution.

Over time measurements often develop stable average behaviour, so that a time-series \( x = \{x_1, x_2, x_3, \ldots\} \) has an average that tends towards a stable value. This is written in a variety of notations in the literature:

\[
\bar{x} = E(x) = \langle x \rangle \equiv \frac{1}{N} \sum_{i=1}^{N} x_i \to \mu,
\]

where \( N \) is the number of data. Few if any of the actual measurements will actually be equal to \( \mu \); rather they are scattered around the average value in some pattern, called their distribution \( P(x) \). The normalization factor is usually chosen so that the area under the curve is unity, giving a probabilistic interpretation.

**Definition 6** (Probability). The probability \( P(x) \) of measuring a value \( x \) in original data set is defined to be the fraction of values that fell into the range \( x \pm \Delta x/2 \), for some class width \( \Delta x \).

\[
P(x) = \frac{N(x - \Delta x/2, x + \Delta x/2)}{N_{\text{total}}}. \tag{3.6}
\]

Here \( N(x, y) \) is the number of observations between \( x \) and \( y \).

This probability distribution is the histogram shown in fig. 3.3.

There are two extremes of distribution: complete certainty (figure 3.4) and complete uncertainty (figure 3.5). If a measurement always gives precisely the same answer, then we say that there is no error. This is never the case is real measurements. Then the curve is just a sharp spike at the particular measured value. If we obtain a different answer each time we measure a quantity, then there is a spread of results. Normally that spread of results will be concentrated around some more or less stable value (figure 3.6). This indicates that the probability of measuring that value is biased, or tends to lead to a particular range of values. The smaller the range of values, the closer we approach figure 3.4. But the converse might also happen: in a completely random system, there might be no fixed value of the quantity we are measuring. In that case, the measured value is completely uncertain, as in figure 3.5. To summarize, a flat distribution is unbiased, or completely
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Figure 3.4: The delta distribution represents complete certainty. The distribution has a value of 1 at the measured value.

Figure 3.5: The flat distribution is a horizontal line indicating that all measured values, within the shown interval, occur with equal probability.

random. A non flat distribution is biased, or has an expectation value, or probable outcome. In the limit of complete certainty, the distribution becomes a spike, called the *delta distribution*.

We are interested in determining the shape of the distribution of values on repeated measurement for the following reason. If the variation of the values is symmetrical about some preferred value, i.e. if the distribution peaks close to its mean value, then we can likely infer that the value of the peak or of the mean is the true value of the measurement and that the variation we measured was due to random external influences. If, on the other hand, we find that the distribution is very asymmetrical, some other explanation is required and we are most likely observing some actual physical phenomenon which requires explanation.
Figure 3.6: Most distributions peak at some value, indicating that there is an expected value (expectation value) which is more probable than all the others.

3.8.1 SCATTER AND JITTER

The term *scatter* is often used to express the amount of variation in the measurements about the mean. It is estimated as the ‘width’ of the histogram $P(x)$. The term *jitter* is often used when describing the scatter of arrival times between measurements in the time series. Decades of artificial courses on statistics have convinced many scientists that the distribution of points about the mean must follow a Gaussian ‘normal’ distribution in the limit of large numbers of measurements. This is not true however: there are ample cases where the scatter is asymmetric or less uniform than the ‘normal distribution’.

3.8.2 THE ‘NORMAL’ DISTRIBUTION

It has been stated that ‘Everyone believes in the exponential law of errors; the experimenters because they think it can be proved by mathematics; and the mathematicians because they believe it has been established by observation’ ([WR29]). Some observational data in science satisfy closely the normal law of error, but this is by no means universally true. The main purpose of the normal error law is to provide an adequate idealization of error treatment that applies to measurements with a ‘true value’ (see section 3.5), which is simple to deal with, and which becomes increasingly accurate with the size of the data sample.

The normal distribution was first derived by DeMoivre in 1733, while dealing with problems involving the tossing of coins; the law of errors was deduced theoretically in 1783 by Laplace. He started with the assumption that the total error in an observation was the sum of a large number of independent deviations, which could be either positive or negative with equal probability, and could therefore be added according to the rule explained in the previous sections. Subsequently Gauss gave a proof of the error law based on the postulate that the most probable value of any number of equally good observations is their arithmetic mean. The distribution is thus sometimes called the
The Gaussian normal distribution, or bell curve, peaks at the arithmetic mean. Its width characterizes the standard deviation. It is therefore the generic model for all measurement distributions.

The Gaussian normal distribution is a smooth curve which is used to model the distribution of discrete points distributed around a mean. The probability density function $P(x)$ tells us with what probability we would expect measurements to be distributed about the mean value $\bar{x}$ (see figure 3.7).

$$P(x_i) = \frac{1}{(2\pi \sigma^2)^{1/2}} \exp\left(-\frac{(x_i - \bar{x})^2}{2\sigma^2}\right).$$

It is based on the idealized limit of an infinite number of points.

### 3.8.3 STANDARD ERROR OF THE MEAN

No experiments have an infinite number of points, so we need to fit a finite number of points to a normal distribution as well as we can. It can be shown that the most probable choice is to take the mean of the finite set to be our estimate of mean of the ideal set. Of course, if we select at random a sample of $N$ values from the idealized infinite set, it is not clear that they will have the same mean as the full set of data. If the number in the sample $N$ is large, the two will not differ by much, but if $N$ is small, they might. In fact, it can be shown that if we take many random samples of the ideal set, each of size $N$ that they will have mean values which are themselves normally distributed, with a standard deviation equal to $\sigma/\sqrt{N}$. The quantity

$$\alpha = \frac{\sigma}{\sqrt{N}}.$$
where $\sigma$ is the standard deviation, is therefore called the standard error of the mean. This is clearly a measure of the accuracy with which we can claim that our finite sample mean agrees with the actual mean. In quoting a measured value which we believe has a unique or correct value (e.g. the height of the Eiffel Tower), it is therefore normal to write the mean value, plus or minus the standard error of the mean:

$$\text{Result} = \bar{x} \pm \sigma / \sqrt{N} \quad (\text{for } N \text{ observations}),$$

where $N$ is the number of measurements. Otherwise, if we believe that the measured value should have a distribution of values (e.g. the height of a river on the first of January of each year), one uses the standard deviation as a measure of the error. Many transactional operations in a computer system do not have a fixed value (see next section).

The law of errors is not universally applicable, without some modification, but it is still almost universally applied, for it serves as a convenient fiction which is mathematically simple\(^1\).

### 3.8.4 Other Distributions

Another distribution which appears in the periodic rhythms of system behaviour is the exponential form. There are many exponential distributions, and they are commonly described in text books. Exponential distributions are used to model component failures in systems over time i.e. most components fail quickly or live for a long time.

The Planck distribution is one example that can be derived theoretically as the most likely distribution to arise from an assembly of fluctuations in equilibrium with a large source (see \[BHRS01\]). The precise reason for its appearance in computer systems is subtle, but has to do with the periodicity imposed by users’ behaviours, as well as the interpretation of transactions as fluctuations. The distribution has the form

$$D(\lambda) = \frac{\lambda^{-m}}{e^{\lambda/T} - 1},$$

where $T$ is a scale, and $m$ is usually an integer greater than 2. When $m = 3$, a single degree of freedom is represented. The shape of the graph is shown in figure 3.8.

Internet network traffic analysis studies (see \[PF95, WPT96\]) show that the arrival times of data packets within a stream has a long tailed distribution, often modelled as a Pareto distribution (a power law) in the asymptotic limit, for constants $\alpha$ and $\beta$:

$$f(\omega) = \beta \omega^{\alpha - 1}.$$

This can be contrasted with the Poissonian arrival times of telephonic data traffic. It is an important consideration to designers of routers and switching hardware. It implies that a fundamental change

---

\(^1\)The applicability of the normal distribution can, in principle, be tested with a $\chi^2$ test, but this is seldom used in physical sciences, since the number of observations is usually so small as to make it meaningless.
in the nature of network traffic has taken place. A partial explanation for this behaviour is that packet arrival times consist not only of Poisson random processes for session arrivals, but also of internal correlations within a session. Thus it is important to distinguish between measurements of packet traffic and measurements of numbers of sockets (or TCP sessions). The power law behaviour exhibited by Pareto tails is often indicative of clustered behaviour. If one event arrives, several tend to arrive in a cluster or burst.

### 3.9 Uncertainty in General Formulae

Suppose we measure the values of $N$ variables that feed into a mathematical expression for something:

$$S = S(x, y, z, \ldots)$$  \hspace{1cm} (3.9)

Assuming that errors are small, we can estimate the effect of an error in one of the parameters on the calculated expression by calculating the gradient (rate of change) of the function at the approximate value of the parameter and by multiplying this by our estimate of the error in the parameter. This tells us the expected error in $S$, given an estimate of the error in $x$. We use the first order Taylor expansion for each variable and then treat each contribution as an orthogonal perturbation and use Pythagoras formula to express the combined error. Knowing the errors $\Delta x$, 

![Chapter 3. EXPERIMENT AND OBSERVATION](image)

Figure 3.8: The Planck distribution for several temperatures. This distribution is the shape generated by random fluctuations from a source which is unchanged by the fluctuations. Here, a fluctuation is a computing transaction, a service request or new process.
\[ \Delta S \equiv \sqrt{\left( \frac{\partial S}{\partial x} \right)^2 \Delta x^2 + \left( \frac{\partial S}{\partial y} \right)^2 \Delta y^2 + \ldots} \quad (3.10) \]

**Example 13.** The average rate of user transactions per second in a database is given by

\[ S = R = \frac{N}{T}, \quad (3.11) \]

where \( N \) is the total number of transactions recorded, and \( T \) is the interval of time over which the measurement was made. We assume that the uncertainty in \( N \) is \( \Delta N \) (caused by the fact that we cannot exactly separate every user transaction from administrative transactions), and that the uncertainty in \( T \) is \( \Delta T \), (caused by not being able to tell the exact moment when the measurements started and stopped, due to context switching). Using the formula above we find that

\[ \frac{\partial S}{\partial N} = \frac{1}{T}, \quad \frac{\partial S}{\partial T} = -\frac{N}{T^2}, \quad (3.12) \]

so that

\[ \Delta S = \sqrt{\left( \frac{1}{T} \right)^2 \Delta N^2 + \left( -\frac{N}{T^2} \right)^2 \Delta T^2}. \quad (3.13) \]

Thus, if the total number of transactions was 1046, with approximately 20 percent (0.2 \times 1046 \sim 200) being administrative transactions, and the time for measurement was 200 seconds, give or take a few milliseconds, then:

\[ \Delta S = \sqrt{(1/200)^2 \times 200^2 + (1046/4000)^2 \times 0.001^2}, \]
\[ \simeq \frac{200}{200}, \]
\[ \simeq 1. \quad (3.14) \]

Thus, we quote the value for \( S \) to be

\[ S = \frac{1046}{200} \pm 1 = 523 \pm 1. \quad (3.15) \]

Note that this is an estimate based on a continuum approximation, since \( N \) and \( T \) are both discrete, non-differentiable quantities. As we are only estimating, this is acceptable.

### 3.10 Fourier Analysis and Periodic Behaviour

Many aspects of computer system behaviour have a strong periodic quality, driven by the human perturbations introduced by users’ daily rhythms. Other natural periods follow from the largest
influences on the system from outside. For instance hourly updates, or automated backups. The source might not even be known: for instance, a potential network intruder attempting a stealthy port scan might have programmed a script to test the ports periodically, over a length of time. Analysis of system behaviour can sometimes benefit from knowing these periods. e.g. If one is trying to determine a causal relationship between one part of a system and another, it is sometimes possible to observe the signature of a process which is periodic and thus obtain direct evidence for its effect on another part of the system.

Periods in data are the realm of Fourier analysis. What a Fourier analysis does is to assume that a data set is built up from the superposition of many periodic processes. Any curve can be represented as a sum of sinusoidal-waves with different frequencies and amplitudes. This is the complex Fourier theorem:

\[ f(t) = \int d\omega f(\omega)e^{-i\omega t}, \]

where \( f(\omega) \) is a series of coefficients. For strictly periodic functions, we can represent this as an infinite sum:

\[ f(t) = \sum_{n=0}^{\infty} c_n e^{-2\pi in t/T}, \]

where \( T \) is some time scale over which the function \( f(t) \) is measured. What we are interested in determining is the function \( f(\omega) \), or equivalently the set of coefficients \( c_n \) which represent the function. These tell us how much of which frequencies are present in the signal \( f(t) \), or its spectrum. It is a kind of data prism, or spectral analyzer, like the graphical displays one finds on some music players. In other words, if we feed in a measured sequence of data and Fourier analyze it, the spectral function show the frequency content of the data which we have measured.

The whys and wherefores of Fourier analysis are beyond the scope of this book; there are standard programs and techniques for determining the series of coefficients. What is more important is to appreciate its utility. If we are looking for periodic behaviour in system characteristics, we can use Fourier analysis to find it. If we analyze a signal and find a spectrum such as the one in figure 3.9, then the peaks in the spectrum show the strong periodic content of the signal. To discover these smaller signals, it will be necessary to remove the louder ones (it is difficult to hear a pin drop when a bomb explodes nearby).

### 3.11 Local Averaging Procedures

One of the most important techniques for analyzing data in time series is that of coarse graining, or local averaging. This is a smoothing procedure in which we collect together a sequence of measurements from a short interval of time \( \Delta t \) and replace them by a single average value for that interval. It is a way of smoothing out random fluctuations in data and extracting the trends. It also used as a way of characterizing the pattern of change in a time series.
CHAPTER 3. EXPERIMENT AND OBSERVATION

Figure 3.9: Fourier analysis is like a prism, showing us the separate frequencies of which is signal is composed. The sharp peaks in this figure illustrate how we can identify periodic behaviour which might otherwise be difficult to identify. The two peaks show that the input source conceals two periodic signals. Such signals might not be obvious, but after analysis we can look for them and their cause.

Computer systems and human systems have often quite different patterns of behaviour. When they are combined, the result is often complex and hence local averaging is a straightforward approach to extracting or suppressing detail about the signal.

Let us define a local averaging procedure using fig 3.10. See also appendix B for more details.

The local averaging procedure re-averages data, moving from a detailed view to a less detailed view, by grouping neighbouring data together. In practice one always deals with data which are sampled at discrete time intervals, but the continuous time case is also important for studying the continuum approximation to systems.

**Discrete time data**

Consider the function $q(t)$ shown in fig. 3.10. Let the small ticks on the horizontal axis represent the true sampling of the data, and label these by $i = 0, 1, 2, 3, \ldots, I$. These have unit spacing. Now let the large ticks, which are more coarsely spread out, be labelled by $k = 1, 2, 3, \ldots, K$. These have spacing $\Delta t = m$, where $m$ is some fixed number of the smaller ticks. The relationship between the small and the larger ticks is thus:

$$i = (k - 1)\Delta t = (k - 1)m. \quad (3.16)$$

In other words, there are $\Delta t = m$ small ticks for each large one. To perform a coarse-graining, we replace the function $q(t)$ over the whole $k$th cell with an average value, for each non-overlapping interval $\Delta t$. We define this average by

$$\langle q(k) \rangle_m \equiv \frac{1}{\Delta t} \sum_{i = (k - 1)\Delta t + 1}^{k\Delta t} q(i). \quad (3.17)$$
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Figure 3.10: A coarse-graining, or local averaging procedure involves averaging over intervals larger than the basic resolution of the data. The flat horizontal lines represent the coarse-grained histogrammatic representation of the function. The scaling hypothesis says that if one ‘zooms out’ far enough, and views the fundamental and coarse-grained representations from a sufficiently high level (\( \delta t \gg \Delta t \)), then they are indistinguishable for all calculational purposes.

We have started with an abstract function \( q(t) \), sampled it at discrete intervals, giving \( q(i) \), and then coarse-grained the data into larger contiguous samples \( \langle q(k) \rangle_m \):

\[
q(t) \to q(i) \to \langle q(k) \rangle_m. \quad (3.18)
\]

CONTINUOUS TIME DATA

We can now perform the same procedure using continuous time. This idealization will allow us to make models using continuous functions and functional methods, such as functional integrals. Referring once again to the figure, we define a local averaging procedure by

\[
\langle q(\bar{t}) \rangle_{\Delta t} = \frac{1}{\Delta t} \int_{\bar{t} - \Delta t/2}^{\bar{t} + \Delta t/2} q(\tilde{t}') \, d\tilde{t}'. \quad (3.19)
\]

The coarse-grained variable \( \bar{t} \) is now the more slowly varying one. It is convenient to define the parameterization

\[
\bar{t} \quad \text{and} \quad \bar{t} = \frac{1}{2}(t + t'), \quad (3.20)
\]

on any interval between points \( t \) and \( t' \). The latter is the midpoint of such a cell, and the former is the offset from that origin.
3.12 REMINDER

Although much of the remainder of the book explores mathematical ways of describing and understanding information from human-computer systems, assuming that observations have been made, one should not lose sight of the importance of measurement. Science demands measurement. Mathematics alone only re-describes what we feed into it. Thus, at every stage of investigation into human-computer systems, one should ask: how can I secure an empirical basis for these assertions?

Applications and Further Study 3.

- Developing critical and analytical thinking.
- Formulating and planning experiments to gather evidence about systems.
- Estimating the uncertainties inherent in observational knowledge.
- Diagnostic investigations into anomalous occurrences.
CHAPTER 4

SIMPLE SYSTEMS

This chapter relates an approach to describing systems and their behaviour in terms of their components, using the ideas of predictability and utility.

4.1 THE CONCEPT OF A SYSTEM

The concept of a system is intuitively familiar to us. In our daily lives, we are surrounded by so many systems that we scarcely notice them or think about them, until they go wrong. From the simplest wristwatch whose mechanical parts cooperate to provide a time service, to public transport systems, to the Byzantine convolutions of our taxation systems which serve to distribute resources throughout a larger social collective, systems pervade society at every level.

A modern computer system is a collection of hardware and software components that cooperate to achieve a goal for users. When users employ computers to carry out a task, the users themselves become a part of the system, both working on behalf of the machine when it prompts them, and instructing the machine on the direction it should take next. If users have access to several computers, which cooperate, then the system is said to be distributed in location. A single computer program can itself be regarded as a system; computer programs often consist of multiple functions and procedures which work together in order to evaluate some larger algorithm. Computer systems can be described using various kinds of diagrams and languages that show where information flows from component to component, and how it changes; a whole field of study has built up around this, and we shall draw upon that field here, since it is a formal framework, which admits analysis.

Any ordinary workplace has the elements of a system also. The concept of a system applies whole organizations, or indeed any subset of an organization that can function independently. This might be a company, a branch office, a computer, a network of computers or even single celled
organism on the keyboard of a computer.

The principles of system design and improvement are quite general, and need not be tied to any one of these examples, but it is useful to adopt the language of computer systems (information systems) in what follows (see fig 4.1). This is both our primary area of interest and a more rigorous language than the corresponding terms of the social sciences, and it ties the discussion immediately to one of its most important applications.

![Figure 4.1: An informal diagram of associations within a system, which shows the main aspects for consideration.](image)

4.2 DATA-STRUCTURES AND PROCESSES

There are many ways to classify systems. At the most basic level one may speak of two kinds of system: those which are dynamic and those which are static.

A static system is often referred to as a data-structure, rather than a system, i.e. it is a systematic organization of its parts or resources (a form of data), which does not change. An archive is an example of this, as is a building, as is the book that you are reading.

**Definition 7 (Data structure).** A data structure is an ordered (systematic) collection of resources or records that are interrelated.

The data form a collection of key-value pairs that also form a ‘graph’ (see chapter 6). The semantics of the data in a functional system are usually implied by the structure.

**Example 14.** A building is a static data structure: it is a regular arrangement of parts which contribute towards a function. A library, or archive, is another example, in which individual records
are organized in an orderly pattern, with a scheme for relocating the information. The book you are reading is a third example. The functional components in these examples are somewhat diverse in their variety, but the all examples share a common feature: they are organized patterns designed to serve a purpose.

Example 15. Many software systems are passed data in a fixed format like JSON or YAML to represent key-value pairs. Databases contain ordered structures, with a fixed structure, to make the retrieval location easy to compute.

For a system to exceed the archival character of a museum, there has to be some activity. A dynamical system is a system which evolves in time with a rate of change; it usually produces something and experiences a number of different operational states, such as running, stopped, broken, waiting, etc. Dynamical systems are a more interesting class of systems, because they open up literally a whole new dimension (time) for organization. Human-computer communities belong to this group.

In order to describe the activity within a system, it is useful to define the notion of a process. A system may comprise several independent processes.

**Definition 8 (Process).** A process is a unit of activity within a system. In order for something to happen, there needs to be something with a freedom to change. A process therefore comprises a set of resources, together with an algorithm (a sequence of operations) for manipulating them. A process is unleashed by the freedoms of the system and restricted by its constraints.

A process can operate on a data structure and alter it. The sum of a process and a data structure is thus a dynamical system.

**Definition 9 (Dynamical system).** A dynamical system is a set of processes which act on a data structure.

A computer program is a process in which an algorithm changes the data in the computer. A maintenance procedure, such as cleaning a building, is a process which changes the state of organization of the building’s resources.

**Example 16.** Active processes: a process is a combination of resources associated with a executed task. It includes a code text, which contains instructions and algorithms for data-manipulation, and it comprises data and resources associated with the task. In a multiprocess environment, each process is an independent object, with its own progress documentation (stack and index markers in computers), so that, if the process should be interrupted, its current state can be saved and later resumed, without loss of integrity.
Example 17. Passive data: the purpose of a process is to manipulate some data. Such data are often arranged in some non-trivial structure, which makes them easily accessible to the algorithms of the process. A filesystem is an example of passive data, as is a database.

4.3 **REPRESENTATION OF VARIABLES**

To describe a system in definite terms, one needs to identify properties of its resources that can change or vary. Without any such variation, a system would be truly uninteresting. There are two kinds of information in a system:

<table>
<thead>
<tr>
<th><strong>Definition 10</strong> (Resource information). The information that is used and produced by the system as part of its functioning. This is the fuel and produce of the system. This information might be sent back and forth between different functional elements in the system, between clients and external service providers, or even between the system and its storage. Resource information may be both dynamical (quantitative) and semantic (qualitative) in nature.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Example 18.</strong> Configuration files, which parameterize the way software behaves is part of the static resource information of control in machinery. Runtime data are the dynamical resources that guide the operation of the system. This includes input and output between human, machine and software.</td>
</tr>
<tr>
<td><strong>Example 19.</strong> Organizational records, customer and sales records, internal communications, personnel movements, etc, all contribute to resource information in the human aspects of a system.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Definition 11</strong> (Algorithmic information). The information on how to achieve the task, or generate the produce of the system is contained in a detailed programme of steps and procedures. This includes control information on which way to branch as as a result of a question, and information about the initial state of the system. Algorithmic information is semantic information.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Example 20.</strong> The source code of computer programs represents the documentation of algorithmic information for computing machines.</td>
</tr>
<tr>
<td><strong>Example 21.</strong> Recipe books and procedural handbooks represent the documentation of algorithms for the human actors in information systems.</td>
</tr>
</tbody>
</table>

Describing the actual characters of a system’s resources requires abstraction. One must use functions which vary with the basic parameters of the system (e.g. time) to represent those properties. The basic properties of a dynamical system are usually labelled $q_i$, for $i = 1, 2, 3, \ldots$. A function is a mapping from the parameter space into a range of values. The range is the set of possible values that the variable can change into.

What might $q(t)$ represent?
1. A number of objects
2. The value of an object
3. An average value
4. The size of an object
5. The shape of an object?

There are no rules about what can and cannot be represented by a variable. The challenge is only to find a sensible and sufficient representation of a phenomenon, to some appropriate level of approximation.

Example 22. In a University or College, students and lecturers follow a system in which they are allocated rooms to be in, at different times. We could choose to analyze the behaviour of this system in a number of ways. One could, for instance, measure the number of students in each lecture room, as a function of time. One could measure the average number of students to get an idea of attendance. One can characterize the rate of work, or how fast the contents of the blackboard change. One could attempt to measure the rate of learning (how would we measure this?). One could measure room temperature, or air-quality and relate it to student attentiveness, and rate of learning. In each case, one must decide what it is one is interested in finding out, and characterize that mathematically in terms of measurable quantities.

The lesson one learn from natural sciences is that models of systems are suitably idealized representations, not exact and unique facsimiles of reality.

4.4 THE SIMPLEST DYNAMICAL SYSTEMS

Complex systems can often be built up by combining simpler ones. An understanding of the behaviour of simple systems is therefore a reasonable place to begin, in order to fathom the greater complexities of more realistic systems. There is something to be learned from even the simplest mechanical device.

The property which characterizes dynamical systems is that their resources change in time; but how? What kind of change can be expected? The theory of change is the study of differential and difference equations. These predict three basic kinds of change:

- Monotonic change (growth, decay)
- An oscillatory change
- Chaotic change
These behaviours can be found in well known places.

Example 23. A simple pendulum is a system which provides a time service. It can go wrong if one of its components fails. The pendulum is a collaboration between a force, a pivot a string and a mass. A pendulum has an oscillatory motion which gradually dies away. It is a combination of oscillation and decay.

It is important to capture the essence of systems in a way that is conducive to analysis. For that reason, we begin with the simplest of mathematical considerations. A mathematical definition of a system is the following:

- A set of variables \( \{ q_i(t) \} \), where \( i = 0, 1, 2, \ldots \) (information describing resources) whose values can change in time.

- A set of rules \( \{ \chi_i \} \) which describes how the variables change with time.

- A definition of the rates \( \{ Dq_i / Dt \} \) at which the difference variables change in time, for deterministic systems, or a probable rate with which transitions occur non-deterministically.

These three things are the basic requirements which are necessary and sufficient to comprise any system that changes in time. This extends from simplest of mechanical devices to the most complex chaotic combinations of elements. To make predictions about a system, one also needs to know what state it was in at some known or initial time. Such information is known as a boundary condition.

Example 24. In a computer system one has variables that characterize state; e.g. the amount of data on a disk, the rate of processing, the number of users, etc. A set of physical rules governs the hardware at the level of electronics, software provides a set of rules for program execution, and policy provides an even higher-level set of rules about how the hardware and software should be used. The system is not static, so we find a rate of change of data on a disk, a rate of change of number of users, and even a rate of change of ‘rate of processing’, as jobs are turned on and off.

Any larger definition of a system which we concoct must contain the basic definition above. In system administration, we are concerned with far more complex systems than can be described with the aid of a few simple variables. One is forced to deal with approximation, as one does in the natural sciences. This is an unusual remedy for computer science, which is more at home with logical propositions and exact theorems. Nonetheless, the lesson of the natural sciences is that complexity abhors precision, and forces one to embrace approximation as a tool for making headway.
4.5 More complex systems

In general we need to describe a complete organization, with interconnecting data structures, and inter-communicating sub-processes. Note that the word organization is ambiguous in this context, in that it describes both an attribute of a system (how well it is organized) and a name for a system (a company or other institution). We shall henceforth limit the word organization to the first of these meanings, and refer to the second by the term enterprise. Adding these notions, we have:

**Definition 12 (System).** A system is a set of resources (variables and processes) and consumers, together with descriptions of how those resources are organized and how they develop in time. This total description defines the arena in which the system develops. It prescribes the possible freedoms one has for change, the constraints imposed externally and internally, for activity within the system.

At a superficial level, we can identify the key elements in common systems.

**Example 25.** A public transport system has a set of resources (busses and trains) \( q(t) \) which are constrained to move on roads or rails, and which run on expended fuel. The details of change in the system are partly determined by policy (the schedule), partly by environmental considerations and critically on the natural laws which govern the physical processes driving the system (all these are in \( \chi \)). The rate of flow of transport \( \frac{Dq}{Dt} \) is related to the overall change in the system.

**Example 26.** A web server is like a query handling system or help desk. These are all systems in which the number of incoming, unanswered requests \( q(t) \) is changing with time. A protocol for handling the requests constrains their expedition by various algorithms that manipulate resources. The rate at which the system expedites requests is \( \frac{Dq}{Dt} \).

4.6 Freedoms and constraints

Any system which has some kind of predictable or regular behaviour is a balance between the freedom to develop and change and a number of constraints which limit the possibilities for change to predictable avenues.

**Definition 13 (Degree of freedom I).** A degree of freedom is a potential for change within a system. Freedom to change is usually represented by a parameter which can be varied over a range of values.

For example, in a service based system, the freedom to accept new clients permits the system to expand.
Table 4.1: Freedoms and constraints in organizations.

<table>
<thead>
<tr>
<th>Freedoms</th>
<th>Constraints</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expansion</td>
<td>Available budget</td>
</tr>
<tr>
<td>Time</td>
<td>Deadline, or limited duration</td>
</tr>
<tr>
<td>Space</td>
<td>Specific location, limited space</td>
</tr>
<tr>
<td>Rate of service</td>
<td>Throughput of bottleneck</td>
</tr>
</tbody>
</table>

Example 27. In a computer system, the freedoms include: memory space, processing time, physical location (server host for job placement), scheduling time, etc. Jobs also have a freedom to be expressed in any number of languages.

Example 28. A program might run equally well on an array of one or twenty computers. The number of computers is thus a freedom. Similarly, a system might have the freedom to increase its use of memory; that is then a freedom of that system.

Example 29. In a computer system, the constraints include the fact that memory is finite, and jobs have a limited quota, CPU processing is finite and has a maximum rate. Jobs have to be compiled into the machine language of the physical processor.

Example 30. In the client-service system above, a constraint could take the form of limited resources for handling client queries (a maximum number that can be expedited per unit time); similarly, it could represent a policy constraint that denies the service to certain clients, or limits their availability to the service. Another constraint is a productivity goal.

At the machine level, it might be that a program only runs equally well on computers with a particular operating system; in that case the choice of operating system becomes a constraint. Similarly, the amount of memory available to a program is normally limited by the total amount on that computer; the amount of memory is thus a constraint on program execution.

Note, the words freedom and constraint are used in a strict sense. Their usage do not imply what advantages or disadvantages they confer on a system. Do not be tempted to bestow these terms with social connotations; e.g., a smart human might creatively use a constraint to his or her advantage, but that is not the same as it being a freedom. For example, a geographical constraint in which an organization is limited to one building might be turned into a positive attribute by arguing that this lends cohesion to the organization. This does not mean that the constraint is really...
a freedom. The advantage here is only made possible by virtue of another freedom, namely the freedom to be creative in that context.

As well as objects which are manipulated and changed, most systems have input and output channels, where information is exchanged with external actors — the “environment”.

4.7 Symmetries

Symmetries are descriptions of what can be changed in a system without affecting the system’s function. Determining what is not important to the functioning of a system is a way of identifying degrees of freedom that could be manipulated for strategic advantage. Knowing about these freedoms might be an advantage to someone managing the system. A change which does not affect a sub-system might nevertheless result in an advantage elsewhere.

**Definition 15 (Symmetry or invariant).** When a change of any variable quantity, whether dynamical or semantic, within a system leaves the outcome of the system invariant, we say that the system and the variable possesses a symmetry or invariant property.

If, for instance, we come up with a system model in which results depend on a specific choice, where in fact that choice does not matter, then we know that model must be wrong.

**Example 31.** If one moves every user of a desktop workstation to a different workstation, the organization will still function in exactly the same way, if the workstations are all alike. This freedom to reseat people might allow groups of workers to sit in close proximity, for verbal communication, or it might allow workers to be spread out to balance the load.

**Example 32.** In a factory or office, workers who are trained to carry out the same jobs can rotate their positions and tasks in shifts, and repostings to other factories or offices. The symmetry between workers means that the outcome will be invariant.

**Example 33.** In so-called cloud computing, or commoditized Infrastructure as a Service (IaaS), software may be moved from host to host by the use of virtual machines, and software process containers. This is called process or machine virtualization. The freedom to move a job from location to location is a symmetry. Not all jobs may have this property, because they are pinned to a location by the need for particular hardware or software dependencies. Dependencies are thus the enemy of symmetry.

**Example 34.** If one swaps all of the busses on route 20 with those on route 37, the service will not be affected, provided the busses are comparable. So buses can be rotated and checked for maintenance in parallel to those in service.
Example 35. If one relabels every file on the system, in all references, it will continue to work as before. Indeed, a file is an invariant abstraction for a changing collection of allocated storage blocks filled with changing data.

4.8 ALGORITHMS, PROTOCOLS AND STANDARD ‘METHODS’

Systems embody a cooperation of parts and are often surrounded by an environment of unpredictable occurrences. To address the cooperation between parts of a system needs to be formalized by defining the mechanisms that contribute to it; to cope with unpredictable, external events, mechanisms and procedures need to be introduced that offer predictability. These requirements are covered by the concepts of algorithm and protocol.

**Definition 16 (Algorithm).** An algorithm is a recipe, or sequence of steps and decisions that solve a particular problem within a system. Algorithms are sometimes referred to as methods in the parlance of programming. A formal definition of an algorithm can be provided in terms of Turing machines (see [LP97]).

An algorithm is a reasoned flow of logic, designed to efficiently perform an operation or sequence of operations. An algorithm is what one finds if one opens up the black-box of an operator and peers at its inner workings. Algorithms are not arbitrary, though there might be several algorithms which solve the same problem. In that situation, a policy decision is required to determine which algorithm is to be used.

**Definition 17 (Protocol).** A protocol is a standard of behaviour, or a strict rule of conduct which ensures that one part of a system behaves in a predictable and comprehensible way. Protocols ensure that one part of a system is able to cooperate with another, and that the integrity of the process is maintained, i.e. information is not lost or misunderstood. A protocol is formally a constraint on a process.

Protocols are used to ensure consistency and to avoid error when executing a process. Typical examples of protocols are used when two communicating parties must understand one another.

Example 36. When a computer sends data over a network, it does so by coding the data into a stream of bit pulses. The machine receiving the message would have no idea how to decode the message unless a pre-agreed standard of behaviour were established in advance: e.g. the first 8 bits are a number, the next 24 bits are three 8-bit characters, and so on. Without such a protocol for interpreting the stream of bits, the meaning of each bit would be lost.
Another example of a protocol is a response plan in case of emergency, e.g. fire, security breach, war etc. The purpose of a strict code of behaviour here is to minimize the damage caused by the event, as well as to mobilize counter-measures. This ensures that all parts of the system are activated and informed in the right order, thus avoiding confusion and conflict.

Protocols are a part of system policy. They are arbitrary, even though some properties of protocols can be analyzed for efficiency and efficacy. They are strategic choices.

Example 37. **Computer security intrusion**: i) freeze the state of the system, ii) gather standardized information and evidence (Copy the current memory to a file), iii) Pull the plug to avoid setting off any logic traps, iv) Report the intrusion to law enforcement. v) Reboot system with a CD-ROM or trusted read-only medium, to avoid logic traps set by an attacker.

The goals of this protocol are: protect the system, secure evidence, repair the problem, and obey the law in that order. If a different set of priorities were in force, the protocol might be changed. It is designed by a person with experience of intrusion, to take into account all of the interests above. An inexperienced person might not understand the protocol, but by following it, the goals of the system will be upheld. The technology of computer security is to a large extent a litany of protocols, designed to avoid unwanted behaviour.

The difference between an algorithm and a protocol is subtle. An algorithm is a sequence of instructions or steps; a protocol is only a specification of what kind of steps is allowed. One is a process specification, the other is a constraint on a specification. Protocols do not make decisions, they are pre-agreed standards.

### 4.9 CURRENCIES AND VALUE SYSTEMS

While autonomous, mechanical and electronic systems can be described purely by simple physical principles, once humans are involved in a system, human values necessarily become part of the equation. This complicates a system, and many engineers find this interaction disturbing because society has an ingrained culture of treating human values as fundamentally different to physical measurements.

Some might even say that the idea of modelling human values, in the same way that one models physical processes, would be disrespectful. As we shall see, however, there is no basic impediment to writing formal rules for human values and concerns; these merely extend the complexity of systems by introducing additional constraints and boundary conditions. Indeed, to represent financial and economic aspects of a system, one already does precisely this.

Human emotion allows us to attach importance, and hence value, to almost anything. Not all values can necessarily be traded, as money or goods can. Happiness, for example, cannot normally be traded, say, for food, but it might be reasonable to say that happiness of a workforce could be
traded for efficiency, in a system with a Draconian work ethic. The key to analysing the interactions between human values and physical resources is to assign to them arbitrary scales which can then be calibrated so that rules can be written down.

4.9.1 **ENERGY AND POWER**

Nature’s fundamental system book-keeping currency is called *energy* in the physical sciences. Energy is simply an abstract representation of the level of activity in different parts of a system. It has an arbitrary value, which is calibrated against the real world by convention, i.e. by using some system of units for measurements, and by adjusting certain constants in physical laws to ‘make it right’.

When parts of a system interact, they are said to exchange energy. Each new interaction has its own equation, and requires a ‘coupling constant’ which calibrates the effect of the arbitrarily dimensioned energy transfer from one part, to the measured effect in the other part. Whatever new age writers might believe, there is really only one kind of energy, just as there is really only one kind of money. People have different conventions for referring to energy or money in different places, but the idea is the same.

Just as saved money can be traded for services, stored (potential) energy can be traded for activity. Physicists have long been dealing with this simple book-keeping concept without questioning its validity, so it should not be a surprise that the same idea can be applied to any form of currency.

**Example 38.** Basic energy requirements are at the heart of all systems, grounded in the physical world. Machines require power and humans require nutrition. A failure to provide these, leads to a failure in the system.

4.9.2 **MONEY**

In days of old, one used silver, gold and other riches to trade for goods. Sufficient gold is still kept in reserves around the world to be able to trade paper currency for its value in gold if the bearer demands it. Today, however, we use money to represent only the promise of something real. Our abstraction of wealth has reached such extremes that we buy and sell shares in the estimated value that a company might have, if its assets were sold. The value of something, in our modern world, is clearly not an intrinsic physical property, like its electric charge, or its mass. It is a fictitious quantity based as much on trends and feelings as it is on physical resources.

In short, money is worth what everyone believes it is worth; no more and no less. It is a sobering thought therefore that our contemporary society and all of its systems run on money. Money is used to measure the cost of building a system, the cost of running it, and the value of what it produces. We speak of *assets* or *resources* to mean the things of value which are recognized by a potential buyer.
How should money be represented in formal (mathematical) models of systems? How does it relate to other measurable resources, like time, space, equipment and so on? Clearly, ‘time is money’ because humans will not work for nothing: we require the promise of reward (money). Space and commodities ‘are money’ because we value anything which is not in infinite supply. But, if the value of these things depends on the fickle moods of the human actors in the system, how can it be used to represent these other things?

To say that a relationship exists between time $t$ and money $m$, for instance, is to say that

$$t = f(m),$$

for some function $f$. In the simplest case, this would be a simple linear relationship:

$$t = km + c,$$

where $k$ and $c$ are arbitrary constants, which can be fixed by calibration. They represent what it costs to buy someone’s time.

The value of objects is arbitrary (a matter of policy), so any relationship could change from system to system, or from time to time within a system. It is not always necessary to think in terms of money. If the issue is that of a cost to the system, then money takes the form of a constraint or limitation.

4.9.3 Social Currency and the Notion of Responsibility

Money is not the only form of abstract currency. Humans value other things too: peer status, privilege and responsibility are all used as measures of social value, every bit as real as money.

The meaning of responsibility is taken for granted, in common speech, but it has several different meanings in the running of systems. If we are to analyse the concept formally, it is necessary to relate responsibility to more tangible concepts.

- **Cause and effect**: For a machine, one says that a component is responsible for an action if the action depends on it somehow. Responsibility is thus associated with an implied trust or dependency. It is simply the law of cause and effect, which is central to the analyses in this text. e.g. the printer is responsible for writing information to paper; electricity is responsible for making a lamp work.

For humans, responsibility has two meanings, and these are somewhat subtle:

- **Responsibility for completing a task (policy)**: This refers to the assignment of a task to some agent within the system (a person or a department, for instance). It is often laced with connotations of a penalty for failure, or a reward for success. e.g. “you are responsible for getting this job done before March!” This is a combination of an arbitrary policy decision.
and a constraint. The correct way to model this is thus as an externally controlled condition, together with the relevant constraint on the system resources (in this case: time). Rewards and penalties can be modelled by introducing a fictitious currency (see below).

- **Responsibility for actions (blame):** As in the case above, this is used to imply the direction of a reward or a punishment associated with the completion, or failure of something in the system. However, this kind of responsibility can be assumed, or transferred from one object to another. A manager or commanding officer will often be made (or held) responsible for a whole department or unit, for instance. This transfer of responsibility is not necessarily related to cause and effect, since the manager’s actions do not necessarily have any direct influence on what transpires. It covers the situation where a leader trusts in the outcome of a hidden process. e.g. an accountant trusts computer software to calculate a correct answer, and holds the software producer responsible for the result, but his boss holds him responsible. This can be modelled as a combination of policy with a fictitious currency scheme.

In modelling human values, one deals with issues such as status, prestige and other emotional considerations. These are social currencies, analogous to material wealth, or resource riches. Social status amongst humans can affect the details of how policy applies to them, and can act as an incentive or a deterrent to certain kinds of behaviour.

**Example 39.** A person with sufficient privilege might be given access to parts of the system which are unavailable to others. Emotional reward can be a sufficient motivation to complete a task, or conversely emotional pressure can be a hindrance or even a factor motivating sabotage. Disgruntled employees have been responsible for the theft of millions of dollars through their abuse of human-computer systems, particularly in the financial sector.

One begins to appreciate the complexity of human-computer communities when one attempts to represent the exchanges that take place on the human side of systems. For computer administrators, this type of modelling is normally only done in connection with the security of the system, i.e. in threat analysis. Software engineers who design software for critical systems need to think about such issues that might lead to human error. Businesses and people-run enterprises such as universities and the military depend critically on the actions of humans; thus social currency tends to dominate these systems.

### 4.10 Open and Closed Systems: The Environment

If we wish to describe the behaviour of a system from an analytical viewpoint, we need to be able to write down a number of variables which capture its behaviour. Ideally, this characterization
would be numerical since quantitative descriptions are more reliable than qualitative ones, though this might not always be feasible. In order to properly characterize a system, we need a theoretical understanding of the system or sub-system which we intend to describe. This is a few important points to be clear about.

Dynamical systems fall into two categories, depending on how one is able to analyze them. These are called open systems (partial systems) and closed systems (complete, independent systems).

- **Open system:** This is a sub-system of some greater whole. An open system can be thought of as a black box which takes in input and generates output, i.e. it communicates with its environment. The names source and sink are traditionally used for the input and output routes. What happens in the black box depends on the state of the environment around it. The system is open because input changes the state of the system’s internal variables and output changes the state of the environment. Every piece of computer software is an open system. Even an isolated total computer system is an open system as long as any user is using it. If we wish to describe what happens inside the black box, then the source and the sink must be modelled by two variables which represent the essential behaviour of the environment. Since one cannot normally predict the exact behaviour of what goes on outside of a black box (it might itself depend on many complicated variables), any study of an open system tends to be incomplete. The source and sink are essentially unknown quantities. Normally one would choose to analyze such a system by choosing some special input and consider a number of special cases. An open system is internally deterministic, meaning that it follows strict rules and algorithms, but its behaviour is not necessarily determined, since the environment is an unknown.

- **Closed system:** This is a system which is complete, in the sense of being isolated from its environment. A closed system receives no input and normally produces no output. Computer systems can only be approximately closed for short periods of time. The essential point is that a closed system is neither affected by, nor affects its environment. In thermodynamics, a closed system always tends to a steady state. Over short periods, under controlled conditions, this might be a useful concept in analyzing computer sub-systems, but only as an idealization. In order to speak of a closed system, we have to know the behaviour of all the variables which characterize the system. A closed system is said to be completely determined.\(^1\)

Suppose we want to consider the behaviour of a small sub-system within the entirety of a much larger system (e.g. a computer on the Internet, or an animal in a complex ecology); first, we have to define what we mean by the sub-system we are studying. This might be a straightforward

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\(^1\)This does not mean that it is exactly calculable. Non-linear, chaotic systems are deterministic but inevitably inexact over any length of time.
conceptual partitioning of the total system, but conceptual decompositions do not necessarily preserve causal relationships (see fig 4.2).

In fact we might have to make special allowances for the fact that the sub-system might not be completely described by a closed set of variables. By treating a sub-system as though it were operating in isolation, we might be ignoring important links in the causal web. If we ignore some of the causal influences to the sub-system, its behaviour will seem confusing and unpredictable.

The principle of causality tells us that unpredictable behaviour means that we have an incomplete description of the sub-system. An important difference between an open system and a closed system is that an open system is not always in a steady state. New input changes the system. The internal variables in the open system are altered by external perturbations from the source, and the sum state of all the internal variables (which can be called the system’s macrostate) reflect the history of changes which have occurred from outside. For example, suppose we are analyzing a word processor. This is clearly an open system: it receives input and its output is simply a window on its data to the user. The buffer containing the text, reflects the history of all that was inputted by the user and the output causes the user to think and change the input again. If we were to characterize the behaviour of a word processor, we would describe it by its internal variables: the text buffer, any special control modes or switches etc.

4.11 RELIABLE AND UNRELIABLE SYSTEMS

**Definition 18** (Unreliable system). An unreliable system is one which attempts to carry out its function without verification or guarantee of success.

Unreliable systems are used either where a trust is placed on the mechanisms of the system to perform their duty, or where a failure is unimportant. For instance, if an unreliable system repeats its actions fairly often, a failure might be corrected at a later time with a high probability. In some cases, it does not matter whether a process fails; we might use an unreliable system to approximately probe or test a random variable, without too much ado.

**Definition 19** (Reliable system). A reliable system is one which tries, verifies and repeats its actions until the result succeeds.

Reliable systems can prove to be expensive, since resources have to be applied to monitor and correct any errors or failures that occur.

**Example 40.** The Internet Protocol (IP) has two control layers: the User Datagram Protocol (UDP) and the Transmission Control Protocol (TCP) that are unreliable and reliable respectively. UDP is used for ‘one shot’ requests, such as name service look ups and route tracing probes,
Figure 4.2: A complex system is a causal web or network of intercommunicating parts. It is only possible to truly isolate a subsystem if we can remove a piece of the network from the rest without cutting a connection. If we think of the total system as $S(x_1 \ldots x_n)$, and the individual subsystems as $s_1(x_1 \ldots x_p)$, $s_2(x_p \ldots x_n)$ etc, then one can analyze a subsystem as an open system if the subsystems share any variables, or as a closed system if there are no shared variables.
where a reply is not necessarily expected. TCP is used for more formal behaviour of network communication protocols where certainty is demanded.

**Applications and Further Study 4.**

- Understanding fundamental issues in system analysis.
- Examining and classifying systems using the concepts described here.
- Isolating the basic behavioural traits of a system.
- Identifying the freedoms and constraints to better understand how a system might be modified or improved.
Chapter 5

Sets, States and Logic

The concept of a state or condition of a system will be central to several discussions. In order to discuss states, we need to have variable quantities that can take on a set of values. The state of any object is its value at a given place and time. Usually variables cannot take on just any value; they take values from a specified set of values.

We need a language that is general enough to be applied to a wide range of situations, but which is specific enough to make clear and verifiable statements about systems. The language of sets and mathematics allows us to state things precisely, and will be of great utility in describing Networks and System Administration.

5.1 Sets

A set is the mathematical notion of a collection of related things. The idea is general and is not limited simply to numbers. Sets describe all manner of useful collections. We must take care not to confuse the term ‘group’ with ‘set’, as these words both have special meanings in mathematics. Sets are denoted by lists enclosed by curly braces \( S = \{ \ldots \} \).

**Example 41.** Example sets include: days of the week:

\[
D = \{ \text{Sunday}, \text{Monday}, \text{Tuesday}, \text{Wednesday}, \text{Thursday}, \text{Friday}, \text{Saturday} \} \quad (5.1)
\]

Types of operating system:

\[
O = \{ \text{Windows}, \text{Solaris}, \text{GNU/Linux}, \text{MacOS} \} \quad (5.2)
\]

Files owned by user mark:

\[
F = \{ \text{INBOX, file.txt, ...} \} \quad (5.3)
\]
Directories owned by mark:
\[ D' = \\{\text{Mail, Documents, \ldots}\} \] (5.4)

System load average values measured in an experiment:
\[ L = \{0.34, 0.42, 0.45, \ldots\} \] (5.5)

Example 42. A set of rules is often called a policy.

The union of two sets is the combination of all elements from both sets, e.g. the catalogue set is the union of directories \( D' \) and files \( F \):
\[ C = F \cup D'. \] (5.6)

The union does not contain more than one copy of an element, so \( \{A, B\} \cup \{A\} = \{A, B\} \). Similarly, files \( F \) are a subset of the catalogue \( C \):
\[ F \subset C. \] (5.7)

If we let \( S \) be the set of all secure policies, and \( P \) be the set of policies in use by an organization, then
\[ P_S = S \cap P \] (5.8)

is the intersection or overlap between these sets (see fig 5.1) and represents the sets that are both secure and in use. If \( S \) is a subset of a set \( X \), then the rest of \( X \) that is not in \( S \) is called the complement and is written \( X - S \) or simply \( -S \) (also written \( \neg S \)).

Example 43. Some common sets include.

- \( \emptyset \) is the empty set, containing no elements.
- \( R^1 \) The set of real numbers (often written simply \( \mathbb{R} \) or with a calligraphic \( R \)).
- \( \mathbb{Z} \) the set of integers.
- \( R^n = R^1 \times R^1 \ldots R^1 \) is the \( n \)-dimensional Cartesian space of real numbers, e.g. \( R^3 \) in three dimensional Euclidean space.

Note that the notation \( \neg \) is used interchangably with \( \text{NOT} \) and the complement operator “\( - \)” for sets.
5.2 A SYSTEM AS A SET OF SETS

To describe a system, we must have a number of objects that can be related to one another and change in some way. Sets enter this picture in two ways:

- There is a set of objects that comprises the components of the system.
- Each component object can change its value or properties, by taking a new value in the set of values that it can possibly take.

Thus, our abstract picture of a system is a set of variable objects, each on which can take on a value from another set of values. These objects are can be related to one another, and the values they take can change according to rules that are determined by the system. The objects are typically areas of memory in a computer system, specific people in a team, or even sets of users who work together. In chapter 6, we shall see that this picture also describes a graph.

Example 44. Strings of symbols or operations (or symbols that represent operations) can be used to formulate some systems. If one can express a system in these terms, the development of the system becomes akin to to problem of transmitting information over a communications channel. This is one of the themes of this book.

Sets of sets the the main way we can talk about scaling of discrete systems.
Example 45. A computer system may be composed of sets of software, running on sets of computers, housed in sets of racks, housed in sets of locations, across sets of countries, etc. Sets of users use sets of computers that may not be co-located but distributed for complicated reasons.

Sets allow us to describe things in a bounded way, without necessarily being localized in a single region. Sets are the most basic and powerful description we have, on which everything else can be based.

5.3 Addresses, Dependencies, and Mappings

Since we need to refer to the elements in a set, we give each element in a system a name or address, i.e. a label that uniquely identifies it. There is thus a mapping from names and addresses to objects (see figs. 5.3 and 5.4). This mapping may form a hierarchy of components connected together as dependencies.

Example 46. In technology, systems are often componentized or modular in construction. That means we can separate parts from one another and understand them partially in isolation. Dependencies are components that are used by larger components. A television and a computer depend on transistors, resistors, capacitors, etc. Software is often packaged as components too, and these packages usually depend on other packages. Certain components are so basic to the operation of software that nearly all other software depends on them. The operating system falls into this category.

Example 47 (Functional mappings and dependencies). In fig. 5.2, we see a mapping from a set of five elements to another set of three elements. In this case, the mapping does not exist for all elements. A similar illustration could be used to create a mapping from the set of computer architectures to the set of operating systems that can run on them: e.g.

\[
O = \{\text{Windows, Solaris, GNU/Linux, MacOS}\} \\
A = \{\text{Sparc, Intel, ARM}\}
\]

(5.9)

So that, if we create a mapping function \(f\), one may write:

\[
o = f(a)
\]

(5.10)

where \(o \in O\) and \(a \in A\), e.g. \(\text{Windows} = f(\text{Intel})\).

The advantage of writing the function is that it can be written for any variable that takes values in the respective sets. This allows us to solve equations for valid mappings, e.g. what are the solutions of

\[
\text{Windows} = f(x)?
\]

(5.11)
Depending on whether the map is *single-valued* or *multi-valued*, there might be zero, one or more solutions.

![Function Diagram](image)

Figure 5.2: A function \( M \) is a mapping from one set, called its domain, to another set called its range. The domain and range of the function do not have to be sets of the same size.

![Time Function Diagram](image)

Figure 5.3: A function of time, \( q(t) \), is a mapping from an arbitrary set of times into the set of real numbers \( \mathbb{R}^1 \).

If we have a continuous set of values, i.e. a set that takes on a different value for any real number parameter, the function can be described in terms of known differentiable functions. See the example in fig. 5.3.

### 5.4 Chains and States

Using the idea of sets, we can define states to be the current values of the objects in the system:

**Definition 20 (State).** A state of the system (also called a microstate) is a value \( q \in Q \) associated with a single addressable entity or location within the system.
A chain is a sequence of events $X_n (n = 0, 1, \ldots, N)$, where each event $X_n$ takes a value $q_i$ ($i = 1, \ldots, I$) called a state at address $n$, belonging to a set of values $Q$. The integers $n$ normally label the development of the process in discrete steps, and are interpreted as time intervals, but they could also be spatial locations, e.g., a sequence of values on a hard disk. The transition matrix $T_{ij}$ is written in a variety of notations in the literature, including the following:

$$T_{ji} = p_{ji} = |\langle q_j | q_i \rangle|^2 = P(X_{n+1} = q_j | X_n = q_i).$$

It represents the probability that the next event $X_{n+1}$ will be in a state $q_j$, given that it is currently in the state $q_i$. By discussing the probability for transitions, we leave open the issue of whether such transitions are deterministic or stochastic. There is a number of possibilities. If $T_{ij} = 1$ at $X_n$, for some $i, j$, the process is deterministic and one may write the development of the chain as a rule

$$X_{n+1} = U_t(X_n, \ldots, X_0),$$

in the general case. If $T_{ij} < 1$ at $X_n$, for all $i, j, n$, the process is stochastic. If $T$ depends only on the current state of the system,

$$P(X_{n+1} = q_i | X_0, X_1, \ldots, X_n) = P(X_{n+1} = q_i | X_n), \quad \forall \ n \geq 1,$$

then the chain is said to be a Markov chain, or memoryless. Markov processes are also called steady state, or equilibrium processes. A system that behaves like a Markov chain has maximum uncertainty, and satisfies the second law of thermodynamics, meaning that ‘entropy’ increases. If $T$ depends on the whole history of $\{X\}$, then it is called a non-equilibrium, non-Markov process.
A state is called persistent if
\[ P(X_{n+m} = q_i | X_m = q_i) = 1, \text{ for some } n \geq 1 \]  
(5.15)
and transient if
\[ P(X_{n+m} = q_i | X_m = q_i) < 1. \]  
(5.16)

The terms periodic, aperiodic and ergodic also describe chains in which the processes return to the same state. Readers are referred to [GS01] for more about this.

Example 48. Computer science, economics, and many other disciplines build on models that are deterministic by default. We often assume that machines and even human systems have simple causal behaviours, but in fact there is a lot of uncertainty or randomness at work. In non-deterministic systems, we can only speak of probabilities of outcomes. In Promise Theory (see volume 2), we start from the assumption that promises may not always be kept, no matter how hard we try, or how well a system has been designed. Quantifying this uncertainty is a job for rational methods. Probabilistic formulations are thus very important.

5.5 Configurations and Macrostates

If we want to collectively talk about the state of all of the objects or locations in a system, at a given time, we use the term configuration. In large systems, a precise system configuration consists of too many independent values to describe in detail and we move to a statistical, averaged description.

Example 49. One of the central issues in system administration is the management of device and software configurations.

**Definition 21** (Configuration). A configuration of the system is the vector of values (microstates) of all its resource variables \( q_i(t,x,\ldots) \), at a given time \( t \), for all positions \( x \) and other parameters so on.

**Definition 22** (Macrostate). A macrostate is an averaged, collective description of a configuration that captures its statistical behaviour in a simplified, high level description. Any function of all the microstates that leads to a summarized value can be called the macrostate of the system. It can therefore be represented as a set of probability distributions for the probabilities that the \( i \)th resource variable is in a state \( s_i \): \( P_i(x = s_i) \).

The concepts of microstates and macrostates are not necessarily unique; they depend on a particular viewpoint of a system, and thus they can be defined, in each case, to discuss a particular issue in a particular way. The main advantage of a description in terms of states is the clarity and definiteness that such a description brings to a discussion.
Example 50. To discuss the security configuration of a computer, it is not necessary to go to the
level of bits. We may take the microstates of the system as being the different permission combina-
tions for the users, e.g. \( q_1 = (\text{read}, \text{write}, \text{mark}) \), \( q_2 = (\text{read}; \text{sally}) \), . . . . A macrostate of the
system is one possibility for the collective permissions of all files and objects of the system, e.g.
(\text{file}1, q_1), (\text{file}2, q_2), . . . .

Example 51. A signal sent from a computer to another, as a binary stream, is represented as a
function \( q(t) \). The values that \( q(t) \) can take, at any time, lie in the set \( Q = \{0, 1\} \), which is the
state space for the data-stream.

Example 52. A user types at a computer terminal, onto a QWERTY keyboard. The data stream
can be represented as a function \( q(t) \), that maps a moment in time to a key typed by the user. The
state space of the stream is the set \( Q = \{q, w, e, r, t, . . . \} \).

Example 53. Data stored on a computer disk are recorded as binary patterns on a series of
concentric rings. Each bit is located physically at a location parameterized by its distance \( r \), from
the centre, and and angle \( \theta \) from some reference line on the disk. The changing binary pattern of
data on the disk is thus a function \( q(r, \theta, t) \) of position and time. At its lowest level of representation,
the state space of the data is the binary set \( Q = \{0, 1\} \).

5.6 CONTINUUM APPROXIMATION

It is often helpful and even important to be able to describe systems in terms of smoothly varying
variables. Just as one would not imagine describing the flow of water in terms of individual discrete
atoms, one does not attempt to describe collective behaviour of many discrete sources in terms of
discrete digital changes. The transition to continuous processes is straightforward. A discrete
chain

\[ X_0 = q_i, X_1 = q_j, \ldots X_n = q_k, \]  \hspace{1cm} (5.17)

is replaced by a function \( q \) of a continuous parameter \( t \), so that a time interval from an initial time
\( t_i \) to a final time \( t_f \) maps into the state space \( Q \):

\[ q(t) : [t_i, t_f] \to Q. \]  \hspace{1cm} (5.18)

The discrete event notation \( X_n \) is now redundant and we can now speak of the value of the state at
time \( t \) as \( X(t) \). \( X \) is the symbol used in most mathematical literature, but we prefer the symbol
\( q(t) \) here. A set of parallel chains, labelled by a parameter \( x \), and time parameter \( t \) is thus written
\( q(x, t) \). The transition matrix is now a function of two times:

\[ T(t, t') = |\langle q(t')|q(t) \rangle|^2 = T(\tilde{t}, \tilde{T}) \]  \hspace{1cm} (5.19)
where
\[ \tilde{t} = t - t', \quad \tau = \frac{1}{2}(t + t'). \] (5.20)

If there is no dependence on the absolute time \( \tilde{t} \), the process is said to be \textit{homogeneous} or \textit{translationally invariant} in time, otherwise it is \textit{inhomogeneous}. A process is said to be memoryless if it does not depend on \( \tilde{t} \), since then it is in a steady state with nothing to characterize how it got there. Invariance under translations of location and time are important in science, because they imply that all states are treated equally, and that this leads to conserved quantities, like the well known accounting parameters energy, transactions, and money.

5.7 THEORY OF COMPUTATION AND MACHINE LANGUAGE

In what we might call Traditional Computer Science, computer systems are described in terms of logical propositions — as automata, working on a usually deterministic set of rules. This area of computer science includes algorithmic complexity theory and automated theorem proving, amongst other things (see for instance [LP97]). Recently the attempt to formalize simple processes has been applied to software engineering, with language-like constructions such as the Unified Modelling Language (UML) (see for instance [Som00]) that apply the forms of rigour to processes that include unpredictability. Although such theories rule a domain of computer science that deals with determinism, this book begins, in a sense, by questioning their broad validity as an approach to human-computer management. Rather than applying a machine model to human-computer interaction, the claim here is that one should apply the scientific, behaviourist approach which has been developed to deal with real world complexity and uncertainty in a systematic way.

Logic is a description of very elementary processes — too elementary to capture the essence of the interaction between networks of humans and computers in actual use; nonetheless, one cannot describe the world without understanding it ‘atoms’. We need logic and reasoning to describe the elementary building blocks before adding the complicating interactions, such as patterns and algorithms. We also need descriptive theories of unpredictable behaviour (computer input is always unpredictable) and of multiple levels of complexity. But as layer upon layer of complexity are compounded, we also need broader scientific ideas that retain determinism in the form of causation, but which abandon the idea of having a complete and exact description about systems.

5.7.1 AUTOMATA OR STATE MACHINES

A finite automaton, or finite state machine, is a theoretical system which responds to changes transmitted to it by a data stream, by signalling \textit{state}. Many mechanisms work in this way. Automata
can be represented or realized in different ways (e.g. cellular automata, push-down automata). The important feature of automata is that they remember state information, i.e. they have a memory of what has happened to them in the recent past. The amount of memory determines the type of automaton.

A finite automaton receives a string of input and makes changes in its internal states (i.e. its memory) depending on the values which it reads at its input. There are physical systems which behave in this way: for instance, multi-state atoms, atoms moving around on metal surfaces etc. If the input is a finite set of photons with frequencies matching different transitions, then that is a simple finite state machine. Some input strings result in final states which are defined to be ‘acceptable’ and all others are unacceptable. Thus a finite state machine can accept or reject strings of digital information. A finite automaton is described by the following components:

1. $Q$ - a complete but finite set of internal states (i.e. memory/registers).
2. $\Sigma$ - a finite alphabet of symbols or digits to be read (e.g. $0, 1; A, B, C, \ldots$).
3. $T = Q \times \Sigma$ - a transition function ($T_{12} = \langle q_1| q_2 \rangle$) of deterministic state mappings, or of transition probabilities.
4. A start state (boundary condition) $|q_s\rangle$
5. A constraint on the allowed transitions.

Automata can do only one type of job: they can distinguish between legal and illegal input. If the system reads in a string, and ends up in an acceptable final state, then the string is said to be legal. If the accepted final state is not amongst the set of legal states, then the input is classified as illegal.

If an input string (a chain) satisfies well-defined rules of construction, it is said to satisfy a grammar. Not all grammars can be parsed by just any automaton. For instance, a simple finite state machine, cannot parse any language that admits an arbitrary level of recursion. The classic example is to think of how you could arrange the states and transitions to parse text containing parentheses.

**Example 54.** A finite state machine can parse a parenthesis embedded string, with maximum nesting level $n$ if it has sufficient internal states. How many states does it need?

$$(\ldots (\ldots (\ldots ))$$

To tackle arbitrary recursion one needs a stack that can remember symbols. The automaton can place a value onto the stack, and retrieve values from it. The stack structure is used in every modern computer because it mimics recursion precisely. If we take the parenthesis example: for every left bracket we add a number to the stack, then for every right bracket we remove one from the stack. The level of the stack is an indicator of which nesting level we are at. At the end of a valid string, the stack should be empty, else the brackets are not balanced. Automata with stacks are called push-down automata.
Example 55. The World Wide Web’s Hypertext transfer protocol, HTTP, is a stateless protocol in the sense that when you visit a web page, it does not remember that you have been there before. However, extensions to the basic protocol using cookies and server-side sessions allow the server to remember the state of transactions that have transpired in the past, e.g. if you have typed in a password, the state changes from “not authenticated” to “authenticated”.

The generalization of the finite state machine (FSM) is the Turing machine and is the model for all modern digital computers. We shall not discuss Turing machines in this book.

![Finite State Machine Diagram](image)

Figure 5.5: A finite state machine representation of a computer process scheduling history.

State machines are a common and useful way of describing certain kinds of systems in both a qualitative or quantitative manner. A state machine that is well known to computer science students is the transition diagram of an operating system process dispatcher (fig. 5.5). This describes transitions between certain states of the system for a given process. These states are not the most microscopic level of describing the system: we do not know the internal code instructions that take place in the operating system of the computer, but these states are characteristic of a ‘black box’ description of the process. The diagram encapsulates rules that the system uses to alter its behaviour.

A similar state machine is found in a rather different system: a service help desk, run by humans (fig. 5.6). The task being performed by these two systems is qualitatively similar to the dispatcher, so the similarity of states should not be a surprise. Of course, humans are not deterministic machines, so they cannot be modelled by exact rules, but the essence of their behaviour can be modelled statistically in any manner that is convenient to the design of a human-computer system (see fig. 5.7). The procedural aspect of the human system can be represented as an automaton, since it is predetermined and has a simple logical structure. Drawing the transition diagram for a finite state machine can be a very useful way of debugging problems in a system.

- Are there states that can never be reached?
- Is it possible for a finite state machine to get into a state from which it cannot emerge?
Figure 5.6: A finite state machine representation for a system administrative help desk.

Figure 5.7: A finite state machine representation for a human being. This is clearly contrived, but this might be a way of modelling the main kinds of activity that humans undergo. A model of transitions between these states is unlikely to be describable by any realistic algorithm, but one could measure the time spent in each state and the likelihood of transitions from one state to another in order to improve the efficiency of the individual’s work pattern.

It might be necessary to change the rules of the system to disallow certain transitions, add more or even add extra states to distinguish between situations that a system cannot cope with. For systems that are probabilistic in nature, so-called hidden Markov models can be used. In a hidden Markov model, there are hidden variables that control how the transitions occur. The transition rules become themselves state dependent.

5.7.2 OPERATORS AND OPERANDS

In order to discuss the active changes made within systems, using a formal framework, we introduce the notions of operators and operands. These terms are encountered routinely in mathematics and in computer science, and have a familiar and well defined meaning.

**Definition 23 (Operator).** An operator is a process which changes the value of a resource variables within the system, by performing an operation on it. It invokes a unit of change $\Delta q$. 
A computer program is an operator which acts on the data within a computer. Human beings are operators when we perform operations. If we want to describe them formally, we need only find a suitable representation for an operator, acting on some operand that mimics the behaviour we need to model.

### Pattern Matching and Operational Grammars

Input to a system is, by definition, unpredictable. If it were not, it could be eliminated and the system could be replaced by a closed automaton. Systems interpret input, usually in the form of symbols, though ‘analogue’ systems often measure continuous input from sensors. To determine the meaning of the data at the input of a system, one must therefore interpret the stream of data and determine its intended meaning. This is a problem that is still a matter of considerable research. There are two main approaches, that are often complementary: grammatical and statistical recognition methods.

If the behaviour of a system is deterministic, it can be described in terms of operations that are executed in some well-defined sequence. A description of this ‘language’ of operation can rightfully be called a machine language. The set of all legal sentences in this language is called the syntax of the language. It does not matter whether humans ever see this language of operation; it exists and is well-defined without ever having to be written down.

The syntax of any language can be modelled by a general theory of its structure, called a grammar. Grammatical methods assume that arriving data form a sequence of digital symbols (called an alphabet) and have a structure that describes an essentially hierarchical coding stream. The meaning of the data is understood by parsing this structure to determine what information is being conveyed. The leads us to the well-known Chomsky hierarchy of transformational grammars (see, for instance, [LP97]).

Using statistical methods of recognition, patterns are digitized and learned, regardless of whether they began in digital form or not. One then gathers statistical evidence about the meaning of previously seen patterns of symbols and tries to use it to guess the meaning of new patterns. This method has been interestingly employed in bio-informatics in recent times ([IDEKM98]) to interpret gene sequences. Even in this case, the idea of grammar is useful for classifying the patterns. A classification of a pattern is a way of assigning one-to-one meaning to it.

The complexity of patterns is related to the level of sophistication required to decipher their structures. Linguist Noam Chomsky defined a four-level hierarchy of languages called the Chomsky hierarchy of transformational grammars that corresponds precisely to four classes of automata capable of parsing them. Each level in the hierarchy incorporates the lower levels: that is, anything
that can be computed by a machine at the lowest level can also be computed by a machine at the
next highest level.

<table>
<thead>
<tr>
<th>State machine</th>
<th>Language class</th>
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<tbody>
<tr>
<td>Finite Automata</td>
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<td>Push-down Automata</td>
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<td>Non-deterministic Linear Bounded Automata</td>
<td>Context-sensitive Languages</td>
</tr>
<tr>
<td>Turing Machines</td>
<td>Recursively Enumerable Languages</td>
</tr>
</tbody>
</table>

State machines are therefore important ways of recognizing input, and thus play an essential part in
human-computer systems.

**Example 56.** Regular expressions are simple regular languages with their own grammar that are
used for matching simple lexical patterns. They are widely used for searching computer systems
with ‘wildcards’ for particular filename patterns. Regular expressions are also used for finding
patterns of input in Network Intrusion Detection Systems.

Because of their regularity and conduciveness to formalization, computer science has seized
upon the idea of grammars and automata to describe processes (see section 13.11). We shall make
some use of these idea, especially in chapter 15, but will not attempt to cover this huge subject in
depth here. Symbolic logics are used to describe everything from computer programs and language
([Log]) to biological mechanisms that describe processes like the vertebrate immune response
([Jer64]). Readers are referred to texts like [LP97, Wat91] for authoritative discussions. For a
cultural discussion of some depth see [Hof81].

5.7.4 PATHWAY ANALYSIS AND DISTRIBUTED ALGORITHMS

In networks or graphs deterministic methods for locating features of the network are often algo-
rithmically complex and many belong to the class NP of algorithms for which there is no known
solution that will execute in polynomial time. Fault isolation is one such problem, as is finding the
shortest path through a network. Approximations are often used to find a reasonable solutions to
these problems.

Distributed algorithms are often discussed in connection with management of networks (see
for instance [SS02]). There is a powerful prejudice towards the use of deterministic algorithms, and
in defaulting to heuristics when these fail to yield desired progress. Computer science has its roots
more in logic than in statistical methods, though some of the latter enter in the field of artificial
intelligence (see, for instance, [Pea00] and [Pea88]). However, increasingly there is a realization
that an algorithmic approach is too elementary for describing systems at a higher level, and thus
we spend little time on discussing such algorithms in this book. A nice overview of the traditional
system management viewpoint is given in [Her94], for instance. Other authors describe algorithmic
tricks for elementary management processes, some of which will make a brief appearance later in
the book (see [Bur04, CS03, CHIK03]).

Example 57. Paxos

Example 58. The CAP conjecture for distributed data consistency, originally postulated by

5.8 A POLICY DEFINED STATE

A theme which recurs in system administration is that of configuration management, i.e. ensuring
that devices, computers and organizations are primed with the correct state configurations in order
so as to behave as we would like. We refer to a description of what we would like a system to do as
a policy, and can imagine that there is an ideal configuration that represents this policy’s wishes\(^1\).
Can we make a system tend towards this state by itself?

In [Bur98b], this idea is called a computer immune system, and the desired state is thought
of as being the ‘healthy configuration’ of the system. If a system will automatically tend to fall
into its own desired configuration, this would require little maintenance (see fig. 5.8). The system
would be self-healing. The task is therefore to design a system with a transition matrix that tends to
lead it towards a fixed, singular configuration, indicated at the centre of the figure. We refer to such
a transition matrix as being convergent\(^2\).

Applications and Further Study 5.

- Describing systems in terms of objects and their classifications.
- Classification of devices and configuration parameters.
- Classification of roles and cliques of components in a system.
- Descriptions of operations and instructions.
- Verification of designs and procedures.

\(^1\) Later we shall advocate more strongly the idea that an ideal state can be associated with policy. For now we can
say that we choose a particular desired state as a matter of policy.

\(^2\) This is sometimes expressed by saying that, in a convergent policy, a repair never makes the system worse.
Figure 5.8: If a system has an ‘ideal’ configuration, we would like this state to be a basin of attraction for the transitions. The state transitions should therefore converge towards a fixed point that we consider to be ‘correct’ according to policy.
Chapter 6

Diagrammatical Representations

Diagrams are useful in planning, designing and understanding systems, because they either conceal details which are not relevant at the architectural level, or amplify details that one normally suppresses.

Diagrams have been used to design and understand systems for thousands of years, in engineering and in medicine, for instance. Complex component diagrams have been in use ever since the first electronic circuits were built. Many themes in electronics have direct analogues in human-computer systems: e.g. flow, amplification, diode (one way flow), resistance, storage, input and output. From the experience of electronic engineering, we know that even fairly simple diagrams can be difficult to understand; huge diagrams are impossible for humans to digest.

Diagrams are helpful as maps, as a blueprint for construction, and even for diagnostics, but they become quickly unwieldy unless they are broken down into manageable pieces. One thus uses a strategy of divide and conquer, or modularity to make systems comprehensible. This results naturally in a hierarchical sub-structure to any large system, in which high levels assume and depend on the correct functioning of lower levels.

It is not easy to formalize diagrams. A diagram is itself a data structure, in which the only resources one has are spatial extent, shapes and colours. Since the space on a page is limited, one usually runs into space limitations long before a formal scheme for organizing that space succeeds in showing anything like a realistic example, i.e. before becoming unwieldy and impossible to grasp. Moreover, there is a limit to the number of ways of making distinctions in an intuitive fashion. For this reason, diagrams are mainly used a visual aids for more formal descriptions using algebra or some kind of functional pseudo-code.
CHAPTER 6. DIAGRAMMATICAL REPRESENTATIONS

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<td>Object size</td>
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<tr>
<td>Shape</td>
<td>Difficulty of recognition</td>
</tr>
<tr>
<td>Colour</td>
<td>Colour perception</td>
</tr>
<tr>
<td>Direction</td>
<td>Only 2 dimensions</td>
</tr>
</tbody>
</table>

Table 6.1: Freedoms and constraints for printed diagrams

There are top-down and bottom-up approaches to diagrams.

6.1 DIAGRAMS AS SYSTEMS

Diagrams are systems in which one uses space as a resource to represent information about something else. An effective diagram, therefore requires a proper allocation of the resource of space to the problem of representing information. The freedoms and constraints one has in a diagram are shown in table 6.1.

Given that a diagram is dependent on these factors, the task is one of how to allocate the space and shapes creatively in order to achieve the goal of the diagram. How is space used? Ideally, every diagram would indicate what its premises are. Some diagrams use space to indicate time, some use space to indicate extent. Some use distance to indicate some measurable property of the system.

Visual representations are, of course, not the only way of representing information. Underwater animals, such as dolphins, communicate mainly by sound, blind people mainly by touch, and so on. In ref. [GC00] the authors employed a ‘sound diagram’ to the problem of representing a computer network. An audio representation presents different limitations, such as ease of distraction from the environment. Electronic diagrams, on computers, can combine sight and sound and other senses into a single representation, but this goes beyond the limitations of a diagram on a printed page.

Example 59. Electrical circuit diagrams: the flow of activity is carried by electricity, but the function of the circuit might be something as abstract as playing music (a radio). The actual function is rather hard to see from the internal algorithms, and yet those details are required at the level of flow. The mechanics of individual electrons is not required, because one trusts that the components behave in a predictable fashion.

6.2 THE CONCEPT OF A GRAPH

There are many situations where we draw dots that are joined together by lines, perhaps with arrows on them, to denote some kind of information. The links between nodes often represent qualities
such as

- $A$ dominates $B$ (directed).
- $A$ depends on $B$ (directed).
- $A$ is associated with $B$ (undirected).

and so on. Note that some of these relationships are one-way (directed) and others are multi-way (undirected).

D. König suggested the name *graph* for all such diagrams, and pioneered the study of their properties. Elementary graph theory is a very useful framework for discussing human-computer systems, because it considers systems with discrete states (the nodes or dots in the graph) that are joined together only if certain criteria are met (i.e. if there is an arc joining the points, perhaps with an arrow going in the right direction).

![Graph Example](image)

**Figure 6.1:** A graph is a general assembly of nodes connected by arcs. It is used to describe many situations in science and mathematics.

A graph with arrows on it is called a *directed graph* and a graph without loops is called *acyclic*; thus a tree structure (so common in computer science) is an acyclic directed graph.

**Definition 25** (Graph). A graph is a pair $(X, \Gamma)$ that consists of a set of nodes $X$ and a mapping $\Gamma : X \rightarrow X$, formed by the arcs or lines between the points $x \in X$.

The degree of any node in a graph is the number of nearest neighbours it has, i.e. the number of nodes that can be reached by travelling along those links that are connected to the node.
**Definition 26** (Degree of a node). *In a non-directed graph, the number of links connecting node $i$ to all other nodes is called the degree $k_i$ of the node. In a directed graph, we distinguish incoming and outgoing degrees.*

Do not be tempted to think of a graph as being necessarily composed of only a sparse set of points with occasional links between them. Even a dense set of points, infinitely close together, such as all the points in a circle form a set and can be mapped onto each other, even if the arcs seem to overlap the points (see fig. 6.2).

![Figure 6.2: A graph mapping points within a disk onto other points within the disk.](image1)

![Figure 6.3: A simple network, represented as an undirected graph.](image2)
A graph may be represented or defined by its adjacency matrix\(^1\). By convention, the adjacency matrix of a network or graph is a symmetric matrix with zero leading diagonal, (see, for example, the graph in fig. 6.3)

\[
A = \begin{pmatrix}
0 & 1 & 1 & 1 \\
1 & 0 & 0 & 1 \\
1 & 0 & 0 & 1 \\
1 & 1 & 1 & 0 \\
\end{pmatrix}
\] (6.1)

**Definition 27 (Adjacency matrix).** A square matrix \(A_{ij}\) whose rows and columns label the nodes of a graph. If a path exists from node \(i\) to node \(j\) then \(A_{ij} = 1\). If the graph is undirected it must have a symmetric adjacency matrix.

If we regard the nodes in a graph as possible states in a system, then the adjacency matrix is also a transition matrix between states, where a 1 indicates the possibility of a transition. For non-deterministic or stochastic systems, one could replace “1” by the probability of a transition taking place.

Graphs or networks come in a variety of forms. Figure 6.4 shows the progression from a highly ordered, centralized structure to a de-centralized form, to a generalized mesh. This classification was originally discussed by Paul Baran of RAND corporation, in 1964 as part of a project to develop a communications system that would be robust to failure, in the case of a nuclear attack (see [Buc02, Bar02] for a review).

### 6.3 PROMISE THEORY

An extension of the graph theory known as Promise Theory\(^2\) [Bur05, BB14] makes use of networks of interactions between nodes that model ‘agents’ and the promises they make some each other. This subject will be dealt with more fully in volume 2, since this theoretical approach has many important qualities, such as the ability to unify intentional aspects of systems with quantitative analysis.

The ‘Promise Theory’, was introduced in 2005 as a way to model distributed systems with complete decentralization[Bur05]. Coupled with abstraction, it offers a looking glass onto the design and management of networks. If we define what a user or application needs from the network we can begin to get away from imperatively controlling the ‘how’ the network functions and instead

\(^1\)Working with graphs in adjacency matrix form becomes decreasingly realistic as the size of the graphs grows beyond thousands of nodes, however this form is very illuminating and is particularly suited to smaller graphs such as those that arise in system administration.

\(^2\)This topic has been added in the second edition.
focus on declaratively describing "what" is required from it. In Promise Theory, network elements act as autonomous agents and collaborate to find the best way to deliver the required function. Promise Theory has become an invaluable measuring stick for describing systems, with semantics and dynamics (intent and outcome) side by side. Promise theory is about what can happen in a collection of components that work together [Bur05, MK05]. It is not a network protocol, but a descriptive algebra. One begins with the idea of completely autonomous agents that interact through the promises they make to one another. It is well-suited to modeling networks [BC04]. Although we cannot force autonomous agents to work together, we can observe when there are sufficient promises made to conclude that they are indeed cooperating voluntarily. Our challenge in this paper, is to translate this bottom-up view into top-down, human managed requirements.

**Definition 28 (Agent).** The term used for the fundamental entities in Promise Theory. Agents may be any active entities in a system. Agents are the seat of intentions, whether directly or by proxy. They are autonomous (independent) in the sense that they may not make promises on behalf of anything except their own behaviour, nor can they be influenced by outside agents without explicitly promised consent.

The promise formalism has a number of features, described in [BB14]. We refer readers to this reference for details. More details will be given in volume 2.
Definition 29 (Promise). A promise is an intention that has been ‘voluntarily’ adopted by an agent (usually channeling a human owner, or perhaps an agreed standardization).

An agent, which only promises to do as it’s told, is dependent or voluntarily subordinated. It has some of the characteristics of a service: an agent makes its intended behavior known to other agents (e.g. I will serve files on demand, or forward packets when I receive them). An imposition is an attempt to induce the cooperation of another agent by imposing upon it (e.g. give me the file, take this packet).

We write a promise from Promiser to Promisee, with body $b$ as follows:

$$\text{Promiser} \xrightarrow{b} \text{Promisee.}$$

(6.2)

and we denote an imposition by

$$\text{Imposer} \xrightarrow{b} \text{Imposee.}$$

(6.3)

Promises and impositions fall into two polarities, denoted by $\pm$. A promise to give or provide a behavior $b$ is denoted by a body $+b$; a promise to accept something is denoted $-b$ (or sometimes $U(b)$, meaning use-$b$). Similarly, an imposition on an agent to give something would have body $+b$, while an imposition to accept something has a body $-b$.

Although promises are not a network protocol, agents can exchange data. To complete any kind of exchange, we need a match an imposition ($+$) with a promise to use ($-$. To form a binding (as part of a contract), we need to match a promise to give ($+$) with a promise to use ($-$. This rule forces one to document necessary and sufficient conditions for cooperative behaviour.

A promise model thus consists of a graph of nodes (agents), and edges (either promises or impositions) used to communicate intentions. Agents publish their intentions and other agents may or may not choose to pay attention. In that sense, it forms a chemistry of intent [Bur13b], with no particular manifesto, other than to decompose systems into the set of necessary and sufficient promises to model intended behavior.

Example 60. The Ethernet protocol may be described simply in terms of promises. The agents that keep promises to send and receive data are the network interfaces. For example, in the Ethernet protocol, interfaces $E_i$ promise to label transmissions with a unique MAC addresses or string of digits.

$$E_i \xrightarrow{+\text{MAC}_i | \text{MAC}_i \neq \text{MAC}_j} E_j \quad \forall i, j$$

When data are transmitted by an interface, the interface keeps its promises to use messages that have (destination MAC address, data). Note: the message is not a promise, the promise governs how the message is handled.

$$E_i \xrightarrow{(+\text{MAC}_j, +\text{data})} E_j$$
Messages are sent ‘fire and forget’ as impositions on to a remote receiver. While all interfaces generally promise to accept any MAC address, (unless they block with MAC access control) only the interface whose MAC address matches the destination in the message doublet actually promises to accept the message voluntarily. Note, there is nothing other than convention to prevent all agents from accepting the data too; this ‘promiscuous mode’ is used for network monitoring, for example.

\[
\begin{align*}
E_{\ast} & \xrightarrow{-\text{MAC}_j} E_{\ast} \quad \forall i, j \\
E_i & \xrightarrow{(-\text{MAC}_j, -\text{data}) \text{ if } (i=j)} E_j
\end{align*}
\]

Since the channel is unprotected, agents effectively promise the data to all others in scope. Moreover, all agents promise to decode the address and the data, but many will discard the results. While this set of promises is scale independent, the assumption that every agent has to be in scope of every transmission does not scale, since it requires messages to be flooded or broadcast to every node (agent), in principle. The primary issue with raw Ethernet is that there are no ways to selectively limit the size of these broadcast domains. This makes the ‘everyone please take a look at this’ approach impractical.

![Figure 6.5: An Ethernet switching function.](image)

In Fig. 6.5 we see two interfaces that promise MAC address 00:00:11:11:11:AA (shortened to AA) and 00:00:11:11:11:BB (shortened to BB). Suppose we wish to send data from AA to BB, then, since the Ethernet is a push-based imposition protocol, only half a contract is needed for emergent delivery, and we leave the rest to trust.

\[
\begin{align*}
E_{AA} & \xrightarrow{+\text{MAC}_{BB}} E\text{switch} \\
E_{\text{switch}} & \xrightarrow{-\text{MAC}_i} E_i \quad \forall i \\
E_{\text{switch}} & \xrightarrow{\text{forward MAC}_{BB}} E_{BB}
\end{align*}
\]

In each point-to-point interaction, the agent has to formally promise to use (-) the delivery service promised by the agent giving (+). This is the algebra of binding. There is no notion of a
permanent virtual circuit, as say in ATM. However, if we add handshaking, a similar story can be told about ATM, Frame Relay, MPLS and other systems.

**Example 61.** The Internet Protocol may also be rendered as in the previous example. IP provides Wide Area Networking by issuing two part addressing to cope with transmission scalability. IP addresses still promise to be globally unique, but are interpreted as doublets.

\[(network \text{ prefix}, \text{ local address})\]

Only addresses with the same prefix are considered in mutual scope for broadcasting, and messages addressed from one prefix to another promise to be forwarded deliberately rather than by ‘flooding’. IP is thus a cooperative effort that builds on promises rather than impositions alone.

To make this work, IP needs two kinds of agent, which fall into different promise roles (see figure 6.6): interfaces (terminating connections), which only transmit and receive data intended for them, and forwarders (called routers or switches) that cooperate with multiple interfaces, and promise to selectively forward data from one interface to another between protected broadcast domain. This acts as a flood-barrier or firewall to packets promised to different prefixed networks.

To model routers, without giving up the interface abstraction, we introduce the concept of a route service (or link service), whose job it is to establish cooperative forwarding between the interfaces.

![Diagram of Internet promises](image)

Figure 6.6: Internet promises. An end-node or leaf and its single interface promises to relay through a ‘router’ which is surrounded by multiple interfaces, thus connecting multiple network branches.

Consider Fig. 6.6. The source node has an address, normally written 128.39.78.4/24. As a doublet, the promises see it in two parts as \(i = (prefix=128.39.78, local=4)\). We’ll call this the source prefix, or, \(j = (prefix=128.39.78, local=1)\) for the router interface. When a message is sent
to an address with a different destination prefix, data are sent by imposition to the interface on the router with the source network prefix (usually the ‘default route’):

\[ I_{source_i} \xrightarrow{+(destination,local),+data} I_{router_j} \]

Each router interface \( j \) promises the connected source interfaces \( i \) to use all such packets, a priori, and to present them to the router (kernel) which keeps the following promises.

\[ I_{router_j} \xrightarrow{-(*,*),-data} I_{source_i} \]

\[ I_{router_j} \xrightarrow{+prefix,+data} Router \]

Similarly, other interfaces connected to the router’s interfaces promise to accept messages from the router that have their prefix:

\[ I_{source_i} \xrightarrow{-\{prefix,source\},+data} Router_j \]

Crucially for messages to escape from a local region, the router promises all IP interfaces to forward messages it receives on one if its own interfaces according to a set of promises which we denote ‘forward’. The router interfaces, in turn, bind to this promise by accepting it.

\[ Router \xrightarrow{+forward} I_{router_j} \]

\[ I_{router_j} \xrightarrow{-forward} Router \]

The forward promise has the following logic:

1. If the prefix of the destination interface is the same as the prefix of one of the router’s interfaces, forward the message onto that interface.

The remainder of the promise requires configuration with knowledge of the wider world.

2. If the prefix of the destination interface is known to an internal database of external knowledge, i.e. the Routing Information Base (RIB), forward the message to the interface known to lead to the desired destination.

3. Send all other message destinations to a pre-decided default interface, where we expect to reach some other router with greater knowledge of how to find the prefixed network.

Note that, like the Ethernet, this algorithm has only emergent behaviour that matches its design goal. It cannot, by direct imposition, assure a successful delivery of messages, because that requires the cooperation of potentially many intermediate interfaces and routing agents. In spite of this apparent lack of control, the Internet works demonstrably well. Trust plays a major role in operations.
6.4 CONNECTIVITY

Let \( \vec{h} \) be a host or node vector whose components are 1 if a host is available and zero if unavailable. The level of connectivity within a graph or closed network can be characterized by an invariant scalar value \( \chi \):

**Definition 30** (Connectivity). The connectivity, \( \chi \), of a network \( N \), is the probability (averaged over all pairs of nodes) that a message can be passed directly between any two nodes. \( \chi \) may be written as

\[
\chi = \frac{1}{N(N-1)} \vec{h}^T A \vec{h}.
\]

(6.4)

\( \chi \) has a maximum value of 1, when every node is connected to every other, and a minimum value of zero when all nodes are disconnected.

The connectivity of a graph is of great importance to systems. It tells us both how easy it is for information to spread throughout the system and how easy it is for damage to spread. This duality is the essence of the security-convenience conundrum. We explore this issue further in the next section.

6.5 CENTRALITY: MAXIMA AND MINIMA IN GRAPHS

Where are the best connected nodes in a graph? These are nodes that we would like to identify, for a variety of reasons: such nodes have the greatest possible access, to the rest of the system. They might be security hazards, bottlenecks for information flows, or key components in a system in which we need to invest money and resources to keep the system running smoothly. From the standpoint of security, important nodes in a network (files, users, hosts) are those that are 'well-connected'. We are thus interested in a precise working definition of 'well-connected' (see [BCE04a, CEM04]).

A simple starting definition of well-connected could be 'of high degree': that is, count the neighbours. We want however to embellish this simple definition in a way that looks beyond just nearest neighbours. To do this, we borrow an old idea from both common folklore and social network theory (see [Bon87]): an important person is not just well endowed with connections, but is well endowed with connections to important persons.

The motivation for this definition is clear from the example in figure 6.7. It is clear from this figure that a definition of 'well-connected' must look beyond first neighbours. Nodes of equal degree have quite different levels of importance depending on their position within the remainder of the graph.
We can now formulate a precise definition of the importance for non-directed graphs using a concept called centrality. We begin by noting that the symmetrical adjacency matrix consists of ones where a node has neighbours and zeros where it does not. Thus multiplying a row of this matrix by a column vector of ones would simply count the number of neighbours for that node. We can use this fact to self-consistently sum the entire graph.

Let $v_i$ denote a vector for the importance ranking, or connectedness, of each node $i$. Then, the importance of node $i$ is proportional to the sum of the importances of all of $i$’s nearest neighbours $N(i)$:

$$v_i \propto \sum_{j=N(i)} v_j.$$  

(6.5)

This may be more compactly written as

$$v_i \propto \sum_j A_{ij} v_j,$$  

(6.6)

where $A$ is the adjacency matrix. We can rewrite eqn. (6.6) as

$$A \vec{v} = \lambda \vec{v}.$$  

(6.7)

Thus the importance vector is actually an eigenvector of the adjacency matrix $A$. If $A$ is an $N \times N$ matrix, it has $N$ eigenvectors (one for each node in the network), and correspondingly many eigenvalues. The eigenvalue of interest is the principal eigenvector, i.e. that with highest eigenvalue, since this is the only one that results from summing all of the possible pathways with a positive sign. The components of the principal eigenvector rank how self-consistently ‘central’ a node is in the graph. Note that only ratios $v_i/v_j$ of the components are meaningfully determined. This is because the lengths $\sum_i v_i^j v_i$ of the eigenvectors are not determined by the eigenvector equation.
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The highest valued component is the most central, i.e. is the eigencentre of the graph. This form of well-connectedness is termed 'eigenvector centrality' (see [Bon87]) in the field of social network analysis, where several other definitions of centrality exist. We shall use the terms 'centrality' and 'eigenvector centrality' interchangeably.

**Example 62.** Consider the graph in fig. 6.8. This has adjacency matrix

\[
A = \begin{pmatrix} 0 & 1 & 0 & 1 & 1 \\ 1 & 0 & 1 & 0 & 1 \\ 0 & 1 & 0 & 1 & 1 \\ 1 & 0 & 1 & 0 & 1 \\ 1 & 1 & 1 & 1 & 0 \end{pmatrix}
\] (6.8)

The matrix has eigenvalues \( \lambda = \{-2, -1.2, 0, 0, 3.2\} \). The principle eigenvector is that associated with the last of these, i.e. that with the highest value.

\[
\vec{P}(A) = (0.43, 0.43, 0.43, 0.43, 0.52).
\] (6.9)

This indicates that node 5 is the most central. The remaining symmetrical nodes are have symmetrical lower values. Thus the principal eigenvector maps the topography of the undirected graph.

![Figure 6.8: An undirected graph with an obvious centre.](image)

Figure 6.9 shows a centrality organized graph of the Gnutella peer to peer network. Peer to peer networks are characterized by having no real centre. The centres here are only marginally higher than their surrounding nodes.

**Definition 31** (Eigenvector Centrality). The Eigenvector centrality of node \( i \) is the \( i \)th component of the principal eigenvector \( \vec{v} \), normalized so that \( \max_i v_i = 1 \). This is used in importance ranking of nodes.
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6.6 RANKING IN DIRECTED GRAPHS

So far we have considered eigenvector centrality in undirected graphs, where its meaning is unique and unambiguous. The same idea can be applied to directed graphs, but with additional subtleties.
The topic of ranking nodes in a directed graph has a history associated with importance ranking by search algorithms for the World Wide Web (see [PBMW98] and [Kle99]).

The arrows on graph edges now identify nodes of three types: sources, sinks and relays.

**Definition 32** (Source). A source is a node from which a directed flow only emerges. In the adjacency matrix, this is represented by a row of 1’s with a corresponding column of 0’s.

**Definition 33** (Sink). A sink is a node that is purely absorbing. In the adjacency matrix has a column of 1’s and corresponding row of zeros.

**Definition 34** (Relay). A relay is a node that has both incoming and outgoing flows.

Principal eigenvector ranking is fraught with subtlety for directed graphs. It does not necessarily have a simple answer for every graph, though it seems to work for some (see section 11.8); nor is there a unique procedure for obtaining an answer for every case. All is not lost, however. Once can still use the spirit of the eigenvalue method to learn about graph structure and importance.

There is a fundamental duality about directed graphs that depends on the direction of the arrows. To understand this, look at the two graphs in figs. 6.10 and 6.11.

![Figure 6.10: A directed graph with a sink, i.e. a node (5) that is absorbing of all flows. This is the complement or dual of fig. 6.11.](image)

In the first of these graphs (fig. 6.10), we have a number of nodes with arrows that point mainly towards an obvious centre. This centre happens to be a sink, i.e. it absorbs the flows. This
diagram has an adjacency matrix $A$. The dual picture (fig. 6.11) is described by the transpose of the adjacency matrix $A^T$, since interchanging rows and columns changes the direction of the arrows. There are thus at least two kinds of importance in a directed graph, and that these are complementary:

- **Sink importance**: a node is important if it gets a lot of attention from other nodes, i.e. if it absorbs a lot of flows. This kind of node is also referred to as an authority, since others point to it and hold it in esteem.

- **Source importance**: a node is important if it influences a lot of nodes, i.e. if it originates many flows. This kind of node is also called a hub because it shoots out its spokes of influence in all directions.

The problems with this simple use of the principal eigenvector are illustrated by the two diagrams in fig. 6.12. These graphs cannot be distinguished by the basic eigenvector method, since there is such great symmetry that all path lengths from the nodes are equal. The eigenvalues are thus all zero and there is no way of finding a most important node, despite the fact that the central node clearly has a privileged position. An interesting approach to directed graphs that is able to distinguish these graphs has been presented by Kleinberg ([Kle99]). The following symmetrized matrices enforce the duality noted above explicitly, by tying together sources and sinks into nearest neighbour ‘molecules’. A good source (hub) is one pointing to many good sinks and a good sink is...
one pointed to by many good sources, at nearest neighbour level. They are guaranteed to have a principle eigenvector.

\[ A_A = A^T A \]  \hspace{1cm} (6.10)

points to a hub (i.e. its principal eigenvector assigns a high weight to a hub or source), while

\[ A_H = AA^T \]  \hspace{1cm} (6.11)

points to an authority (i.e. its principal eigenvector assigns a high weight to an authority or sink), while

\[ A \leftrightarrow A_H = AA^T \]
\[ A^T \leftrightarrow A_A = A^T A. \]  \hspace{1cm} (6.12)

See section 11.8 for an example.

In the hub-authority viewpoint, the matrices attach importance explicitly to local clusters of sources and sinks, but why not longer range dependencies? Another approach is used by the PageRank algorithm (see [PBMW98]) in which stochastic noise is added to the actual adjacency matrix in order to generate loops that resolve the eigenvalue problem. The success of the procedure is well known in the form of the search engine Google, but it is also an arbitrary procedure that is now patented and out of public scrutiny. The issues surrounding importance ranking will likely be resolved in future work, allowing methods to be used as organizing principles for systems of all kinds.
6.7 Applied Diagrammatical Methods

A variety of heuristic diagrammatical forms are in use.

- **Maps**

  Maps (or ‘mind maps’) are a loose representation of everything in a system. They are used as a basis for identifying relationships and structure, as well as simply documenting all the relevant parts (see fig. 6.13).

![Figure 6.13: A ‘mind map’ of an enterprise, showing a brain-storming approach to understanding the elements that are important, and their relationship to one another.](image)

Note this is not a dependency diagram. It is simply a diagram of associations.

- **Flow diagrams**

  These were common in the early days of programming and are still used sometimes for representing simple algorithms graphically. Flow diagrams show the causal sequence of actions and the decision branches in simple processes; they are a graphical pseudo-code and thus provide a very low level picture of a system. For large or complex systems, flow diagrams become unwieldy and more of a hindrance to comprehension than an aid (see fig. 6.14). The Unified Modelling Language (UML) attempts to extend this idea to make diagrams express a strict grammar.

- **Transition diagrams**
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A system of distinct states is called a Finite State Machine (see section 5.7.1). It is formally represented as a directed graph. Finite state machines are at the centres of many systems (see fig. 5.5). They represent a coarse type of memory of context in a system. Transition diagrams are related to functional structure diagrams. If one labels being in each function as a state of a process, then they represent the same information.

- **Functional structure diagrams**
  A structure diagram is a chart of the independent functions within a process and the flow of information between them. They are sometimes called Jackson diagrams ([Jac75]).
  A structure diagram describes a chain of command and maps the independent methods in a process, showing how they relate to one another. For instance, the functional declaration

  ```
  begin function A
  do function B
  do function C
  end function
  ```

  would be drawn as in fig. 6.16. By the same token, it charts the flow of resources in a system between the high level objects within. This diagram limits its view to the top level structure, so its value is limited; it is most useful for top-down approaches to system design.

- **Dependency diagrams**
  Dependency diagrams are the basic tool for fault analysis. Each arrow in a dependency diagram should be read ‘depends on’ (see fig. 6.18 and fig. 6.19). If a component or object in a system fails, all the components that point to it will also fail. A fault tree is a special case of a dependency diagram.
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Figure 6.15: An excerpt of a functional diagram showing the economic (resource) organization of an enterprise. In many organizations this doubles as a chain of command diagram, since control is often chosen to flow with the dissemination of resources, though this need not be the case.

Figure 6.16: A simple Jackson diagram for a function with two dependencies.

In these diagrams we see the repeated significance of the process of classification and subclassification of objects., The role of object orientation and type distinction is demonstrated.

• **Entity relation diagrams**

Entity Relation (ER) diagrams are used in the description of database tables, for example, in the Structured Query Language (SQL). They describe the interrelationships between objects of different types, as well as the cardinality (number of objects that are involved in each type of relationship) that are possible between objects of different types (see fig. 6.20).

Object diagrams that describe object-orientated programming relationships are simplified forms of ER diagrams.

• **Petri-nets and stochastic networks**
Figure 6.17: An excerpt from a functional diagram showing the resource organization of a project, within a larger enterprise. This diagram takes a two dimensional view of an organization, by plotting the two degrees of freedom project development versus dependency structure.
Petri nets (see [DA94, MMS85]) are a graphical tool for modelling discrete event systems, e.g. network services. They are similar in concept to flow diagrams, but are able to describe concurrency, synchronization and resource sharing. Stochastic networks are a related form of network for modelling discrete stochastic events. These topics are beyond the scope of the book.

Diagrammatical methods are only representations of systems and the systems they represent can often be described more compactly in other forms, such as by algebraic rules. The utility of diagrams lies mainly in human understanding and pedagogy; even graph theory relies more on algebraic construction than on pictures. Some attempts have been made to use diagrams to describe system behaviour in a rigorous way. The Unified Modelling Language (UML) is one such attempt.

Applications and Further Study 6.

- Identifying the main scales and structures in a system.
- Classifying system components.
- Visualizing relationships and roles between the components.
- Evaluating the importance of components within systems.
- Finding weak points or inconsistencies in systems.
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Figure 6.19: An excerpt of a dependency diagram for an enterprise, such as that from figs 6.13 and 6.15.

Figure 6.20: Entity relation diagrams describe the structure of tabular data relationships. Each square box is a tabular object with references to fields shown as ellipses. The links show $N : M$ relationships.
CHAPTER 7

SYSTEM VARIABLES

To go beyond pictorial representations and be able to make quantitative statements about systems, we must develop a representation of system properties in terms of variables. One of the difficult notions to understand in the science of systems is how to describe their changing properties. In particular, the transition from discrete, digital changes to smooth differentiable functions is unfamiliar in computer science. Both types of variable are required to describe human-computer systems.

7.1 INFORMATION SYSTEMS

Any system can be thought of as an information system (i.e. as the abstract development of a set of variables), simply by describing activity with the help of an abstract representation, i.e. by viewing a change performed by the system as being a change in a description of the system’s basic resources. This is a one to one mapping. The resources themselves describe and are described by variables, which yield information.

Example 63. A factory which manufactures cars is an information system, because it receives input in the form of steel and changes the pattern of the steel into new shapes, described by a certain amount of information, before emitting cars as output. We can describe this system in terms of the actual physical cars that are produced, and the resources that go into them, or we can describe it on paper, using numbers to represent the flow of items. Accountants regularly consider businesses as information systems; accountants do not get their hands dirty on the factory floor.

The converse is also true: in order to represent information, we must encode it in a physical pattern, represented in some physical medium (brain cells, paper, computer storage, steel etc). To create a pattern, there must be an attribute of some basic resource which can vary, such as colour,
height, size, shape, etc. and these attributes belong to physical objects. Thus it would be wrong to try to divorce systems from their physical limitations, even though we are interested in abstracting them and speaking only of their information. Many of the limitations in systems arise due to the physical nature of their representation.

**Example 64.** Money is a form of information which conveys the power of purchase between individuals. Money began as a form of ‘I owe you’ (IOU) note during barter, where immediate goods for trade were not available. Later it was embodied by physical coins and notes, with no intrinsic value. In modern society, money is created electronically by banks through loans, without any basis in a physical reality. Money is information of accounting. Its value lies in its availability, non-specificity (fungibility), and the widespread acceptance by others in exchange for goods and services.

7.2 **Addresses, Labels, Keys and Other Resource Locators**

In order to speak of change of location, we need to be able to measure and parameterize location. This is not necessarily like measuring things on a ruler. If the locations are named rather than measured, i.e. if it is the name rather than the distance which counts, then one does not use a continuously varying parameter like $x$ or $t$ to describe it, but rather a discrete number or label.

Once the medium for representation of information change has been identified, there is the question of where and when the change takes place in that medium. Describing the when and where in a system is called the parameterization of the system.

**Low Level Parameters**

Ultimately, any system is bounded by physical limitations. In the real world, there are only two variables which can parameterize change in a medium: location (space) and time. Physical objects exist only in space and time. Thus, at the lowest level, a physical dynamical system can only consist of values of the systems’ resources at different spacetime locations. Mathematically, we denote this as functions which vary in space and time:

$$q(\vec{x}, t).$$

(7.1)

**Example 65.** Morse code is a pattern of sounds in time. A picture, or shape is a pattern of material in space. A construction site is a pattern in both space and time, since it has a definite form in space, but the form changes with time.

While space and time are sufficient to describe any system, they are too low-level to be satisfactory in the description of abstract systems. One can create abstract freedoms, such as the
orientation of a non-trivial shape, or the loudness of a sound, by building on ideas of space and time (this is what physics does), but it is not always necessary to refer to such low level details.

**Derived Addresses**

In the virtual world of abstract information, and hence of human-computer systems, there are other ways of parameterizing spatial change (location):

- Geographical location of data.
- Memory location inside a computer.
- Internet address.
- Identity of a container.
- Ownership.

Each of these examples is an *address*, i.e. a label which denotes the location of a resource. We may thus parameterize a change in terms of the value of a resource at an arbitrary address and a particular time. Mathematically, we write this as a function:

\[ q(A, t). \]  

(7.2)

**Higher Level Patterns: Associations**

A common generalization of the idea of a resource address is employed in abstract information systems, such as databases, where one does not wish to refer to the physical location of data, but rather to an abstract location. Instead of thinking of a pattern as being a function of a particular location \( q(A, t) \) one can use a reference key. The key itself is just a pseudonym-label or alias for the detailed physical address:

\[ q(A, t) \rightarrow q(k, t), \]  

(7.3)

where the key \( k \) can be any collection of labels or ‘coordinates’ in whatever abstract space one cares to imagine. e.g.

\[ q(k, t) \rightarrow q(\text{building, town, country, floor, project, t}) \]  

(7.4)

Now, instead of thinking of pattern and structure as varying with address and time, one views it as varying with different values of the abstract key. Thus we can say that a system is a dynamical function of one or more abstract keys.

The *key* forms an abstract representation of the properties of the data structure within the system; one creates an *association*, or *associative relation* between a *key* and a resource *value*.
located by that key. Note that, in writing \( q(k, t) \), the time at which the change takes place is itself simply a label identifying the time at which the value was true.

**Example 66.** The arrival of incoming packets, on a network connection, is a signal that can be described by a function \( q(t) \), that varies according to the chosen representation of the signal, e.g. a binary signal with \( q \in \{0, 1\} \).

**Example 67.** Consider a pattern of data in the memory of a computer, or on some storage medium, described by \( q(\vec{x}, t) \), where \( \vec{x} \) labels the objects of which the pattern is composed. The pattern might be a software package, or an image etc. Since the data change over time, at some rate, this function also depends on time.

**Example 68.** Suppose that the software package in the previous example is determined by \( n \) criteria or tests that determine its integrity or correctness. The package thus has \( n \) state variables, encoding \( 2^n \). Let us give new coordinate labels to all such packages on the system, \( \vec{X} \), and consider the variable \( Q(X, t) \), where \( Q \in \{Q_1, \ldots, Q_n\} \). This variable now describes the state of software packages over time. The value describes the relative integrity of the system.

**Example 69.** A graph is characterized by \( n \) nodes, linked together. Any property of graph nodes can be represented by vectors of objects \( \vec{v} = (v_1, v_2, \ldots, v_n) \). Any property of links in the graph can be represented by a matrix \( A_{ij} \) of appropriate objects or values.

Describing systems formally using variables with particular representations is limited only by the imagination. We are free to do whatever is helpful or productive.

### 7.3 Functional Dependent Relationships

What variables should we look to to characterize the behaviour of a system? If we assume that the outcomes function correctly (with the correct semantics) then all that is left is performance. In any network of parts, i.e. system, communication is the main throttle to performance. This suggests several \( q(t) \) worth following:

- **Latency** of communication: what delays between stimulus and response?

- **Traffic intensity**: what is the total amount of traffic as a fraction of the total capacity at each moment over repeated social intervals? It is well documented that the human working week drives patterns of traffic[BHS98, BR00, BHRS01]. This is called the traffic intensity (see section 12.5).

- **Number of errors** detected per unit time, relative to some semantic probe. In general we have to construct probes to measure specific semantics. These are called unit tests and acceptance tests in software development.
Sharing collisions contention events in resource sharing.

Special probes to measure breaches of policy. These may relate to security breaches or performance envelope breaches.

The semantics of variations in these variables are easy to encode, by measuring the distributions of values to detect anomalies, so that automated policy responses can respond immediately to regulate and maintain system state. Automated machine learning is one way to do this.

Example 70. In server management, the system monitoring engine CFEngine’s monitor daemon performs continuous machine learning of variables, gathering distribution statistics. Human administrators may then express the semantics of anomalous measurements and automate triggered responses. By setting a policy for relative conditions of the system, one turns system sampling and measurement into event generation for automated response[Bar06, BB05a].

An important use for variable quantities is to express relationships between measurables. Comparisons and relationships between quantities are the basic tools one has for expressing conditions of a system, and stating policy about them. To be able to express and enforce what we want, we must be able to compare what is measured with what we specify.

Example 71. In system performance, configuration and resource usage, we make various comparisons:

- Rate of job arrivals (<, >, ≥, ≤, =) rate of processing.
- State \( q(x, y, z, t) > q(x', y, z, t) \).

Example 72. Some variables are related to one another in linear combinations:

- Total system communications capacity \( C_T = \sum_{i=1}^{N} C_i \).
- Average traffic at location \( x \) is the sum of traffics from locations \( x' \in S \): \( T(x) = \sum_{x'} T(x') \).

Linear combinations of variables are often useful in parameterizing systems where hidden relationships occur.

Example 73. Suppose one finds that the probability of a program crash is a function of the number of users logged on to a computer and the number of processes being executed in separate measurements. One might observe that most processes are started identically for all users (e.g. Window manager processes) and that only negligible differences are measured between users. In this case we notice that \( N_{\text{procs}} \propto N_{\text{users}} \) and thus:

\[
P_{\text{crash}}(N_{\text{procs}}, N_{\text{users}}) \to P_{\text{crash}}(N_{\text{procs}}).
\]

(7.5)

Simplifications arise from a knowledge of relationships.
7.4 DIGITAL COMPARISON

What is the difference between two system configurations \( C_1 \) and \( C_2 \)? This is a question that is frequently asked in connection with device management. Rather than testing whether a machine configuration is consistent with a policy (which normally involves only approximate or fuzzy classifiers), it is common to compare a system configuration \( C(x, t) \), at location \( x \) and time \( t \), to a reference system \( C_0 \) and characterize the difference in terms of the number of items that do not agree:

\[
\Delta C(x, t) = C_0 - C(x, t).
\]

(7.6)

One can question how useful it is to compare of dynamic system \( C(x, t) \) to a static snapshot \( C_0 \), but we shall not discuss that here. How shall we make such a comparison? What is the meaning of the difference symbol in eqn. (7.6)? Is it a numerical difference or a difference of discrete sets? The ability to compare configurations depends on the nature of the variables being compared: are they continuous or discrete? Although we shall later argue for the use of continuous variables in chapter 16, most comparisons are made digitally.

**Example 74.** Consider two configurations that are coded with symbols \( A, B, C, \ldots \):

\[
C_0 = \{A, D, F, G, \ldots\}
\]

\[
C(x_1, t) = \{B, D, F, A, \ldots\}.
\]

(7.7)

These configurations can be compared symbol by symbol.

To define a measure of distance between two such configurations, one can take a variety of approaches; there is no unique answer. Instead, we make a definition that suits our purpose. Two strings differ if their symbols do not match. We define a distance function (or metric) to define the distance between differing symbols. The most common distance function is the Hamming distance, which is a count of the number of bits that differ in the binary coding between two strings. A variation on this for higher level coding is the function for comparing symbol \( q \) with reference symbol \( q' \) is:

\[
d(q, q') = \begin{cases} 
a & q = q' \\ 
0 & q \neq q' \end{cases}
\]

(7.8)

This is a linear function, thus the distance grows additively for strings of many symbols. The distance between two configurations is thus:

\[
D(Q, Q') = \sum_{q \in Q, q' \in Q'} d(q, q').
\]

(7.9)

The straightforward comparison of strings is a naive way of comparing configurations that assumes only substitution errors in the symbol string. In general, we can have differences that are:
• Insertions.
• Deletions.
• Substitutions.

If one relates the differences or ‘errors’ between configurations to random processes, then one can only speak of the probability of an error. A knowledge of underlying mechanisms can allow the construction of a transition matrix (see section 5.4) that measures the likelihood of a transition from one state to another. This allows us to define a different kind of distance that is symbol dependent:

\[
d_{\text{sub}} = -\log \left( \frac{P(q \rightarrow q')}{P(q \rightarrow q)} \right) \\
d_{\text{del}} = -\log \left( \frac{P(q \rightarrow \emptyset)}{P(q \rightarrow q)} \right) \\
d_{\text{ins}} = -\log \left( \frac{P(\emptyset \rightarrow q)}{P(q \rightarrow q)} \right).
\]  

(7.10)

This is sometimes called the Levenshtein distance (see [BC95] and [OK98] for an intelligent discussion of pattern comparison with generalized differences).

**Applications and Further Study 7.**

• Describing any system or phenomenon quantitatively.
• Quantitative analysis allows us to study scalability of systems to changes in parameters.
• Comparing systems with different characteristics and determining the ‘distance’ or metric that measures this distance.
• Refining algebraic formulations of a problem in order to better understand its structure.
• Manipulating parameter choices in systems and exploring consequences.
• Pattern recognition in the layout of and input to systems for identification of system problems (anomaly detection).
CHAPTER 8

CHANGE IN SYSTEMS

Change is probably the most important quality of systems. A system that cannot express change cannot perform a function and is therefore trivial. We need a way to describe changes that occur in human computer systems; the natural language for this is mathematics, since it is both expressive and precise.

8.1 RENDITIONS OF CHANGE

Change can be represented in many forms. We choose the mode of description that is most convenient on a case by case basis. Some examples include:

EVENTS AND TIME

In a continuous system, time is an essentially mysterious dimension: a theatre in which things happen, without explanation. Essentially we have no idea why changes occur in systems. The information about the changes in information is not available in most systems of interest. We merely accept that change happens. In a discrete system, change defines time. This is because clocks are a part of the system, and if the clock doesn't change, then we cannot measure time[Bur14]. In the end, we simply accept the existence of ‘events’ in which states change their value.

In most system descriptions, there is someone outside the system observing it as an ‘open system’ from without, surrounded by so many changes that we need to separate out notion of time from what happens in the system. This allows us to separate change from a definition of external time. This is the usual approach to talking about human-computer systems. So, from now on, we shall assume that time exists independently of transactions and changes within a system.
Eventually, this assumption will get us into trouble, especially in defining the concept of relativity and consensual agreement in systems [Lam78, Lam01].

**Example 75.** Event driven systems, also called reactive systems, are driven by the arrival of events, usually network packet arrivals. Web servers are an example of this, in which web transactions are initiated by remote clients. The emergency services are also event driven. They remain in a largely dormant state until an event triggers them to life.

**Time series**

The time series is one of the simplest ways of representing change graphically (see [BJR94]). See, for example, figs 3.11 and 10.5. Time series are most easily plotted for data that are dense, or almost continuously varying, however streams of discrete symbols also form time series. Time series are especially useful in the continuum approximation to discrete systems, since they can be approximated by known functions, e.g. in Fourier analysis.

**Transitions, transactions and rollback**

Systems that change irregularly between well-defined states are often described in terms of transition tables. Finite state machines and Markov processes fall into this category. Transition systems are incredibly important in information theory and in physics. Any system can be described in terms of transition tables. Transition tables are essentially like the adjacency matrices of graphs. They tell us what the possible transitions to neighbouring states (nodes) are, given that we are already in a particular state (at a particular node of the graph). Transitions can be deterministic or non-deterministic (see section 8.2).

**Example 76 (Rollback).** The assumption that it is possible to reverse changes, or create generic ‘undo’ buttons in arbitrary software systems, remains a persistent myth amongst software developers, system designers, and system administrators in all areas of computing. The term ‘rollback’ is often used to describe this form of repair, usurped from its original usage in database transaction theory [RS87, WV01]. In current usage, rollback refers to the act of undoing what has been done; it is intimately related to checkpointing [LNP90, LNP91, PKD97, ABL+95, PT01], version control and release management. The impossibility of rollback was proven for open systems in [BC11].

**Invariance to change: stability in fixed points**

Changes in a system sometimes result in it converging towards a preferred state that is not easily altered again — such a state exhibits some kind of stability and this makes the system easy to

---

1The method of Green functions is the continuous generalization of transition matrix approach to continuous systems. See ref. [Bur02a].
predict; other times it results in chaotic, unpredictable behaviour that has no compact description. If a system wanders into a state from which it does not emerge without outside help, the state is said to be a fixed point of the system. This idea is interesting in management, since it represents a notion of stability. See also section 5.8.

8.2 DETERMINISM AND PREDICTABILITY

A deterministic process is one that is governed by rules that are always obeyed with complete certainty. For instance, it is generally assumed that the law of gravitation is a deterministic process: if one releases an object in a gravitational field, it falls (every time). In a deterministic system, if a system variable has value \( q(t') \) at some earlier time and value \( q(t) \) at some later time, then the probability of a transition from the value \( q(t') \), given that the earlier value was \( q(t') \) is unity:

\[
P(t, t') = P(q(t)|q(t')) = 1.
\]  

(8.1)

This certainty about the outcome of the process implies that we can use past knowledge to predict behaviour in the future. The system behaves in the same way under identical conditions, each time it is measured. In practice, this applies only to very simple systems, or systems that are isolated from external influences.

Non-determinism means that the probability of making a successful prediction about the system \( P < 1 \). The transition matrix for the process:

\[
P(q(t)|q(t')) < 1,
\]  

(8.2)

for any \( t, t' \). At each time-step there is a probability distribution indicating the likelihood of obtaining possible measured values. The distribution \( P(q) \) is the probability of value choice \( q \) at a given time. Since it is a probability distribution,

\[
\int dq P(q) = 1.
\]  

(8.3)

Non-determinism implies that we must make an educated guess about what is likely to happen in the future. This means there is uncertainty about what we can expect. The uncertainty is a direct result of what we do not know about the system. Even if there is an underlying deterministic system, it is of such complexity that we cannot realistically predict everything that will happen.

There are many ways in which randomness or unpredictability can enter into systems. One common assumption is that randomness follows the pattern of a Gaussian or ‘normal’ distribution:

\[
P(c) \propto e^{-(c-\mu)^2/2C}.
\]  

(8.4)

This would be equivalent to the assumption that the variable we were measuring had some ‘true value’ \( \mu \) that varied at random by about \( \pm \sqrt{C} \) in either direction. This model approximates some phenomena well, but not all.
Figure 8.1: The evolution of a function occurs as its value changes at each time step. In a deterministic process, the choice at each time step $dt$ is selected by a pre-determined function $U(t, t + dt)$, the result would always be the same, if one rolled back time and repeated the measurements; i.e. we can predict the future. In a non-deterministic system, the value of the function is picked at random, so that if one rolled back time and tried again, the outcome could be different.

Example 77. A transmitted network signal is a variable that has a ‘true’ or intended value, i.e. the signal that we are trying to send. This can pick up random noise along the way. Such noise is often well approximated by a Gaussian random error. Indeed, that is the assumption behind the Shannon formula for the capacity of a communications channel:

$$C = B \log \left( 1 + \frac{S}{N} \right)$$  \hspace{1cm} (8.5)

Example 78. The arrival of packets at a network switch is a random process, but there is no ‘correct’ or ‘true’ value to this number. There will be a probability distribution of values from different customers on different arms of the switch, but even this distribution might change slowly over time.

8.3 Oscillations and Fluctuations

Few systems are ever truly constant as parameters such as time are allowed to vary, but several systems exhibit change that averages out to nothing. Two examples of this that provide a good illustration of the difference between deterministic change and non-deterministic change are oscillations and random fluctuations. An oscillation is a periodic pattern of change that can be
expressed as a relatively simple combination of sine and cosine waves:

\[
q(t)_{\text{osc}} = \sum_{n} \sin(\omega_n t + \phi_n),
\]

for various constant circular frequencies \(\omega_n\) and phase shifts \(\phi_n\). The oscillation is deterministic because we have written down its exact functional form and can therefore predict its behaviour at any time with complete certainty.

**Example 79.** Consider a simplistic model of arriving traffic to a Web server over the course of a week. By measuring the actual arrival of requests at the server, we find a complicated pattern that must be described as a random variable since the requests are sent by a broad number of independent sources at unpredictable times (see fig 8.2). This behaviour is clearly complicated, but for the purpose of estimating system load one might try to approximate it by something simpler. We might try a test function of the form:

\[
f_1(t) = 2 \sin(t) + \cos(2t)
\]

See fig. 8.3. This function does not really resemble the actual random data, but it has some similar features. We see that there is a general decay in activity towards the end of the week, and we could
try to model this by adding an exponential decay term to our approximate model:

\[ f_2(t) = (2 \sin(t) + \cos(2t))e^{-t/4} \]  

(8.8)

The figure starts to take on some of the general features of the actual measurements, but it is still a long way off being a good approximation. On the other hand, the extreme simplicity of the function \( f_2(t) \) might outweigh its crude form.

Sometimes it is useful to model the average properties of a fluctuating function using a deterministic oscillation, as in the example above.

**Definition 35 (Fluctuation).** A fluctuation is a change in a random variable. Fluctuations \( \delta q(t) \) are sometimes measured relative to the mean value of a variable, i.e.

\[ \delta q(t) = q(t) - \langle q(t) \rangle. \]  

(8.9)

Random fluctuations are characterized by probability distributions \( P(q) \), i.e. the measured likelihood \( P(q) \) that the variable has value \( q \), or the cumulative distribution,

\[ F(Q) = P(q \leq Q). \]  

(8.10)

### 8.4 Rate of Change

Rates of change are important for modelling the dynamical interplay between competing processes, at large scales, so we shall need to be able to describe these changes. In physics, one nearly always assumes that we can use differential formulations of time dependence, because of the ubiquity of deterministic approximations and infinite scale resolution. For the kinds of random variables that...
we frequently meet in human-computer systems, there are no smoothly varying quantities in the raw data: the data fluctuate randomly. This makes the description of change more subtle.

In a continuum approximation, the description of rates is an easy matter: we have the derivative or gradient operator, whose effect is given by

$$\frac{\partial q(t, x_i)}{\partial t} \equiv \lim_{\Delta t \to 0} \frac{q(t + \Delta t, x_i) - q(t, x_i)}{\Delta t} \tag{8.11}$$

for the rate of change in time (speed). If there are other approximately continuous dependent labels $x_i$, or $i = 0, 1, 2, \ldots$, then there will also be partial derivative for these:

$$\frac{\partial q(t, x_i)}{\partial x_i} \equiv \lim_{\Delta x_i \to 0} \frac{q(t, x_i + \Delta x_i) - q(t, x_i)}{\Delta x_i} \tag{8.12}$$

For continuous functions, the limit $\lim_{\Delta t \to 0}$ is well defined, but for stochastic variables it is not. However, one can employ the continuum approximation as described in section 8.5 to approximate the local average behaviour by a smooth function for convenience, or simply use the definition above without the limit for a finite interval $\Delta t$.

The error incurred by assuming that these derivatives are actual smooth functions, i.e. $\Delta t \to 0$ and $\delta x_i \to 0$ over an interval of time $T$ is of the order $\Delta t/DeltaT$, from the continuum approximation.

Derivatives are used to find the extrema of a function, i.e. maxima, minima and inflection points, that satisfy:

$$\frac{\partial q}{\partial x_i} = 0, \tag{8.13}$$

(see fig. 8.5).
The second derivative or acceleration of the variable
\[ \frac{d^2 q(t)}{dt^2}, \frac{\partial^2 q(t,x_i)}{\partial x_i^2}, \] (8.14)
describes the curvature of the function, and is commonly used to determine the nature of turning points in a plot.

- At a minimum, the curvature is positive, i.e. the function is concave:
  \[ \frac{dq}{dt} = 0, \frac{d^2 q}{dt^2} > 0 \] (8.15)
- At a maximum, the curvature is negative, or the function is convex:
  \[ \frac{dq}{dt} = 0, \frac{d^2 q}{dt^2} < 0 \] (8.16)

Generalizations of these for several dimensions can be found in any book on analytical geometry.

We shall have frequent use for the idea of a saddle point in describing processes of competition (see fig. 8.5). A saddle point can be thought of as a region of a function in which one parameter is maximized while another is minimized, i.e. a saddle is both the top of a hill and the bottom of a valley. This structure occurs in ‘tug of war’ contests between different processes that share a common resource: one player is trying to maximize gains and the opposing player is trying to minimize the first player’s gains. This is a basic scenario in Game Theory (see chapter 19).

Figure 8.5: Turning points characterize the extrema of functions: (a) is a maximum, (b) is a minimum and (c) is a saddle point, or minimax.

In discrete change systems, like information systems, and transactional systems, changes are not described in terms of a time variable, but as sequences. Thus instead of writing \( X(t) \) as a smooth differentiable function, we would write a sequence: \( X_1, X_2, X_i, \ldots, X_N \), for events \( i = 1, 2, \ldots, N \). This is true in information analysis (see chapter 9), where we deal principally with Markoff processes.
CHAPTER 8. CHANGE IN SYSTEMS

8.5 APPLICATIONS OF THE CONTINUUM APPROXIMATION

In dealing with probabilities and statistical phenomena, we must distinguish between what is true over short times and what is true over long times. Defining this distinction is central to defining the average properties of systems, as experienced by users. One example is Quality of Service and what this means to different parties, at different scales. If there is a natural separation of time-scales then we can employ an approximation in which we consider the average system behaviour to be smooth and continuous, up to a limited resolution.

- \( \Delta t \): the interval at which we sample the system. The distribution of possible outcomes is approximately constant over such a small time scale. Even though each measurement contributes to defining the probability distribution of measured values, it would take many more measurements to change the distribution significantly.

- \( T \): the interval over which we can expect the distribution of values to change significantly. This is usually several orders of magnitude greater than the sampling time \( T \gg \delta t \).

Do we want quality of service at the level of seconds, minutes or days? This is an important issue. In systems that exhibit approximate stability (i.e. non-chaotic and non-self-similar systems), it is usually possible to separate the deterministic behaviour of system’s average behaviour, from the non-deterministic ‘fluctuation’ behaviour of microscopic details. Suppose that random requests arrive at intervals of the order \( \Delta t \) and that large scale variations in traffic levels occur over times of the order \( \Delta T \), where \( \Delta T \gg \delta t \) (see fig. 8.6).

Then, schematically, we may approximate the service rate function \( R(t) \), in terms of the average rate \( R_{av} \) for all time, as

\[
R(t) = R_{av} + R_{av} f(t) s(t)/2, \tag{8.17}
\]

where \( s(t) \) is a slowly varying function and \( f(t) \) is a rapidly varying function, both of maximum order 1:

\[
\max f - \min f \sim 1 \\
\max s - \min s \sim 1. \tag{8.18}
\]

The ‘fast’ fluctuation function \( f(t) \) modulates the average level about the mean value, so its average value over one of the larger time intervals \( T_n \) is zero:

\[
\int_{T_n}^{T_{n+1}} f(t) \, dt = 0. \tag{8.19}
\]

For example, \( f(t) = \sin(2\pi N \Delta t / \Delta T) \), for some positive integer \( N \). The slowly varying change in traffic, on the other hand, grows only slightly over the same interval:

\[
\int_{T_n}^{T_{n+1}} s(t) \, dt \equiv \Delta s \sim \frac{\Delta t}{\Delta T}. \tag{8.20}
\]
Figure 8.6: Short and long time scales represent actual and averaged variation in service rate. At the microscopic level, there is much noisy variation that averages out at over larger periods. Thus, the average service rate, over such a long interval is approximately constant (indeed, it tends to a constant as $\Delta T \to \infty$):

$$\langle R(t) \rangle_n = \frac{1}{\Delta T} \int_{T_n}^{T_{n+1}} R(t) \, dt = \bar{R} \left(1 + O\left(\frac{\Delta t}{\Delta T}\right)\right). \quad (8.21)$$

The uncertainty in rate is quantified by the range of values measured over all quality of service time intervals $\Delta T$:

$$U = \sqrt{\frac{\sum_n \langle R(t) \rangle_n - \langle R(t) \rangle_{\infty}}{N}^2}$$

$$= O \left( \frac{R_{av} \Delta t}{\Delta T} \right). \quad (8.22)$$

Clearly this approaches zero as $\Delta T \gg \Delta t$. This simply shows us that we can define any kind of service behaviour as stable and as fulfilling our ‘quality’ requirements, just by choosing a low enough time resolution. Thus, Quality of Service has no meaning unless we specify how large $\Delta T$ actually is.

A comment is in order for this reasoning. It has been observed that some service level patterns, such as Ethernet network traffic follow a self-similar pattern with very large variances over large time-scales (see [LTWW94, WP98]). This makes the idea of a quality time-scale very hard to implement in terms of averages, because the variances are so large that $\Delta T$ needs to be impractically large to get a clean separation of scales. See section 10.9 for further discussion on this.
CHAPTER 8. CHANGE IN SYSTEMS

8.6 UNCERTAINTY IN THE CONTINUUM APPROXIMATION

If we assume that a system changes smoothly and with infinite resolution, then we must also be realistic about when that assumption will fail to live up to reality. We must estimate the uncertainty in our estimate of the true value.

Example 80. Many Service Providers and companies that sell services claim guarantees on the level of service that they can provide, in spite of the inevitable occurrence of random events that limit the predictability of their assumptions. It is important to understand that service is about changes which occur in time, and thus time is an essential element of any service level agreement. If we focus on shorter and shorter intervals of time, it becomes impossible to guarantee what will happen. It is only over longer intervals that we can say, on average, what has been the level of service and what is likely to be the level of service in the future. We must therefore specify the time-scale on which we shall measure service levels.

Example 81. A Service Level Agreement for UUCP network connectivity could agree to transfer up to 10 MB of data per day. This is an easy goal, by modern standards, and it hardly seems worth including any margin for error in this. On the other hand, a Digital Subscription Line (DSL) network provider might offer a guaranteed rate of 350 Mbs (Mega-bits per second). This is a common level of service at the time of writing. But what are the margins for error now? If each customer has a private network telephone line, we might think that there is no uncertainty here, but this would be wrong. There might be noise on the line, causing a reduction in error-free transmission rate. When the signal reaches the Service Provider’s switching centre, customers are suddenly expected to share common resources, and this sharing must maintain the guarantees. Suddenly it becomes realistic to assess the margin for error in the figure 350 Mbps.

Example 82. A University Professor can agree to grade 10 examination papers per day. It is not clear that the level of interruptions and other duties will not make this goal unreasonable. The level of uncertainty is much higher than in a mechanistic network switch. We might estimate it to be 10 ± 3 exam papers per day. In this case, the Professor should include this margin for error in the contract of service.

Uncertainty is an important concern in discussing ‘Quality of Service’ (QoS); it is calculated using the ‘Theory of Errors’. Error theory arises from experimental sciences where one assumes, with some justification, that errors or uncertainties occur at random, with a Gaussian profile, about some true value. The Gaussian property basically ensures that errors are small or do not grow to an arbitrarily large size, compared to the rate of change of the average. However, whether or not a phenomenon really has a Gaussian profile or not, error handling techniques can be used to estimate uncertainties provided there is a suitable separation of time-scales. If there is not, the system must be regarded as unstable and therefore no guarantee can be made (see section 10.9). In section 3.9, the method of combining uncertainties is presented.
Example 83. Consider the rate of arrival of data $R$, in bytes, from the viewpoint of a network switch or router. The measurables are typically the packet size $P$ and the number of packets per second $r$. These are independent quantities, with independent uncertainties: packet sizes are distributed according to network protocol and traffic types, whereas packet rates are dictated by router/switch performance and queue lengths. The total rate is expressed as:

$$\lambda = rP.$$  \hspace{1cm} (8.23)

Using the method of combining independent uncertainties, we write:

$$\lambda = \langle \lambda \rangle + \Delta \lambda$$
$$r = \langle r \rangle + \Delta r$$
$$P = \langle P \rangle + \Delta P,$$

and

$$\Delta \lambda = \sqrt{\left(\frac{\partial \lambda}{\partial P}\right)^2 \Delta P^2 + \left(\frac{\partial \lambda}{\partial r}\right)^2 \Delta r^2}.$$ \hspace{1cm} (8.24)

Now, Asynchronous Transfer Mode (ATM) packets have a fixed size of 53 bytes, thus $\Delta P_{ATM} = 0$, but Ethernet or Frame Relay packets have varying sizes. An average uncertainty needs to be measured over time. Let us suppose that it might be 1kB, or something of that order of magnitude.

For a Service Provider, the uncertainty in $r$ also requires measurement; $r$ represents the aggregated traffic from multiple customers. A Service Provider could hope that the aggregation of traffic load from several customers would even out, allowing the capacity of a channel to be used evenly at all times. Alas, traffic in the same geographical regions tends to peak at the same times, not different times, so channels must be idle most of the time and inundated for brief periods. To find $r$ and $\Delta r$, we aggregate the separate sources into the total packet-rate:

$$r(t) = \sum_i r_i(t)$$ \hspace{1cm} (8.25)

The aggregated uncertainty in $r$ is the Pythagorean sum:

$$\Delta r = \sqrt{\sum_i \Delta r_i^2}$$ \hspace{1cm} (8.26)

The estimated uncertainty is

$$\Delta \lambda = \sqrt{r^2(\Delta P)^2 + \langle P \rangle^2(\Delta r)^2}$$ \hspace{1cm} (8.27)

Since $r$ and $\Delta r$ are likely to be of similar orders of magnitude for most customers, whereas $\Delta P < P$, this indicates that uncertainty is dominated by demand uncertainty, i.e.

$$\Delta \lambda \approx \langle P \rangle \Delta r.$$ \hspace{1cm} (8.28)

This uncertainty can now be used in queueing estimates.
8.7 CAUSATION IN SYSTEMS

Causation is the identification of a precondition that is necessary though not necessarily sufficient for an event to occur. In a deterministic system, we assume that there is always causation in every time step, by virtue of system design. If we push the button, the light turns on. Naturally, this assumes that all the parts of a system are in a working state. In the language of promises, we need all components to keep their promises in order for causation to propagate. This topic becomes increasingly subtle and thus we defer a detailed discussion until we have a satisfactory formalism to cope with it, in volume 2.

Probabilistic causation is a troublesome topic, because we assume that causation implies determinism. In probabilistic inference, there is a long standing discussion about the difference between causation and correlation of events. This is an issue that mixes semantics with the dynamical channels of influence, because inference methods necessarily coarse grain (or aggregate) data into lumps that obliterate a precise record of causation. One approach is to look for changes, and how a change at one location precedes a change at another location[Pea88, Pea00]. However, as we shall see in the next chapter, detecting change is also a matter for semantic interpretation, requiring a chosen scale of measurement, and the wilful aggregation of signals into intentionally chosen approximate categories that ultimately affect the definition of causal influence. This is the subject of information.

Applications and Further Study 8.

- Identifying mechanisms and character of changes in systems.
- Identifying pathways of causation (cause and effect).
- Estimating uncertainties and graininess inherent in changes.
- Relate cause and effect by a specific model.
- Quantifying the limitations of a description in terms of variables (approximation).
CHAPTER 9

INFORMATION AND INFLUENCE

An important concept in describing human-computer systems is the information encoded in changes that take place during their operation. This is one way of measuring the work that is carried out by the system. In the administration of systems, one needs to concept of information in several situations: to be able to match change with counter-change by sending the opposite information (configuration maintenance), as a principle of maximization or minimization for modelling randomness (disorder or predictability), and as a measure of the wastage in human-computer systems due to the transmission of uncontrolled information (noise).

The formulation of information in symbolic terms is important to system configuration and maintenance because, any problem that can be described in this way can be analyzed using the tools of stochastic error correction. Information is where the stochastic meets to deterministic.

9.1 WHAT IS INFORMATION?

The idea of information is rather subtle and is used in several different ways. The study of information began in the 1930’s and 1940’s by mathematicians such as Church, Turing, Von Neumann and Shannon. What one calls information theory today was largely worked out by Claude Shannon (of Bell Labs) and published in 1949 under the title *The Mathematical Theory Of Communication*. He defined the mathematical measure of information, called the entropy, and devised many of the core theorems used in information theory today.

Information theory is about the representation, interpretation and transmission of patterns of symbols (data). We attach meaning to patterns of change and call the result information. However, it is vital to distinguish meaning from the amount of information that represents it.

Example 84. *In Morse code, combinations of the digits ‘.’ and ‘–’ are used to ‘mean’ letters of*
Morse code uses strings of up to six dots and dashes to represent every single letter of the English alphabet. This seems to be rather inefficient, but there are only two symbols that can be communicated in Morse: it is a binary encoding. Clearly, the number of symbols, of a given alphabet, required to represent the same amount of information is important.

**Example 85.** *In the Unix operating system, the symbol CTRL-C means ‘interrupt program’, while the symbol # means ‘what follows is a comment and should be ignored’.*

Each of the symbols above conceals a whole series of actions that are carried out, as part of their interpretation. What is significant is that both meanings could be compressed into a single symbol, in the appropriate context. Context is very important in coding meaning in symbols, but symbols can be transmitted by the same rules, regardless of their interpretation; thus they form the basis of information.

### 9.2 Transmission

Patterns of data are mainly of interest when they are transmitted from a source to a receiver, over some channel of communication: e.g.,

- Text read from page to brain.
- Morse code sent by telegraph or by lantern.
- By memo or letter.
- As speech transmitted acoustically or by telephony.
- Copy data from hard-disk to memory.
- Copy data from memory to screen.

In each case, a pattern is transferred from one representation to another and perhaps retranslated at destination. Another way that data are transmitted is to copy data from one place to another. In computer administration, this is a way of making backups or of installing systems with software from a source repository. When data are copied, there is a chance that noise will cause errors to be injected, so that the copying is not performed with complete fidelity. The model of transmission again provides a model for discussing this. Data might be sent:

- From one place to another.
- From the past into the future, without moving.
In the first case, data are transmitted by copying, e.g. during a system backup. In the latter case, data do not move, but chance events (cosmic rays, moving magnets, accidental destruction) can that compromise the integrity of the data, e.g. data stored on a hard disk, or the programs and tasks that work in the system cause the data to evolve deterministically.

### 9.3 INFORMATION AND CONTROL

The ability to control a system implies an ability to stabilize it by countering change. In stochastic systems, such change is represented by the information content of the stochastic environment. To control such an environment requires a counter-input of information of comparable complexity. Information thus gauges the likelihood of one’s ability to control a system. If the informational entropy of the environment is much greater than the information content of the regulation scheme (e.g. the information content of policy rules), then regulation cannot be guaranteed. We might define the controllability of a system by the ratio of information input from the environment to the information contained in its control policy:

$$C = \frac{I_{\text{environment}}}{I_{\text{policy}}}.$$  \hfill (9.1)

### 9.4 CLASSIFICATION AND RESOLUTION

How shall we define information? We must contend with:

- Distinguishing and classifying patterns of symbols.
- Space-time coordinates and units. (Is a long beep the same as a short beep? Quantization/digitization)
- The meaning of redundancy and repetition.

Information arises from the abstract interpretation of changes in a medium.

To build a more precise picture of what information is, we begin with a signal $q(t, x, ..)$ which is a function or field which changes in time or space. We shall consider only time, as though we are receiving a signal from a stationary antenna, e.g. a radio signal. The signal is really a pattern formed by a disturbance in a physical medium. Our everyday experience leads us to believe that there are two types of signal $q(t)$:

- ‘Analogue’ or continuous functions $q(t)$.
- Digital or discontinuous functions $q(t) = \sum_i \theta(t - t_i)Q_i$. 

We shall see that this distinction is artificial but that, in this distinction, lies the central essence of what information is about. An analogue signal is a limiting case of a digital signal.

![Figure 9.1](image-url)

Figure 9.1: Coarse-graining or digitization is a coordinatization of the continuous signal.

In order to say anything about the signal, we have to map it out by placing it on a discrete grid of coordinates. At some arbitrary level, one decides not to subdivide space-time any further and one reaches a limit of resolution. This choice can result in a loss of fidelity in describing the pattern, if the pattern is denser than the coordinate grid (see figures). This is coordinatization is called digitization, or coarse graining of the signal (see figure 1). Such a process always takes place, even if one is not conscious of it. For instance, the eye automatically digitizes images for the brain since there is a finite number of cells on the retina.

Information must be defined relative to this set of coordinates since it is the only means we have of describing change. Let us begin by assuming that the detail in the signal is greater than the resolution of the grid. We do the following:

- Divide up the time axis into steps of equal width $\Delta t$. Here we shall look at an interval of time from $t = 0$ to $t = N\Delta t$, for some $N$.

- Divide up the $q$-axis into $C$ classes $Q_i = [Q_i^-, Q_i^+]$, which touch such that $Q_{i+1}^- = Q_i^+$. Digitization means that, whenever the function $q(t)$ is mostly inside a cell $Q_i$, its value is simply represented by the cell. We have compressed the detail in the square region $Q_i$ into a single representative value $i$, over an interval of time. There are good digitizations and bad digitizations (see figures 2,3,4). Nyquist’s sampling law tells us that the interval widths need to the half the width of the ‘finest change’ in the signal. In Fourier language, the sampling rate must be twice that of the greatest frequency we wish to resolve.
Figure 9.2: A poor digitization cannot sensibly determine the value of the signal within the cells.

Figure 9.3: A well-suited digitization without loss. This signal can be represented with just two classes, i.e. binary digitization.

Figure 9.4: The same signal, as in figure 3, this time digitized into 6 classes.
Example 86. CD players sample at 44.1 kHz and Digital Audio Tape (DAT) samples at 48kHz: the limit of human hearing is about 20 kHz when we are young, and falls off to about 12 kHz as we grow old.

In the physical world all information is digital if we examine it with sufficient resolution. Electrons are assumed to be indivisible, energy levels are really discrete. Even if we could devise a fully continuous representation, it would not be useful for carrying information because we would have to distinguish between an infinite number of different values, which would be noise.

The digits $Q_i$ are regarded as the basic units of information: they are a strict model for representing change. If $C = 2$, we have binary digits $\{Q_1, Q_2\} = \{0, 1\}$ etc, or bits.

9.5 Statistical Uncertainty and Entropy

In communicating commands and information, we associate meaning primarily to discrete events, or symbols. These might be words, or numbers or even pictograms and glyphs. In western languages, we have grown used to a fixed phonetic alphabet that is used to build up words. In the administration of human-computer systems, communication occurs by several methods:

Example 87. At the level of the computer, messages are passed to the CPU as instruction opcodes that are read as a stream of words from the memory of the computer. The size of each instruction is known to the processor, so that it always knows where the current instruction starts and ends. The symbol lengths are all the same size, since some instructions contain data and others do not.

Example 88. The English language alphabet consists of letters A, B, C, D, E, F, G, H, I, J, K, L, M, N, O, P, Q, R, S, T, U, V, W, X, Y, Z, together with a number of punctuation symbols: , ; . This is a basic set of symbols for forming messages in English. This is the alphabet used to convey information in written communication. In verbal communication, words can be broken down into phonemes or sound digits, though we known from computer speech that this is not always very realistic. Voice is a continuous (non-digital) process.

Example 89. In system administration, commands and communication between humans and computers are often conveyed in a specific terminology of code words, almost like a military code book. Special words are used: reboot, format, crash etc. These have special meanings and therefore act as single letters in a command alphabet. If we replaced them by their first letters: R, F, C, etc. the same meaning could be conveyed in the context of system administration\(^1\). Unix commands have a special alphabetic structure:

\[\text{command -option1 -option2 ...}\]

\(^1\)Indeed, technical workers have the uncanny habit of replacing all natural language with three letter abbreviations!
The set of commands is a finite set that can be labelled by a single digit for each command. Each option (−v, −w, etc.) is also a digital symbol that communicates information to the computer with standard interpretation. Windows commands are often conveyed through menu selections and button selections. In this case each button or menu item is a digit, with an interpreted meaning.

Example 90. The communication of simple configuration instructions by the Simple Network Management Protocol (SNMP) makes use of an alphabet of configuration codes with standard meanings, defined in Management Information Base (MIB) models for a device. A MIB defines an effective symbolic alphabet for sending streams of instructions with the modifiers: read, write, and trap.

In each of the actual examples above, we see communication – however complex its actual representation – reduced to the transmission of codes or symbols with special meanings. As long as the symbols can be distinguished, each symbol reduces the uncertainty or increases the information of the receiver about the sender. Such a stream of codes or symbols is the basis for nearly all instructions in a human-computer system.

We therefore need to recognize each code or symbol in a data stream. As we saw earlier, a signal \( q(t) \), digitized into symbols, or digits, by a digitization procedure (see fig. 9.5).

![Figure 9.5](image)

**Figure 9.5:** Coarse-graining or digitization is a labelling of the signal. First we divide up the signal variability into discrete symbols or finite sized blocks (first picture), and then we approximate the true signal to the coarse representation. Collapsing the digit-blocks into a histogram, by counting the number of occurrences (second picture) gives a graphical representation of the average numbers of digits in the signal. By scaling this histogram, one obtains a probability distribution for the signal. This is used to calculate the entropy, or average information content.

---

Even where speech is involved, or English text, we tend to reduce the number of possibilities to a few simple symbolic cases by inventing forms to fill out, boxes to tick, or special verbal jargon that reduces the amount of talking required to convey meaning.
is coded into the channel by the variation of the signal. This is digitized into a string of digital symbols (called a message) which occur in a certain order, with a particular frequency. When a digit occurs $n_i$ times out of a total of $N$ digits, where

$$\sum_i n_i = N,$$

one may say that the probability of the symbol’s occurrence was

$$p_i = \frac{n_i}{N}.$$  

(9.3)

The probability distribution of symbols in the signal is found by collapsing the image in fig. 3.3 into a histogram, as shown in the figure. Note that, it is assumed that only a single symbol can occur at any one time, i.e. that the channel is a serial channel.

This probability distribution displays the average amount of uncertainty in the symbols being transmitted. If the distribution shows that a message was concentrated entirely around a single symbol, then the uncertainty is very small. If it shows that all symbols were used in equal numbers, then the uncertainty is very large. To convey a lot of information in a message, we need to use different symbols, thus uncertainty in the average use of symbols is a measure of information content. This information is measured by the entropy:

$$H[p] = -\sum_{i=1}^{A} p_i \log_2 p_i,$$

(9.4)

where $A$ is the number of symbols in the alphabet. The entropy is measured in ‘bits’, or binary digits, if the logarithm is base 2. Note that the entropy is a scalar functional of the probability distribution.

If we define the expectation value of a function $g = \{g_i\}$, on this discrete space, by

$$\langle g \rangle_p = \sum_i p_i g_i,$$

(9.5)

then the entropy may also be written as the expectation value of the logarithm of the number of symbols per message length:

$$H[p] = \langle -\log_2 p \rangle.$$  

(9.6)

If one further assumes that the transmission of symbols is discrete in time, so that each symbol has the same duration in time, then this has the interpretation of an average rate of bits per second.

**Example 91.** If we have a value $x$, we need $\log_2(x)$ binary digits (bits) to represent and distinguish it from other values, e.g. 8 requires at least $\log_2(8) = 3$ bits, 32 requires at least $\log_2(32) = 5$ bits, and so on. If we use base 10 numerals, 8 requires at least $\log_{10}(8) = 0.9$ i.e. 1 symbol to
represent it, while 16 requires at least $\log_{10}(16) = 1.2$ i.e. 2 symbols to represent it. Thus the amount of information that needs to be distinguished to call a value $x$ includes the values of values less than this that must be coded separately, and the length of the symbol string measures the amount of information required to code the value in some kind of digits.

The entropy is a single scalar characterization of a probability distribution. Since it collapses all of the changes of the signal into a single number, it cannot distinguish equivalent signals, i.e. signals with the same frequency distributions of symbols in them. However, its interpretation as a transmission rate is very useful for considering the efficiency of communication.

Suppose we consider the transmission of information over a communications channel, then there is information both at the input of the channel and at the output. Under ideal conditions, the average information entering the input would be the same as the average information leaving the output; however, this is not necessarily the case if the channel is affected by noise. Noise is simply unpredictable information, or extra uncertainty, which enters the system from the environment and changes some of the symbols.

Fig. 9.6 shows how the distribution of symbols in a message can be altered by its transmission over a channel. The entropy may be used to characterize the average information content at the

Figure 9.6: Information passing through a noisy channel. The frequency distributions of symbols can be altered by noise introduced by the environment, as the data pass through the channel. Ideally, these two distributions would be equal. In the worst case, they would have nothing in common.
input $I$ and at the output $O$.

$$H[I] = -\sum_{i=1}^{A} p_i(I) \log_2 p_i(I),$$
$$H[O] = -\sum_{i=1}^{A} p_i(O) \log_2 p_i(O),$$

(9.7)

where $p_i(I)$ are the probabilities at the input, and $p_i(O)$ are the probabilities at the output. Note that $I$ and $O$ label sets of symbols, whose elements are the alphabet of digits.\(^3\) If transmission of information is perfect, then,

$$H[I] = H[O].$$

(9.8)

This can only occur, if the information extracted from the output is predictable using the information at the input, i.e. if they are correlated. If $p(I)$ and $p(O)$ are independent events, this cannot be the case, since output and input would be unrelated.

### 9.6 Properties of the Entropy

The entropy $H$ has the following properties:

- It is continuous in the $p_i$.
- When the $p_i$ are equal to $1/C$, it is a monotonically increasing function of $C$, meaning that information is proportional to resolution.
- If the signal is completely ordered, i.e. $p_j = 1$, and $p_{i \neq j} = 0$, then the entropy has a minimum value of zero. This tells us that a trivial signal which never changes carries no information.
- If the signal is maximally randomized so that all the $p_i$ are equal to $1/C$, then the entropy takes on its maximum value $\log_m C$. This tells us that a lot of information is needed to characterize a signal with a lot of change.
- It is independent of the route by which the final probabilities were achieved. (Shannon III)

\(^3\)In this picture, the transmission must be a congruent mapping of symbols: the same symbol set should be used at the input and output of the channel. Usually the input and output alphabets are assumed to be the same; however, if the channel is encrypted or otherwise encoded, this need not be the case. For now, we shall assume that the number of symbols is $A$ in both cases, and that the symbols map congruently, in the same sequence.
CHAPTER 9. INFORMATION AND INFLUENCE

Information is about the length of the shortest message one could send that exactly describes a string of data to someone else, i.e. so that they could reproduce it with complete accuracy.

It is important not to confuse information with regularity of a signal, or orderliness. We might think that a maximally random, noisy system (like a the fuzzy dots on a television screen with no signal) has no information worth speaking of, whereas a system which is very ordered does convey information. This is not true; in fact, the noisy television screen contains so much information that extracting meaning from it is difficult. This is a human cognitive problem.

9.7 UNCERTAINTY IN COMMUNICATION

To discuss how information is transmitted, we must broaden the discussion to allow for unreliability in copying data. For this, we introduce the joint probability matrix, which describes the probability that sending digit $Q_i$ from $A$ actually results in symbol $Q_j$ being received by $B$, for any $i, j$.

Suppose there are two sets denoted $A$ and $B$, there is a joint probability matrix

$$p_{ij}(A, B) = p_{ij}(A \cap B), \tag{9.9}$$

which specifies the probability that the $i$-th event will be measured in $A$ in conjunction with the $j$-th event is measured in $B$. In the case where $A$ and $B$ are mutually independent events, i.e. the dual event occurs entirely by coincidence,

$$p_{ij}(A, B) = p_i(A)p_j(B) = p_{ji}(B, A). \tag{9.10}$$

In this case, the probabilities factorize and the combined probability is the overlap of the two sets, i.e. the product of the probabilities of the individual events, i.e. the two sets have to have some common elements by coincidence (see fig. 9.7). Such a factorization is not possible if $A$ and $B$ are more deeply related. In general, one can only say that

$$p_i(A) = \sum_j p_{ij}(A, B) \tag{9.11}$$

$$p_j(B) = \sum_i p_{ij}(A, B). \tag{9.12}$$

These are called the marginal distributions, and are formed by summing over all other indices than the one of interest. It is natural to apply joint probabilities to the input and output of a communications channel. Since we are interested in communication, the worst case scenario is when the probability of an event at the output occurs completely independently of an event being sent at the input. In that case, the output is purely ‘noise’, and the only chance of seeing the same data at input and output, is when there is a coincidental overlap

$$p_{ij}(I, O)_{\text{worst-case}} = p_i(I)p_j(O). \tag{9.13}$$
The ideal case is when a transmission is reproduced with perfect fidelity; i.e. $p_{ij}(I, O)$ is a diagonal matrix

$$p_{ii}(I, O)_{\text{best-case}} = p_i(I) = p_i(O),$$

(9.14)
i.e. the probability distributions at input and output are identical.

Using the joint probability of events at the input and output, of a communications channel, we can construct a joint entropy,

$$H(I, O) = - \sum_{i,j=1}^{A} p_{ij}(I, O) \log_2 p_{ij}(I, O),$$

(9.15)
which measures the information in the whole channel, between input and output. As an expectation value, it may be written

$$H(I, O) = \langle - \log_2 p \rangle_{p(I, O)},$$

(9.16)
where the probability set which performs the weighting is specified for clarity. If the input and output are independent (worst case), then one has

$$H(I, O) = - \sum_{i,j=1}^{A} p_i(I)p_j(O) \log_2 p_i(I)p_j(O),$$

$$= - \sum_{i} p_i \log_2 p_i - \sum_{j} p_j \log_2 p_j$$

$$= H(I) + H(O).$$

(9.17)

In general,

$$H(I, O) \leq H(I) + H(O),$$

(9.18)
i.e. the uncertainty (or information) in the joint system is less than that of the two ends combined, because some of the information overlaps, or is common to both input and output.
CONCEPTUAL ENTROPY AND INFORMATION FLOW

The joint entropy measures a kind of correlation of events, but it is not a measure of communication, since it does not specify the causal direction of the transmission. We need a way of saying that a certain symbol arrived at the output, because it was sent into the input. For this, we define the conditional probability of measuring $A$, given that we are certain of $B$:

$$p_{ij}(I|O) \equiv \frac{p_{ij}(I,O)}{p_j(O)} = \frac{p_{ij}(I \cap O)}{p_j(O)} = \frac{n_{ij}(I \cap O)/N}{n_j(O)/N}, \quad (n_{ij} \in (I \cap O)) \tag{9.19}$$

This measures the likelihood that the $i$-th symbol was presented to the input, given that the $j$-th symbol was measured at the output. In the case where the input and output are completely independent, the numerator factorizes, and $p(I|O) \rightarrow p(I)$, i.e. the knowledge of the output makes no difference.

Another way of interpreting the set $p_{ij}(I|O_j)$, for all $i, j$, is to ask: how many different input distributions give rise to a given output distribution? The sketch in fig. 9.8 shows the regions of the symbol spaces which give support to this quantity. One may also write,

$$p_i(A) = \sum_j p_{ij}(A|B)P_j(B) = \frac{n(A)}{(n(A) + n(B))}. \tag{9.20}$$

The conditional probability represents the overlap region, scaled by the space of possible outputs.

Figure 9.8: The receiver’s viewpoint: the conditional probability of input, given that the output is known. This is the space of possible overlaps, given that the space of total possibility is now restricted to those that belong to known outputs. The normalizing factor in the denominator is thus reduced, and the conditional probability is greater for the knowledge of the output.
The conditional entropy is defined by

\[ H(I|O) = - \sum_{i,j} p_{ij}(I,O) \log_2 p_{ij}(I|O) \]

\[ = -\langle \log_2 p(I|O) \rangle_{I\cap O} \]  

(9.21)

It is a scalar value, measuring the average variability (information) in \( I \), given that the input overlaps with the output, i.e. given that there is a causal connection between input and output. The meaning is slightly non-intuitive. Although the conditional probability is restricted to the output region, on the right hand side, the averaging is taken over the joint probability space (see fig. refvenn3) and thus receives support from the whole region. This measures the average uncertainty that remains about the input, given that the output is known. That uncertainty exists because of the possibility of noise en route.

We are assuming (hoping) that the probabilities \( p(I) \) and \( p(O) \) are not going to be independent, since that would be an uninteresting system, so the joint probability is not merely the overlap region, but the whole space of inputs and outputs. The region which now overlaps with the specific output distribution is only one point, but there is still uncertainty in the input.

The conditional entropy is a measure of information transmitted, because it contains the noise which can be picked up along the channel. To filter out the noise, we consider one final quantity: the relative entropy.

**Relative entropy and mutual information**

The relative entropy is a measure of the information which is common to two possibility spaces. Consider two spaces \( P \) and \( Q \),

\[ H(P/Q) \equiv + \sum_P P \log \frac{P}{Q}. \]  

(9.22)
This is a measure of the distance between \( P \) and \( Q \). Although it is not a true metric, it has the property of being zero when \( P \) and \( Q \) are identical. If we apply this to the input and output of the communications channel, the result is called the mutual information, or common entropy.

\[
H(I;O) = H(I/O) = \sum_{ij} p_{ij}(I,O) \log_2 \frac{p_{ij}(I,O)}{p_i(I)p_j(O)}.
\]  
\text{(9.23)}

The semi-colon is used to indicate that this quantity is a symmetrical function of the input and output. It compares two situations: a source and a receiver that are completely independent (\( Q \)), and a general source and receiver that are connected by a partially reliable communications channel. It measures the average reduction in uncertainty about the input that results from learning the value that emerges from the output. In other words, it represents the what passes along the channel between input and output.

We can express the mutual information in a number of different ways. Noting that:

\[
p(I,O) = p(O)p(I|O) = p(I)p(O|I),
\]  
\text{(9.24)}

we have:

\[
H(I;O) = \sum_{ij} p(I,O) \log_2 \frac{p(I|O)}{p(I)}
\]
\[
= H(I) - H(I|O).
\]  
\text{(9.25)}

This has the form of the information (uncertainty plus signal) at the input minus the uncertainty in the input, given a definite output. In other words, it is the part of the input which was transmitted to the output, or the likelihood of transmission (the fidelity). Another representation is:

\[
H(I;O) = \sum_{ij} p(I,O) \log_2 \frac{p(O|I)}{p(O)}
\]
\[
= H(O) - H(O|I).
\]  
\text{(9.26)}

This is the information arriving at the output (signal plus noise), minus the uncertainty information at the output, given a fixed input (i.e. the noise picked up along the way). Again, this is the likelihood of correct transmission. A third form can be found from

\[
H(I,O) = H(I|O) + H(O).
\]  
\text{(9.27)}

Using this in eqn. (9.25), we obtain,

\[
H(O; I) = H(I) - (H(I,O) - H(O))
\]
\[
= H(I) + H(O) - H(I,O).
\]  
\text{(9.28)}

This is the sum of information entering the input and leaving the output, minus the total amount of independent information in the system. What is left, must be the information in the overlap region (see fig 9.10), i.e. the information which is common to both input and output.
Figure 9.10: The entropies visualized as measures on the space of all possible input and output signals. The Venn diagram shows where the mutual information received support from the input-output probabilities; it does not strictly represent the magnitude of the entropy.

9.8 A GEOMETRICAL INTERPRETATION OF INFORMATION

The idea of classification, or quantization leads to the idea of states $Q_i$. We say that the signal, is in state $Q_i$ at time $t$. These states are linearly independent: matter how many times we add together $Q_0$, we will never get $Q_1$ or $Q_2$... The effect of a message is thus to draw a vector in this space (fig. 9.11):

$$I = \sum_{i=1}^{C} n_i \hat{e}_i = n_1 \hat{e}_1 + n_2 \hat{e}_2 + ... n_C \hat{e}_C.$$  

(9.29)

This vector summarizes a new kind of state: an average state. It does not preserve the details of the path which was taken in order to reach the point. In fact there is a large number of equivalent paths, of the same length, to the same point.

_The number of equivalent paths is a measure of how much effort one must expend to describe the actual path, i.e. it is a measure of the information represented by the path._

$I$ summarizes the number of units of information in the signal, since each unit vector is a character or digit in the message. The metric distance of the such a vector

$$|I|^2 = I^i \delta_{ij} I^j = I \cdot I$$  

(9.30)

is a measure of the number of digits, but that is not a good measure of the information. The difference of two message vectors is a measure of how many digits were different (without respect to ordering). This is related to a quantity called the Hamming distance, which represents the number of symbols which differ between two messages. We shall return to this when considering how messages are coded.
Figure 9.11: A binary message may be drawn as a path in a two dimensional lattice. A 26 dimensional lattice would be needed to represent a word in English.

As digits are received and plotted in the lattice, we build up an average description of the message. If the probability of receiving digit $i$ is $p_i$, then clearly, after $N$ time units, we have received $N$ digits, and:

$$i = \frac{1}{N} = \sum_{i=1}^{C} \frac{n_i}{N} \hat{e}_i = \sum_{i=1}^{C} p_i \hat{e}_i.$$  \hspace{1cm} (9.31)

$p_i$ is the amount of time the signal spends within class $Q_i$. Suppose we reduced the number of classes of $Q_i$ to one: then there would only be a single path (ordering irrelevant), and only one kind of digit would be possible. Only the number of digits (length) can then convey information. If we had an infinite number of classes or dimensions, then there would be an infinite number of equivalent paths and the amount of detail would be infinite. No two signals would ever be the same in practice, and the signal would be indistinguishable from noise.

Let us call the number of equivalent paths between two points in this lattice $h$, (the ‘hopelessness’ or uncertainty of finding the right path, i.e. the correct ordering of digits, given only the final
state). It is possible to find a formula for this number:

\[
    h(N) = \frac{(\sum_{j=1}^{C} I_j)!}{\prod_{k=1}^{C} (I_k!)}. \tag{9.32}
\]

This grows very rapidly with the Euclidean distance \(|\vec{d}|\) in this message lattice.

\[
    |\vec{d}| \equiv d = \sqrt{\sum_{i=1}^{C} (I_i)^2}. \tag{9.33}
\]

If we consider messages over a fixed number of time intervals \(N\Delta t\), then we can express \(h\) as:

\[
    h(N) = \frac{N!}{(N p_1)! (N p_2)! \ldots (N p_C)!}. \tag{9.34}
\]

This is a large and awkward number to deal with, so we look at its logarithm \(H\). This represents the length of a string in base \(m\) digits that would be able to code the number. This is an important measure, because it is the smallest number of digits that one needs to be able to label the exact path, and therefore distinguish it from all the other alternative paths:

\[
    H_N = \log_m h. \tag{9.35}
\]

Moreover, by assuming that \(N\) is large (high resolution), then we can Stirling’s approximation to write this in a simpler form, without awkward factorials. Using:

\[
    \sum_i \log_m (N p_i)! \simeq N \log_m N - N
\]

we get

\[
    H_N = -N \sum_i p_i \log_m p_i \equiv N \langle i \rangle. \tag{9.37}
\]

To get the fractional uncertainty per digit, we divide by \(N\). Shannon called this quantity the informational entropy, or average information. He did not derive it in this way, but on general grounds using similar combinatorial properties. It represents the average information needed to distinguish the exact message from a random message with the same probability distribution. It is measured in units of \(m\)-ary digits:

\[
    H = H_N / N = -\sum_i p_i \log_m p_i. \tag{9.38}
\]
This geometrical interpretation is very helpful in interpreting the entropy. It can be extended to the conditional entropies also. Imagine that the picture in fig. 9.11 leads to a slightly different picture at the receiver, due to errors in transmission. Over time, there is an ensemble of similar pictures that might be transmitted with various probabilities. There is thus an additional uncertainty between \( I \) and \( O \) that is not caused by the information in the message, but by unreliability of the channel. A person receiving a message knows what they have received from the sender, so they are not completely ignorant. That knowledge reduces their uncertainty about the message from a random jumble of digits to one of the possible causes, given our knowledge of the probable distortion by the channel:

\[
H(I, O) \rightarrow H(I|O) = H(I, O) - H(O).
\] (9.39)

The conditional entropy picks out the ensemble of possible causes, i.e. the alternative messages that could reasonably have given rise to the message received. From eqn. (9.39) we see that this is the maximum possible uncertainty in the channel \( H(I, O) \) minus the uncertainty that distinguishes a specific message from a random jumble of symbols at the receiver \( H(O) \). The number of possible causes for the message that arrives is therefore of the order

\[
E \sim m^{H(I|O)},
\] (9.40)

where \( m \) is the base of the logarithm in the entropy \( (m = 2 \) for binary digits). We should never forget that the entropy makes statements about average properties, not about specific instances.

### 9.9 Compressibility and size of information

Defining the expectation value of a vector \( L_i \) by

\[
\langle L \rangle = \sum_{i=1}^{C} p_i L_i = \text{Tr}(pL),
\] (9.41)

it is possible to view the entropy as the expected uncertainty per digit in the message:

\[
H = \langle - \log_m p \rangle = \langle \log_m p^{-1} \rangle = \langle \text{probable digits} \rangle.
\] (9.42)

**Example 92.** Consider a message variable which has 16 possible measurable values or states that occur with equal probability. In order to label (and therefore describe) these outcomes, we need 4-bit strings, since \( 2^4 = 16 \). The information/entropy is

\[
H = - \sum_{i=1}^{16} \frac{1}{16} \log_m \frac{1}{16} = \log_m 16.
\] (9.43)
If we take binary digits \( m = 2 \) (alphabet length 2), we get \( H = 4 \), showing that the average information per message needed to communicate the behaviour of the random variable over time is 4 − bits, as we assumed. This shows that, if we choose log-base 2, the answer comes out in bits. This is the uncertainty per digit in the message, since all digits occur with equal probability.

Suppose we encoded the information on a string of DNA (alphabet length 4), then the result would be \( m = 4 \), \( H = 2 \), or 2 DNA characters (A,C,T,G). \( m \) is the alphabet size of the message.

**Example 93.** Consider a human-computer monitoring system in which the probabilities of eight different faults, \( \Delta_i \) \((i = 1 \ldots 8)\), are found over time to occur with the following probabilities:

\[
P = \left( \frac{1}{2}, \frac{1}{4}, \frac{1}{8}, \frac{1}{16}, \frac{1}{64}, \frac{1}{64}, \frac{1}{64}, \frac{1}{64} \right).
\]

Notice the degeneracy of the last four values. The entropy or information of this distribution is

\[
H = -\frac{1}{2} \log_2 \frac{1}{2} - 4 \frac{1}{64} \log_2 \frac{1}{64} = 2 \text{ bits}.
\]

If we want to communicate a message that has this distribution of probabilities on average (i.e. we are looking for long-term average efficiency) to an operator, it looks as though we’ll need 3-bits in order to label the outcomes of the eight values. However, we do not need to use the same number of bits to code each of the values. It makes sense to code the most probable fault symbol using the smallest number of bits. Suppose we use the bit strings

\[
\begin{align*}
\Delta_1 &= 0, \\
\Delta_2 &= 10, \\
\Delta_3 &= 110, \\
\Delta_4 &= 1110, \\
\Delta_5 &= 111100, \\
\Delta_6 &= 111101, \\
\Delta_7 &= 111110, \\
\Delta_8 &= 111111,
\end{align*}
\]

With this coding, we have distinguished both the distribution of the digits and their positions with a minimum amount of information. The lengths of these strings are \( L = (1, 2, 3, 4, 6, 6, 6, 6) \), and the expectation value of the length, given the probabilities is

\[
\langle L \rangle = \sum_{i=1}^{8} p_i L_i = 2 \text{ bits}.
\]

This illustrates Shannon’s theorem that the entropy is a lower bound on the average length to which any digit of a message can be compressed. In other words, if wait until we have received \( N \) such fault reports, each coded according to the scheme above, then the total amount of data will be approximately \( 2N \) bits long.
9.10 INFORMATION AND STATE

Informational entropy tells us about the statistical distribution of digits produced by a system. It measures the average state of the system over an ensemble of measurements, i.e. it is a cumulative result. The details of the current state are mixed up with all other measurements so that specific information about present state is lost.

One of the surprises of information theory is that one does not usually need to know the precise state of a system in order to be able to describe the system’s properties over time\(^4\). This is an important point, because it means that we do not need to keep infinitely many records or logs of what happened to a system in order to understand its history: it is possible to compress that information into its statistical essence.

If we imagine that a string of symbols, with a certain entropy, is generated by a finite state machine, or other computer program that remembers state, then the entropy also tells us about the average state of the state machine. We cannot tell what state the system is in by looking at the entropy; however, we can tell, at least in principle, that a change of state has occurred within the time resolution of our measurements.

Consider the histogram in fig. 9.12, representing a system with three states: active, waiting and terminated. If a command symbol is transmitted, causing this system to change state to ‘active’, then the column for the probability for ‘active’ gets relatively taller and the others get a little shorter. This changes the entropy. If, however, on average we receive equally many commands for each state, then distribution does not change, and we say that the system is in a steady state.

\(^4\)This has been one of the most difficult ideas to accept for authors trying to model the system administration process.
**Definition 36** (Steady state). A system is said to be in a steady state if the entropy of the transitions in the system is constant over statistically significant times. Usually the only stable steady state is one of maximal entropy.

If we want to change the average state of a system, we need to send it a persistent command signal, consisting of many symbols of the same type (or subset of the whole). This will tend to raise the level of one or more of the histogram columns at the expense of the rest, and ‘sharpen’ the distribution. In the language of uncertainty, a persistent signal reduces our uncertainty about the state of the system, since it forces the signal to be in a known state more often.

This result, although seemingly innocuous, is of the utmost importance to system administration; let us express it in three different ways:

- A stable average state requires no external information to be input to maintain it.
- A random sequence of commands, with maximal entropy, causes no change in the statistical state of a system over long times.
- A sustained system reconfiguration requires the input of statistically significant, low entropy information over long times.

Here ‘long times’ means long enough to be statistically significant in relation to any noise or residual randomness. Referring to fig. (8.6), we require a signal of the order of time $\Delta T$ to make a significant impact. We shall return to the issue of configuration changes in chapters 15 and 16.

### 9.11 Maximum Entropy Principle

The concept of entropy characterizes the uncertainty or ‘bluntness’ of statistical distributions. If we consider the statistical distributions in fig. 3.4 and in fig. 3.5, then the complete certainty of fig. 3.4 characterizes an absolute minimum of entropy, whereas the completely uncertain fig. 3.5 represents maximal entropy.

If we plot the entropy as a function of the probability of two events, it has the form of fig. 9.13. This shows the shape of the entropy function. More general distributions are multi-dimensional generalizations of this. They key point is that there is a maximum value for the entropy that is indicated by a turning point, in the middle of the graph. There is also a minimum value at the symmetrical points $p = 0$ and $p = 1$, but there is no turning point here. This means that, if we try to maximize the entropy by looking for the stationary points of this curve, the result will only find the maximum value, not the minimum (which is always zero, from the form of the function).
Let us work out the maximum entropy distribution for a histogram with $C$ classes. We do this using the method of Lagrange. Let the Lagrangian be
\[
L = -\sum_{i=1}^{C} p_i \ln p_i - \alpha \left( \sum_{i=1}^{C} p_i - 1 \right), \tag{9.48}
\]
where $\alpha$ is a Lagrange multiplier, or parameter that enforces the constraint that the sum of probabilities is 1, and we use the natural logarithm for convenience.
\[
\sum_{i=1}^{C} p_i = 1. \tag{9.49}
\]
Maximizing this with respect to all parameters gives:
\[
\frac{\partial L}{\partial p_i} = -\ln p_i - 1 - \alpha = 0
\]
\[
\frac{\partial L}{\partial \alpha} = \sum_{i=1}^{C} p_i - 1 = 0. \tag{9.50}
\]
This is solved for $p_i$ by

$$p_i = e^{\alpha - 1}$$

$$\sum_i p_i = Ce^{\alpha - 1} = 1$$

i.e. $p_i = \frac{1}{C}$. \hfill (9.51)

In other words, the completely flat distribution is the case of maximum entropy, in which all probabilities are the same.

If we maximize the uncertainty about a system, entropy leads us to the distribution of values that contains the least planning (the fewest assumptions), or the most randomness. This is not particularly interesting until we apply additional constraints that tend to make the entropy less than this value. This turns out to be a powerful tool.

**Definition 37** (Maximum entropy distribution). A *maximum entropy distribution* is that produced by a maximally random variable that is constrained by a function $\chi(p) = 0$. It is found by maximizing the Lagrangian function

$$L = -\sum_{i=1}^{C} p_i \ln p_i - \alpha \left( \sum_{i=1}^{C} p_i - 1 \right) - \beta \chi(p),$$

and solving for $p_i$. \hfill (9.52)

**Example 94.** Find the least clustered (most distributed or robust) network, as a function of node-degree, given that we have a fixed number of links $L$ to join the nodes together; i.e. where do we place $L$ cables between a number of cities in order to have the best distribution of resources.

We begin by defining the probability of finding a node of degree $k$,

$$p_k = \frac{n_k}{N} = \frac{\text{Nodes with degree } k}{\text{Total number of nodes}}.$$ \hfill (9.53)

To count the number of links in the graph, as a function of $k$, we note that every node of degree $k$ has $k$ links attached to it, but only half the link is attached since the other half is attached to another node. Thus the number of links is related to $k$ and $n_k$ by

$$L = \sum_k \frac{1}{2} k \times n_k.$$ \hfill (9.54)

Our constraint is thus

$$\chi(p) = \sum_k \frac{1}{2} k \times n_k - L = \sum_k \frac{1}{2} N p_k - L = 0.$$ \hfill (9.55)
The Lagrangian is therefore

\[ L = - \sum_{i=1}^{C} p_i \ln p_i - \alpha \left( \sum_{i=1}^{C} p_i - 1 \right) - \beta \left( \sum_{k} \frac{1}{2} Nkp_k - L \right), \quad (9.56) \]

Maximizing this function gives:

\[ \frac{\partial L}{\partial p_k} = - \ln p_k - 1 - \alpha - \beta \sum_{k=1}^{1} \frac{1}{2} Nk = 0 \]

\[ \frac{\partial L}{\partial \alpha} = \sum_{k=1}^{p_k - 1} = 0 \]

\[ \frac{\partial L}{\partial \beta} = \sum_{k} \frac{1}{2} Nkp_k - L = 0. \quad (9.57) \]

This has the solution, from the first two lines in eqn. (9.57)

\[ p_k = e^{-\frac{1}{2} N\beta k} \sum_{k} e^{-\frac{1}{2} N\beta k}. \quad (9.58) \]

It is an exponential distribution, i.e. nodes of large degree are exponentially suppressed (\( \beta > 0 \) else the distribution is not normalizable for arbitrarily large \( k \)) relative to nodes with small \( k \). To express this in terms of the constant \( L \) we can perform a final mathematical trick. Suppose we define the generating function

\[ Z = \sum_{k} e^{-\frac{1}{2} N\beta k}, \quad (9.59) \]

then we can express

\[ L = - \frac{\partial}{\partial \beta} \ln Z. \quad (9.60) \]

Thus \( \ln Z = -L\beta \). The value of \( \ln Z \) cannot be evaluated exactly, but it is constant, and we can call it \( \ln \equiv -\zeta \) for convenience, so that \( \beta = \zeta / L \), thus we have:

\[ p_k = e^{-\frac{1}{2} N\zeta k/L} \sum_{k} e^{-\frac{1}{2} N\zeta k/L}. \quad (9.61) \]

If we express \( L \) as a fraction of \( N \), this can be simplified even further. This probability distribution is called the Boltzmann distribution after the physicist L. Boltzmann who discovered its importance in physics. It tells us that most nodes should have small values of \( k > 0 \), and exponentially fewer large nodes should exist, for maximum entropy or least clustering. Clearly this makes sense – the fewer large links, the less clustering there will be. The question is why should be any large degree nodes? The reason is clearly that we have fixed the number of links and the number of
nodes to be constants, and there are limits to how many links we can fit into a small number of
nodes – some of them will have to be of higher degree if \( L > N \). If \( N/L \) is small, this exponential
falls off only slowly and there will be larger numbers of higher degree nodes. Thus, this is the least
clustered network we can build.

Maximum entropy distributions occur in many situations where one wishes to make the least
possible assumption, or invoke the greatest range of possibility in a system.

9.12 Fluctuation Spectra

One application of maximal entropy distributions is in the characterization of fluctuating random
behaviour in systems.

**Definition 38** (Fluctuation spectrum). A fluctuation spectrum is a probability distribution of values
that can be assumed by a random variable. i.e. if \( P(q) \) is the fluctuation spectrum of the variable,
then the average value over all times is:

\[
\langle q \rangle = \int dq P(q)q. \tag{9.62}
\]

The maximum entropy principle allows us to characterize the likely signal behaviour of random
events, such as data requests arriving at a server, given the known constraints under which the
system can behave. We do this by assuming maximal entropy for \( P(q) \), subject to any boundary
conditions.

**Example 95.** In ref. [Bur00a], this method was used to model the observed fluctuation spectra of
network services, such as the World Wide Web, using a periodic model for requests based on the
working week (see fig. 2.1). The resulting distribution was a Planck exponential distribution (see
fig. 9.14). There are many maximum entropy distributions — as many as there are constraints to
apply — and they can often be related to simpler forms.

9.13 Propagation of Influence

When a change in one part of a system necessarily precedes a change in another, we use the term
causation. Our understanding of causation is intrinsically linked to approximation into symbol
categories, as information. If we aggregate all changes into a binary signal ‘something happened’,
then causation becomes increasingly vague. If we trace very precise channels of information from
atomic and isolated parts, we can pinpoint channels of causation with much greater plausibility.
The dependence of causation on approximation and aggregation means that it is scale dependent. If
we look at a coarse scale, it might be impossible to distinguish the order of prior events from final evidence. This is a fault of the methodology, not a proof that causation is non-existent.

If we can reduce a system to a network of low level atomic parts, then each point to point interaction may lead to transmission of influence, and thus propagation of causality. However, even this might not be quite what we expect. Isolation of neat linear stories is not possible in general, especially in non-linear, strongly coupled systems. Thus causation can become circular.

**Example 96.** In computer monitoring, we are usually forced to obtain information from kernel resource metrics. When fault finding, we look for changes in these measured values as evidence of causation. A change in user behaviour might lead to a change in the processes running on a computer. The change in processes leads to a change in the measured aggregate resource consumption. This seems reasonable. However, contention for resources may also lead to feedback on the processes, causing them to alter their behaviour, possibly even crash. Thrashing is such behaviour that can even prove fatal to a process. It results from interactions between parts, not from a single causal channel, thus we cannot definitely say that process changes cause performance changes; it is also true that performance changes may cause process changes. There is causation in both directions, at different scales.

In the latter example, these observations can be reduced to a discussion about whether the active parts of a system are able to keep certain promises or not. For a full discussion, we defer to volume 2.
Applications and Further Study 9.

- Quantitative discussion of the flows of information and instruction in a system.
- Measurement of workflow in a system.
- Gauging the controllability of a system.
- Determining how focused, constrained (low entropy) or distributed (high entropy) a process or structure is.
- Maximization of entropy for balancing the idea of “what can happen will happen” with the known constraints.
CHAPTER 10

STABILITY

One of our fundamental premises about systems is that medium term stability, allowing for long term change, is a desirable concept. Systems must be predictable for long enough to perform their intended function. This applies both to the human and machine parts of a system. However we still need a quantitative description of what such stability means, and on what timescales.

10.1 BASIC NOTIONS

If we place a ball at the crest of a hill, the slightest movement will cause it to roll down to the bottom. A ball placed at the top of a hill is a mechanical system which is said to be unstable to small perturbations, i.e. a small push changes the character of the system. By contrast, a ball placed at the bottom of a valley or trough is said to be stable to perturbations, because a small push will only take it a short way up the hill before it rolls back down again, preserving the original condition of the system.

Stability is an important idea in systems. If a small change can completely alter a system, then its usefulness is limited. A bomb is a chemical-mechanical system which can only be used once, because it is impractical to reset it to its original condition once it has exploded. Instability affects a great many dynamical systems, from financial systems, to computers, to social systems. The idea of building stability into systems is thus of central importance to human-computer administration.

Modern fighter jets are built with an inherent instability under flight, unlike passenger jets which are inherently more stable. Fighter jets are much more maneuverable because of this ability to lapse into instability. The price one pays for this is a much more essential and risky regulation requirement that maintains the system right at the edge of instability, allowing rapid, controlled change, but flirting with rapid, uncontrolled change.
10.2 Types of Stability

There are various notions of stability, but they all have in common a search for states that are not significantly altered when we perturb the system by a small amount. We can write most notions of stability in the generic form:

**Definition 39 (Stability).** A stable state is one that is preserved, up to a multiplying factor, when perturbed by some operation. Let \( Q \) be a state of a system, and let \( \hat{\Delta}_\lambda \) be an unspecified operator that perturbs \( Q \) by an amount \( \lambda \) in some parameter. If the generic perturbation leads to the same state \( Q \) multiplied by a scale factor \( \Omega(\lambda) \), for some function \( \Omega \), the state may be described as \( \Omega \)-stable under this operation; i.e. the result of a perturbation

\[
\delta Q \equiv (\hat{\Delta}_\lambda(Q) - \Omega(\lambda)Q) = 0.
\]  

(10.1)

This definition associates stability with stationary variations. This is not a rigorous definition, since the descriptions of \( \hat{\Delta} \) and \( \Omega \) are vague; however, it expresses the general concept behind a broad range of ideas of stability. In some cases, it is desirable to demand restrictions on \( \Omega(\lambda) \).

10.3 Constancy

The simplest kind of stability is constancy (\( \Delta = \Omega = 1 \)). Mathematically, we express this by saying that a system variable does not change at all with respect to its parameters:

\[
q(t, x_1, x_2, \ldots) = \text{const} = q.
\]  

(10.2)

We can change time and the other parameters of the system \( x_i \), but such change has no effect on \( q(t, x_i) \) because the function is a trivial one. Such constancy is a rather simplistic viewpoint that is rarely true of anything but the simplest systems, but it is a convenient approximation in many cases. A slightly more realistic viewpoint is to only expect constancy on average.

Average constancy allows a system to be dynamic and to change over short time intervals, in an unspecified way, as long as it changes back again so that the average result is zero over longer times:

\[
\langle q(t) \rangle_{\Delta t} = \frac{1}{\Delta t} \int_{-\Delta t/2}^{+\Delta t/2} q(t)dt = \text{const}.
\]  

(10.3)

An oscillation is a deterministic example of this; random fluctuations are a non-deterministic example. This kind of stability allows us discuss situations where there are fluctuations in the behaviour of system that average to zero over time. In this case we must ask: how long do we have to observe the system before the fluctuations will average out.
An enhancement of the previous case is to allow the average of a function to change by a small amount, i.e. to exhibit a trend, by varying slowly at rate that is much slower than fluctuations that almost average out. This is called a separation of scales, and is discussed in section 8.5.

Finally, a more advanced notion of stability, in dynamic systems that exhibit fluctuations is statistical stability. Here we ask the question, in a system that is stochastic, i.e. exhibits unpredictable fluctuations, with a particular statistical distribution of values, are there certain statistical distributions that are the natural result of system behaviour? Are some distributions more stable than others, if we change the system slightly? Such distributions summarize the stability of the management scale, without disallowing the minutiae of the system.

10.4 Convergence of behaviour

Systems at any scale can exhibit oscillatory or random behaviour. It is sometimes a desirable property and sometimes an undesirable one. An oscillation with fixed frequency (such as a swinging pendulum or a daily task list) is said to be in a steady state, even though it is changing in time in a regular way. Such a steady state is also called a limit cycle, if it is the result of a process of convergence towards this final steady state from an irregular pattern. Generally we are interested in systems that enter into a steady state, i.e. either a static or dynamical equilibrium.

Another possibility is that oscillations die out and leave a static state. In order to converge to a static state, oscillations must be dissipated by a drain on the system, that behaves analogously to the friction in a pendulum.

Example 97. Circular dependencies often result in oscillations that need to be damped out. Suppose host $A$ requests a result from a database that is located on host $B$, but the data in the database running on host $B$ is located physically on host $A$ and is shared by a network service. A single request from host $A$ results in traffic

$$A \rightarrow B \rightarrow A \rightarrow B \rightarrow A.$$  \hspace{1cm} (10.4)

Moreover, a timeout, or failure could result in longer oscillations of period $A$ – $B$. A local cache of data could be used to dampen these oscillatory convulsions and relieve the network of the unnecessary oscillatory behaviour, or the system could be reorganized to alter the flow of communication between the client, the disk and the database.

Example 98. The build up of temporary disk files on a computer system can result in an escalating space problem that can be kept under control by introducing a counterforce that makes the behaviour oscillatory. Human work patterns have a daily rhythm, so daily tidying of garbage can keep the build up within manageable levels. Thus, in this example we are trying to achieve a steady limit cycle from a behaviour that is originally divergent, by introducing a countermeasure.
Example 99. Tit for tat reprisals between users and/or administrators are ping-pong oscillations that could escalate or, in principle, never terminate. A prolonged bout of tit for tat is warfare.

We have two possibilities for modelling converging oscillatory behaviour: discrete or continuous models. Discrete changes models use graphs or chains of discrete changes can be used to trace the possible changes in the system. Cycles can then be identified by looking at the topology of the graph of allowed transitions. One way to define convergent behaviour is to consider the transition function $T_{ij}$ of the system that determines what state $q_j$ a system that is in a state $q_i$ will make a transition to. Suppose that, no matter what state we are in, a number $n$ of transitions will bring us into a definite state $q_c$:

\[
(T_{ij})^n q = q_c \\
T_{ij} q_c = q_c
\]

(10.5)  
(10.6)  
(10.7)

Such a transition function $T_{ij}$ may be called convergent. Such a state $q_c$ is called a fixed point of the function (see section 10.13).

If we are willing to ignore the discrete details of the system, over longer times, then another picture of this can be drawn for smooth and differentiable systems (the continuum approximation). Let $q$ be the state of the system and $\gamma$ is the rate at which it converges to a stable state. In order to make oscillations converge, they are damped by a frictional or counter force $\gamma$. The solutions to this kind of motion are damped oscillations of the form

\[
q(t) \sim e^{-\gamma t} \sin(\omega t),
\]

(10.8)

for some frequency $\omega$ and damping rate $\gamma$. Three cases are distinguished: under-damped motion, damped and over-damped motion. In under-damped motion $\gamma \ll \omega$, there is never sufficient counterforce to make the oscillations converge to any degree. In damped motion the oscillations do converge quite quickly $\gamma \sim \omega$. Finally with over-damped motion $\gamma \gg \omega$ the counter force is so strong as to never allow any change at all.

<table>
<thead>
<tr>
<th>Under-damped</th>
<th>Inefficient: the system can never quite keep oscillations in check.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Critically-damped</td>
<td>System converges in a time scale of the order the rate of fluctuation.</td>
</tr>
<tr>
<td>Over-damped</td>
<td>Draconian: changes are stopped before completing a single cycle.</td>
</tr>
</tbody>
</table>

An over-damped solution to system management is rarely acceptable. An under-damped solution will not be able to keep up with the changes to the system made by users or attackers.
10.5 MAXIMA AND MINIMA

The extrema of smooth functions can be used to define stability. The extrema are found at the turning points of functions, where the rate of change of the function is smallest.

\[
\frac{dq(\lambda)}{d\lambda} = 0.
\]  

(10.9)

Thus, both maxima and minima exhibit local sluggishness to change; however, only minima are stable to perturbations.

Maxima and minima can also be defined for non-smooth domains, such as the nodes of a graph, provided there is a function which defines a value \( \phi_i \) on each node \( i \). For instance, if we have a graph formed from a discrete set of nodes, a node is a maximum if it has a greater value \( \phi_i \) than any of its neighbouring (adjacent) nodes.

**Definition 40** (Local maximum). If \( \phi \) is a mapping from a domain \( i \) to some range, then a node \( i \) in the domain is a local maximum of the mapping if the nearest neighbours \( j \) of node \( i \) have strictly lower values \( \phi_j \) than node \( i \) itself; i.e., if \( A_{ij} \) is the adjacency matrix of the graph, a local maximum satisfies:

\[
\phi_i > \phi_j, \quad \forall \{j | A_{ij} \neq 0\}.
\]  

(10.10)

This definition works equally well for sparse or dense sets, i.e., for functions of a continuous variable or for discrete graphs.

10.6 REGIONS OF STABILITY IN A GRAPH

Another way of talking about the stability of structures is to define regions of persistence in mappings (like the eye of a storm, amongst all the connections). If a mapping leads us into a natural resting place, the graphical equivalent of a basin or minimum, then we can use that to define stability.

There are two distinct notions of stability for graphs that are very useful for us throughout this book.

**Definition 41** (Internal stability). A set \( S \in X \) in a graph \( (X, \Gamma) \) is said to be **internally stable**, if no two nodes in \( S \) are connected, i.e., if

\[
\Gamma S \cap S = \emptyset.
\]  

(10.11)

In other words, internal stability says that a region of internal stability consists of points that are cannot be exchanged for one another (fig. 10.1). None of them have any arrow of link between
CHAPTER 10. STABILITY

Figure 10.1: An internally stable set is a set of peers within a graph, that are unconnected by any single hop. Once we get into an internally stable state, we cannot get into any other internally stable state without backtracking out of the internally stable region.

them that might identify one as being better than another in the sense of the mapping $A$. This property makes them ‘as good as each other’ as end points in a journey through the graph. In the theory of games, that we shall turn to later, this property is used to mean that none of the points in an internally stable set dominate any of the others, thus they are all equally valid solutions to the game.

Figure 10.2: An externally stable state is accessible to all points outside the externally stable region, by a single hop.
Definition 42 (External stability). A set \( T \subseteq X \) in a graph \( (X, \Gamma) \) is said to be **externally stable** if every node \( x \notin T \), outside of \( T \), satisfies
\[
\Gamma x \cap T \neq \emptyset,
\]
(10.12)

i.e. the image of every node inside of \( T \) lies outside of \( T \), or conversely: every point outside of \( T \) (in the complement set \( X - T \)) maps into \( T \) by the inverse mapping
\[
X - T \subseteq \Gamma^{-1}T.
\]
(10.13)

External stability tells us that a stable set is a place, like a local minimum, where the graph connections lead us into an end point. In other words, at least one node inside the externally stable set is ‘better’ than any node outside the set, in the sense of the mapping \( A \).

Definition 43 (Kernel). A set in a graph that is both internally and externally stable is said to be a **kernel** of the graph. The kernel is free of loops and contains all points \( x \in X \) for which \( \Gamma x = \emptyset \).

The importance of the kernel for systems is that \( \Gamma \) can be regarded as a mapping of states, i.e. of transitions between states of a system — and we are interested in having these transitions converge towards some stable end state. The kernel of a graph contains all the states that can be regarded as being such end states\(^1\). Internal stability tells us the possible candidates that are at the end of a sequence of arcs, and that once we arrive inside the set, we have chosen one of the set. External stability tells us that we can always get to one of those points from outside the region, thus the region is accessible to the whole system. If we require both, then the conclusion is that the kernel is the set of states that is accessible to the whole system and is stable under single hop perturbations.

The kernel is an important concept of equilibrium in games. The notion of a kernel was introduced into game theory by Von Neumann and Morgenstern ([NM44]) as a proposal for the solution of a game, for finding preferential strategies. Since a point in the kernel is internally stable, no other point is preferable to it; moreover, since it is externally stable it is preferable to any place outside the kernel. Not all graphs have kernels, but kernels are guaranteed in a number of cases (see, for instance, [Ber01]).

Note that although we will always be able to reach a stable node in the kernel, from any point in the system, we will not necessarily be able to find the ‘best’ node according to some extra criterion, such as centrality or some other ‘hidden variable’. The graph mapping \( \Gamma \) itself does not distinguish between the nodes in the internally stable set — they are all valid end states. However, by jumping out of the region and back into it through multiple hops, we can find alternative points in the region.

---

\(^1\)Note: the kernel of the mapping is not to be confused with the ‘Heat Kernel’ generating functional of the graph (see[Chu97]) that is related to algebraic geometry and field theory.
that might satisfy additional criteria. This means that other notions of ‘preferable’ could be allowed
to select from the set of states in the kernel of the graph.

10.7 GRAPH STABILITY UNDER RANDOM NODE REMOVAL

The stability of regions and graph structure to node removal has been discussed by various authors.
See [AB02] for a review. The stability of local regions in a graph is somewhat unpredictable to the
removal of nodes. If we use centrality to define regions (see section 6.5), then it might seem that the
removal of the most central nodes in a network would have the most damaging results. However,
some graphs with random graph properties are extremely invulnerable to random node removal,
including the most central nodes. Peer to peer graphs are examples of this: because they have
no true centre (the maxima are rarely very ‘high’ above the rest), their structures remain largely
unchanged by random node removal. Other structures, such as hub configurations are extremely
susceptible to node removal, since the most central points are of significantly greater importance to
the connectivity of the graph. Node removal is clearly more serious to the network as a whole, but
perhaps not to individuals (see fig. 10.3).

Figure 10.3: Network tolerance to node removal: nodes are more important than connectors.

One might be tempted to use centrality as a measure of stability, however there is no direct
correlation between the centrality values and the connectivities, since the scale-free eigenvector
values do not retain information about total numbers of nodes in a graph, thus tiny graphs of a
few nodes could have similar centrality values to a huge graph. Centrality is only of interest as a
relative measure for connected components with a fixed number of nodes.
Susceptibility to node removal can be gauged by examining the degree distributions of the nodes

\[ P(k) = \frac{n(k)}{\sum_k n(k)}, \]  

(10.14)

where \( n(k) \) is the number of nodes with \( k \) connected neighbours. Studies show that random node removal of a fraction \( f \) of a graphs nodes has varying effects depending on this degree distribution (see [AB02] for a review). Large networks undergo phase transitions from states of being connected to being fragmented when critical fractions of nodes are removed.

Scale free (power law) networks have node degree distributions

\[ P(k) \propto \frac{1}{k^\alpha}, \]  

(10.15)

for some positive constant \( \alpha \). These are especially robust to random node removal, and it is known that peer to peer graphs have this structure (see [Bar02]). Other reports of self-similar and power law behaviour can be found in connection with the World Wide Web (see [BA99, BAJ00, A JB99, GHH91]).

Attacks by deliberate targeting of the largest nodes are more efficient and breaking up graphs than random failures. Studies typically show numbers at around the order of magnitude of ten percent level for fragmenting graphs. A greater fraction of nodes must be destroyed to break up a highly connected graph (see section [AB02]).

10.8 Dynamical equilibria: compromise

A ball sitting at the trough of a valley is said to be in a state of static equilibrium. The forces acting upon it are balanced, and the result is that nothing happens. A ‘tug of war’, on the other hand, is only in a state of dynamic equilibrium when the two groups pulling on the rope are not moving. Disk storage is a tug of war.

Example 100. A computer storage system under normal usage, is being filled up with data by the actions of users, but there is no counterforce which frees up any of the space again. This is not in equilibrium. A garbage collection service can maintain a dynamic equilibrium if the average amount of storage in use is constant over time.

Example 101. In a car park (parking lot), cars come and go. The rate of arrival and the rate of departure determine whether the total number of cars is increasing, decreasing or in balance. If the number of cars is in balance, then we say that a dynamic equilibrium has been reached.

Equilibrium can be mechanical, chemical, or statistical. By implication, equilibrium is a balance between opposing forces. In a dynamical equilibrium, there is motion which is continually being reigned in check by other processes.
10.9 Statistical stability

A system that settles into a predictable average behaviour, that withstands perturbations to its fluctuation distribution, can be said to exhibit statistical stability (see [Hug95, Sat99]).

Suppose that a system is characterized by a measurable \( q(t) \) that varies with time (the same analysis can be applied to any other control parameter). As the state \( q(t) \in \{ q_1, q_2, \ldots, q_i \} \) varies, it changes value within a set of possible values. In a deterministic system, the pattern of change is predictable at any given moment; in a non-deterministic system it is unpredictable. Either way, if there is sufficient regularity in the behaviour of the system — so that a knowledge of the past can predict an outcome in the future — then we can characterize the change over time by plotting the distribution of values as a histogram (see fig 10.4) that represents the probability \( P(q) \) of measuring a given \( q_i \).

![Figure 10.4: The probability distribution of values \( q_i \) for a given measurable characterizes the average spread of values that occur over time. If this distribution maintains the same form, as the system develops then it can be called a stable distribution. For systems with random changes, there is only a few possible distributions that are stable over long times: these are the Lévy distributions and the Gaussian ‘normal’ distribution.](image)

But what happens when we mix together different signals, each with their own distribution of values? The result will surely be a different distribution that is neither the one nor the other, but are there any special distributions that are the stable end result of a lot of mixing of this kind? These would represent the limits of predictability for otherwise unpredictable systems. A. Cauchy and P. Lévy asked this question: what distributions have the property that, when perturbed by mixing, they retain their essential features? Or, is there a \( P(x) \) such that the convolution of two signals with the same \( p(x) \) results in the same distribution:

\[
P(q) = \int dq' P_1(q) P_2(q - q')
\] (10.16)
CHAPTER 10. STABILITY

Another way of expressing this is that, if \( q_1(t) \) and \( q_2(t) \) are random variables with stable distributions, then so is \( q_1(t) + q_2(t) \). The Gaussian distribution is one solution to this problem; the other solutions are called the stable Lévy distributions. Some of these have infinite variance and therefore cannot truly represent the behaviour of real systems. However, they do describe systems approximately\(^2\).

The Lévy distributions are denoted by \( L_\alpha(q) \) (\( 0 < \alpha \leq 2 \)), and most of them cannot be written down as analytical expressions in terms of \( q \). The Fourier transforms (or characteristic functions) of the distributions

\[
L_\alpha(k) = \int dq \, e^{iqk} L_\alpha(q)
\]

(10.17)

(called their characteristic functions) can be written down however. The symmetrical distributions have characteristic functions of the form

\[
L_\alpha(k) = \exp (-c_\alpha |k|^\alpha),
\]

(10.18)

for constants \( c_\alpha \). These allow one to work out the asymptotic behaviour for large \( q \), which turns out to follow a power law behaviour:

\[
L_\alpha(q) \sim \frac{\alpha A}{|q|^{1+\alpha}}, \quad q \to \pm \infty.
\]

(10.19)

for constant \( A \). Two exceptions exist that can be integrated straightforwardly to obtain an analytical form (see appendix D): one is the so-called Cauchy distribution:

\[
L_1(q) = \frac{c_1/\pi}{q^2 + c_1^2};
\]

(10.20)

the other is the Gaussian

\[
L_2(q) = \frac{1}{2\sqrt{\pi c_2}} e^{-q^2/4c_2}.
\]

(10.21)

For other values of \( \alpha \), the full asymptotic form can be used for large \( q \to \infty \):

\[
L_\alpha(q) = \sum_{n=1}^{\infty} \frac{(-1)^{n+1}}{\pi} \frac{c_\alpha^n}{q^{1+n\alpha}} \frac{\Gamma(1+n\alpha)}{\Gamma(n+1)} \sin(\pi \alpha n/2).
\]

(10.22)

and for small \( q \to 0 \):

\[
L_\alpha(q) \sim \frac{1}{\pi} \sum_{m=0}^{\infty} \frac{(-1)^m}{(2m)!} \frac{q^{2m}}{2^{m+1} c_\alpha^m} \Gamma \left( \frac{2m+1}{\alpha} \right).
\]

(10.23)

The range of values of \( \alpha \) that leads to stable behaviour is limited to \( 0 < \alpha \leq 2 \), due to the scaling behaviour noted in section 10.10; \( \alpha \) must exceed 0 in order to have non-negative

---

\(^2\)Exponentially truncated forms of the Lévy distributions are sometimes used to discuss realistic examples.
probabilities, and must be less than or equal to 2 or else there is only short-range dependence of the data and the distribution has insufficient ‘memory’ to form a stable distribution. This phenomenon is sometimes called $\alpha$-stability and the parameter $\alpha$ is related to the Hurst parameter $H = 1/\alpha$ described in section 10.10.

Investigating statistical stability allows us to determine whether probabilistic management policies are sustainable over time. For example, if the fluctuations in a system are not stable, then we can make no guarantees about the system in the future.

10.10 SCALING STABILITY

A related form of stability to the statistical stability is stability under scaling. This is also called scale-invariance, and it asks a more specific version of the same question as in section 10.9: if we have a fluctuating variable $q(t)$, how does the size of the fluctuations depend on how closely we examine the system. If we view the system through a ‘magnifying glass’ by focusing on small times, we might see large fluctuations, but if we reduce the resolution by stepping back from the system and examining large time intervals, how does the size of fluctuations change in relation to our resolution. A scale transformation on time is called a dilatation.

If the relative size of fluctuations is the same at all scales, the system is said to exhibit statistical self-similarity. The scaling hypothesis, for a function $q(t)$, under a dilatation by an arbitrary constant $s$, is expressed by:

$$q(st) = \Omega(s) q(t).$$

(10.24)

In other words, the assumption is that stretching the parameterization of time $t \rightarrow st$, leads to a uniform stretching of the function $q(t)$, by a factorizable magnification $\Omega(s)$. The function retains its same ‘shape’, or functional form; it is just magnified by a constant scale.

This property is clearly not true of an arbitrary function. For example, $q(t) = \sin(\omega t)$ does not satisfy the property. Our interest in such functions is connected with dynamical systems which exist and operate over a wide range of scales. Physical systems are always limited by some constraints, so this kind of scaling law is very unlikely to be true over more than a limited range of $s$ values. Nevertheless, it is possible to discuss functions which, indeed, scale in this fashion, for all values of $s$, as an idealization. Such functions are said to be scale invariant, dilatation invariant, or self-similar.

Exact self-similarity, for all $s$, is only a theoretical possibility, and has led to the study of fractals, but a similarity in statistical profiles of functions is a real possibility over finite ranges of $s$. This is a weaker condition, which means that the behaviour of a complete system $S$ is invariant, but that $q(t)$ itself need not be.

$$S[\Omega^{-1}(\alpha) q(\alpha t)] \rightarrow S[q(t)].$$

(10.25)
From eqn (10.24), the symmetry between \( q(t) \) and \( \Omega(s) \), tells us that
\[
q(x) \sim \Omega(x),
\]
(10.26)
i.e. that they must possess similar scaling properties. In fact, \( q(t) \) and \( \Omega(s) \) must be homogeneous functions, in order to satisfy this relationship:
\[
q(t) = t^H, \\
\Omega(s) = s^H,
\]
(10.27)
for some power \( H \). In other words, one has a general scaling law:
\[
 s^{-H} q(st) = q(t).
\]
(10.28)
We apply this to locally averaged functions:
\[
 s^{-H} \langle q(st) \rangle = \langle q(t) \rangle.
\]
(10.29)
The exponent \( H \) is called the Hurst exponent, after the Hydrologist H.E. Hurst who studied such behaviour in the flows of the Nile river. It can be estimated for real data by noting that, over an interval \( \Delta t \),
\[
\langle \max(q(t)) - \min(q(t)) \rangle_s \Delta t = s^H \langle \max(q(t)) - \min(q(t)) \rangle_{\Delta t},
\]
(10.30)
i.e.
\[
H = \frac{\log \left( \frac{\langle \max - \min \rangle_{s \Delta t}}{\langle \max - \min \rangle_{\Delta t}} \right)}{\log(s)}.
\]
(10.31)
Note that this estimator will give an incorrect answer for exponential functions \( \exp(t^n) \) that increase monotonically; it should only be used on genuine time series. For the Gaussian distribution, we have \( H = \frac{1}{2} \), since
\[
(\max - \min) \frac{1}{\sqrt{2\pi}\sigma^2} \exp \left( -\frac{q^2}{2\sigma^2} \right) \sim \frac{1}{\sigma},
\]
(10.32)
and
\[
\sigma(st) = \frac{1}{\sqrt{s}} \sigma(t),
\]
(10.33)
thus
\[
H_{\text{Gauss}} = \frac{\log(\sqrt{s})}{\log(s)} = \frac{1}{2}.
\]
(10.34)
The parameter $H$ is related to the parameter $\alpha$ from the previous discussion in section 10.9 on $\alpha$-stability by $H = 1/\alpha$. The Hurst exponent characterizes several properties of the signal $q(t)$’s correlation functions. The auto-correlation is sometimes referred to as the ‘memory function’ of the system.

For $\frac{1}{2} < H < 1$ processes have long-range dependence, or are correlated significantly with values of the function in the distant past. For $H = \frac{1}{2}$ observations are uncorrelated, and for $0 < H < \frac{1}{2}$ they have short-range dependence and the correlations sum to zero (see [Ber94] for details).

For $H \geq 1$ the second moment of the probability distribution (the variance $\sigma^2$) diverges as a further symptom of long-range dependence. This means that the system can have significant numbers of fluctuations of arbitrarily large size.

Statistical self-similarity as been observed in network traffic ([LTWW94]).

**Example 102.** Time series measurements\(^5\) from a local area network measured over an eleven weeks span at 5 different time scales of up to a week, give the following root mean square values for the Hurst exponent, with 10% tolerance (see fig. 10.5 and table 10.1). Here we find no values lower than $H = \frac{1}{2}$, which means that all of the results belong to potentially $\alpha$-stable distributions. Incoming Web traffic, in particular has a high value of $H$ which tends to suggest a long tailed distribution that signifies very large fluctuations at all the measurable scales. However, this is is slightly misleading and is the result of a stable non-equilibrium process that is superimposed on the fluctuating signal as shown in [BHRS01].

The Hurst exponent measures apparent self-similarity. Self-similarity is sometimes linked to long-tailed distributions that are typical of the asymptotic Lévy distributions. However, care is needed to draw any connection between estimates of the Hurst exponent and estimates of the distribution of values measured from limited amounts of data. Heavy tailed distributions can be caused by monotonically growing fluctuations during the sampling period, for instance, without any scale-invariant behaviour in the fluctuations. It is not just the distribution of values that is important to scale-free behaviour, but the order in which the distributed values occur. Thus self-similar behaviour is not the same as stability of fluctuation distributions. This is the peril of basing conclusions of average measures.

### 10.11 Maximum entropy distributions

Entropy measures the even distribution of resource consumption in a system. A high entropy means a high degree of resource fragmentation. Maximum entropy distributions are also a kind of stability criterion. By maximizing entropy, we assure a most probable distribution, in the limit of long

\(^5\)These measurements were made using the cfenvd daemon in cfengine ([Bur93]).
Table 10.1: Estimated Hurst exponents for time series of traffic intensities for different traffic types, measured on a single computer over several weeks. Human activities such as numbers of users follow a basically Gaussian profile $H = \frac{1}{2}$, while local and wide area network driven measurements show higher values of the Hurst exponent. Measurements like these are sometimes used to show evidence of self-similar behaviour in computer service patterns, but closer study of the data is required to draw this conclusion.

<table>
<thead>
<tr>
<th>Variable</th>
<th>$\frac{1}{2} &lt; H &lt; 1$</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of users</td>
<td>0.5</td>
</tr>
<tr>
<td>Root processes</td>
<td>0.6</td>
</tr>
<tr>
<td>Other processes</td>
<td>0.7</td>
</tr>
<tr>
<td>Diskfree</td>
<td>0.8</td>
</tr>
<tr>
<td>Load average</td>
<td>0.6</td>
</tr>
<tr>
<td>Incoming netbiosns</td>
<td>1.6</td>
</tr>
<tr>
<td>Outgoing netbiosns</td>
<td>1.8</td>
</tr>
<tr>
<td>Incoming netbiosdgm</td>
<td>1.4</td>
</tr>
<tr>
<td>Outgoing netbiosdgm</td>
<td>1.5</td>
</tr>
<tr>
<td>Outgoing netbiosssn</td>
<td>1.6</td>
</tr>
<tr>
<td>Incoming nfsd</td>
<td>2.0</td>
</tr>
<tr>
<td>Outgoing nfsd</td>
<td>2.3</td>
</tr>
<tr>
<td>Incoming smtp</td>
<td>1.6</td>
</tr>
<tr>
<td>Outgoing smtp</td>
<td>2.0</td>
</tr>
<tr>
<td>Incoming www</td>
<td>2.5</td>
</tr>
<tr>
<td>Outgoing www</td>
<td>1.1</td>
</tr>
<tr>
<td>Incoming ftp</td>
<td>1.7</td>
</tr>
<tr>
<td>Outgoing ftp</td>
<td>2.2</td>
</tr>
<tr>
<td>Incoming ssh</td>
<td>1.5</td>
</tr>
<tr>
<td>Outgoing ssh</td>
<td>1.4</td>
</tr>
<tr>
<td>Incoming telnet</td>
<td>1.2</td>
</tr>
</tbody>
</table>
times. It is a limiting point that a system will tend to by virtue of randomness. Put another way, chance will never favour a different configuration once we have arrived in a maximum entropy configuration, so this provides an important reference point for a future state of any system. See section 9.11 for more about this topic.

10.12 EIGENSTATES

Eigenvalues are especially stable solutions of simultaneous linear equations. For an $N \times N$ matrix $M$, eigenvectors $\vec{v}_\lambda$ and their associated eigenvalues $\lambda$ satisfy the matrix equation

$$M \vec{v}_\lambda = \lambda \vec{v}_\lambda, \quad (10.35)$$

or in component form:

$$M_{ij} \vec{v}_{\lambda}^j = \lambda \vec{v}_{\lambda}^i. \quad (10.36)$$

Put another way, when the matrix $M$ acts on certain vectors, the vectors can become longer or shorter (by a factor $\lambda$) but they still point in the same direction. If this were to be true for any vector $\vec{v}$, it would only be true for diagonal matrices; however, every non-singular $N \times N$ matrix has
this property for a special set of $N$ linearly independent vectors, and for special values $\lambda$. These vectors are the eigenvectors and eigenvalues of the matrix.

Eqn. (10.35) has the form

$$\delta \vec{v}_\lambda = (M \vec{v}_\lambda - \lambda \vec{v}_\lambda) = 0,$$

and thus defines a set of stable vectors. For $\lambda = 1$, the vectors are invariant, or constant under the perturbation by $M$.

**Example 103.** Consider the adjacency matrix of a simple graph (fig 10.6). The adjacency matrix

![Figure 10.6: A three node graph with symmetry about node 2.](image)

plays a dual role for a graph: both as a representation of the connectivity between adjacent nodes and as a recipe for summing over nearest neighbours (see section 6.5). Consider the simple three node graph, where nodes (1,2) and (2,3) are joined by two links. The adjacency matrix is given by

$$A = \begin{pmatrix} 0 & 1 & 0 \\ 1 & 0 & 1 \\ 0 & 1 & 0 \end{pmatrix}$$

(10.38)

This has eigenvalues $\lambda = \{0, \pm \sqrt{2}\}$ and corresponding eigenvectors:

$$\begin{pmatrix} v_1 \\ v_2 \\ v_3 \end{pmatrix} = \left\{ \begin{pmatrix} 1 \\ 0 \\ -1 \end{pmatrix}, \begin{pmatrix} \frac{1}{\sqrt{2}} \\ \frac{1}{\sqrt{2}} \\ -\frac{1}{\sqrt{2}} \end{pmatrix}, \begin{pmatrix} \frac{1}{\sqrt{2}} \\ -\frac{1}{\sqrt{2}} \\ \frac{1}{\sqrt{2}} \end{pmatrix} \right\}.$$

(10.39)

The structure of the eigenvalues and eigenvectors reflects the symmetry of the graph, and the principal eigenvector (that belonging to the highest eigenvalue) has its highest value for node number 2, indicating that it is the most ‘central’ node in the graph.

Sometimes eigenvectors and eigenvalues have more subtle meanings that arise from the mathematical structure that underlies a problem. One must always be careful in interpreting mathematical results, in terms of the assumptions that are entered at the start.
Example 104. Consider two service departments or servers $d_1$ and $d_2$ that work together share the load of work between them, with manager $m$ (fig 10.7). Both departments have a limited capacity $C$ and can only just cope with the total work load alone. If both worked flat out, they could muster a total of

$$C_{\text{max}} = \sqrt{C_1^2 + C_2^2},$$

(10.40)

as independent units. However, in order to provide for full redundancy, we must not use up this total capacity, but instead require $C_1 = C_2 = C_{\text{tot}}$. Now, to balance their load and provide redundancy they split the workload between themselves in such a way that either one could take over at a moment’s notice, by changing a management parameter $\theta$.

$$d_1 \rightarrow d_1 \cos \theta + d_2 \sin \theta$$

(10.41)

$$d_2 \rightarrow -d_1 \sin \theta + d_2 \cos \theta.$$  

(10.42)

Are there any stable or preferred solutions to this problem? It is helpful to draw this situation geometrically by defining a vector:

$$\begin{pmatrix} d_1 \\ d_2 \\ m \end{pmatrix} \rightarrow \begin{pmatrix} v_1 \\ v_2 \\ v_3 \end{pmatrix}$$

(10.43)
in three dimensional space.

Consider a matrix $M$ representing a two dimensional rotation about the $m$ axis. The manager can choose to rotate the load from one department to another by changing the angle $\theta$. What are the privileged vectors for this perturbation matrix?

$$M = \begin{pmatrix} \cos \theta & \sin \theta & 0 \\ -\sin \theta & \cos \theta & 0 \\ 0 & 0 & 1 \end{pmatrix} \quad (10.44)$$

The characteristic equation for the eigenvalues is

$$\det | M - \lambda I | = 0,$$

(10.45)

giving

$$\lambda = 1, \exp(\pm i\theta),$$

(10.46)

and eigenvectors

$$\begin{pmatrix} v_1 \\ v_2 \\ v_3 \end{pmatrix} = \begin{pmatrix} 1 \\ i \\ 0 \end{pmatrix}, \begin{pmatrix} 1 \\ -i \\ 0 \end{pmatrix}, \begin{pmatrix} 0 \\ 0 \\ 1 \end{pmatrix}. \quad (10.47)$$

In this example, two of the eigenvalues are complex numbers, but this should not distract from the interpretation of the result. For $\lambda = 1$, we find an eigenvector along the management rotation axis. This tells us that the management axis has a privileged status in this system: it is independent of any change by varying $\theta$. The remaining two eigenvalues and eigenvectors appear to be lines pointing at 45 degrees into the complex plane. If we substitute these values into the eigenvalue equation $M \vec{v} = \lambda \vec{v}$, we find that it is identically satisfied, i.e. no conditions are placed on $\theta$ or a specific configuration. This makes sense, from our original system design: the rotational symmetry was introduced to provide redundancy, or independence of configuration, so any balance is as good as any other.

This last example illustrates an important point: eigenvectors and eigenvalue solutions fall into two categories: solution by linear identity and solution by invariance (symmetry), so eigenvalue solutions will always tell us the intrinsic, stable or invariant properties of the perturbation matrix. Eigenvectors are important in dynamics because systems often have preferred configurations that are described by their eigenvectors.

\footnote{The presence of $i = \sqrt{-1} = \exp(i\pi/2)$ in $(1, \pm i)$ signals a phase shift between the preferred solution for $d_1$ and $d_2$ of 90 degrees ($\pi/2$ radians). This is exactly reflected in the placement of $\cos$ and $\sin$ in eqn. (10.42), since sine and cosine are phase shifted by 90 degrees with respect to one another.}
10.13 FIXED POINTS OF MAPS

In the theory of dynamical systems, stationary points where derivatives vanish are sometimes called fixed points, however there is a more fundamental and interesting definition of fixed points that is of great importance to average stability of human-computer systems.

Any function \( x' = f(x) \) defines a mapping from some domain of values to a range of values. In a continuum description, the domain and range are normally subsets of the Euclidean space of real numbers \( \mathbb{R}^n \). A fixed point \( x^* \) of a mapping \( f(x) \) is any point that maps onto itself:

\[
f(x^*) = x^*.
\]

(10.48)

In other words, if a system finds itself in \( x^* \) by iteration of this mapping, it will remain there. This is clearly a definition of stability, since:

\[
\delta Q^* = (f(Q^*) - Q^*) = 0.
\]

(10.49)

We are particularly interested in functions that determine the time development or running of a system, since \( Q^* \) is a natural choice for a configuration of a system that is required to be stable.

In this section we describe, without proof or detailed explanation, two notions of fixed point that relate to continuous functions: the Brouwer fixed point theorem and the Kakutani fixed point theorem\(^5\).

**Theorem 1** (Intermediate values). Let \( f : [a, b] \to \mathbb{R}^1 \) be a continuous function, where \([a, b]\) is non-empty, compact, convex subset of \( \mathbb{R}^1 \) and \( f(a) f(b) < 0 \), then there exists an \( x^* \in [a, b] \) such that \( f(x^*) = 0 \).

This theorem makes the simple point that if two points lie in the range of a continuous function, and one point is positive and the other is negative, then the function has to cross the \( f = 0 \) axis, by virtue of its continuity. The only way that a function would not satisfy this property is if it were broken into disjointed pieces. Although the theorem uses the value \( f = 0 \) this construction, the result remains true if we relabel the axes to any values. The important point about this theorem is a corollary. If we restrict the domain and range to the unit interval (for convenience)

**Corollary 1.** Let \( f : [0, 1] \to [0, 1] \) be a continuous function, then there exists a fixed point \( x^* \in [0, 1] \) such that \( x^* = f(x^*) \).

This tells us that, for a function that is convex over the unit interval, and therefore covers the range \([0, 1]\), the function must cross the 45 degree line \( x = f(x) \) at at least one point \( x^* \). This leads us to Brouwer’s theorem, which is a generalization of this idea for more general sets.

To describe the two fixed-point theorems, we need to use mathematical terms describing sets: convex and upper semi-continuous\(^6\).

\(^5\)Several excellent explanations of these results exist, often in books on the Theory of Games. The World Wide Web also has numerous helpful pages on these theorems.

\(^6\)Also called upper hemi-continuous in some texts.
Figure 10.8: The Intermediate Value Theorem says that a continuous function must cross a line that passes between two points that lie in the function’s range. The figure makes it obvious why this must be true.

**Theorem 2 (Brouwer).** Let \( f : S \rightarrow S \) be a continuous mapping from a non-empty convex, compact set \( S \subset \mathbb{R}^n \) into itself (see fig. 10.9). Then there exists an \( x^* \in S \) such that \( x^* = f(x^*) \).

The convex property is important here: if the upper and lower portions of the curve \( F \) were not connected and convex then the mapping would not necessarily intersect the 45 degree line and thus there would be no fixed point. (The set of rotations of a circle by irrational angles has no fixed point, since the set has gaps.)

In system administration we are often interested in procedures that map some input state to some output state. Arbitrary discrete sets of operators need not map convex sets of states. This places limits on what kinds of operator can have a stable outcome. One of the challenges of system administration is determining ways of *ensuring* that a system has a fixed point.

**Example 105.** Discrete operations do not necessarily have fixed points, but a continuum approximation created by a local averaging procedure can interpolate a non-convex set of operations into a convex one, allowing a virtual fixed point to be defined, on average. This is one of the main themes of this book. It is the essence of the maintenance theorem (see section 16.8) and it is a major reason for abandoning a strictly discrete formulation of systems.

Kakutani’s generalization of Brouwer’s theorem allows us to map not just points to points, but sets to sets, or points to sets. A *correspondence* is a mapping from a point to a set. This allows us to have not just stable points for simple mappings from point to point, but also in mappings that are formed by sewing together disjointed pieces into a consistent union.
Figure 10.9: Brouwer’s Fixed Point Theorem. At some point, a function that satisfies the convexity conditions must cross the diagonal line. Thus a mapping from one axis to the other must have a fixed point.

**Theorem 3** (Kakutani). Let \( F : S \to S \) be an upper semi-continuous correspondence, or mapping from a non-empty convex, compact set \( S \subset \mathbb{R}^n \) into itself (see fig. 10.10), such that, for all \( x \in S \), the set \( F(x) \) is convex and non-empty, then \( F(\cdot) \) (which can now have a multitude of values as it is a set) has a fixed point \( x^* \in S \) such that \( x^* \in F(x^*) \).

The importance of this theorem is in being able to identify fixed points even in functions that map sets that are more complicated than one-to-one mappings: we can talk identify regions in parameter spaces.

The theorem tells us that there is at least one point \( x = P \) within the values laid out along the axes that map that meets itself within the regions that are defined by the correspondence. In fig. 10.10, there is a range of values where the 45 degree line intersects the region that are fixed points. Thus we have a fixed subset.

The importance of this final theorem is that it applies to objects (sets) that are general enough to be able to describe policy in a continuum approximation. Although the figure illustrates this for a subset of \( \mathbb{R}^1 \), i.e. a single parameter, the regions can be multi-dimensional, so we can identify stable sub-spaces. It is seldom that we can identify a single ‘correct point’, but a correct region is a more likely prospect in a complex system of many variables. Note, however that these theorems apply to continuous functions, and thus apply to the continuum approximation of human-computer systems, which in turn applies on average. That is another reason why regions are important – we must allow for uncertainty. Discrete systems do not necessarily have stable fixed points.

**Example 106.** An important equilibrium for defining system policy is based on the notion of a fixed point of the graph of all rational strategy preferences (see below). This is not stability
Figure 10.10: Kakutani’s Fixed Point Theorem. The Brouwer theorem can be generalized to regions, provided the regions touch and have the same convexity properties. A point on the horizontal axis now maps to a range of values within itself indicated by the brace. Had the two regions not touched and been intersected, there the system would have been doomed to oscillate between the two regions and never stabilize.

of the system’s evolution, but stability of the choice of end point in the kernel under different policy decision criteria. This equilibrium is best known as the Nash equilibrium, or Kakutani fixed point (see [Mye91] for an excellent introduction) of the preference graph. It looks for a subset of states that can be regarded as a limit points of competing decision criteria. In a two-person zero sum game, this corresponds to the minimax solution that is used below. The idea of an ideal configuration $Q^*$ for a system can be defined as a fixed point of the ‘response matrix’ for mapping non-ideal states onto ideal ones. This matrix is readily defined in terms of convergent mappings ([Bur95, BR97]),

$$Q' = R(Q),$$  \hspace{1cm} (10.50)

However, the convergence property is not enough to select a stable base state for a convergent process, because convergence can be applied to any state. In order to prevent configuration loops and find the set of self-consistent fixed points that can be identified with policies, we must solve

$$Q^* = R(Q^*).$$  \hspace{1cm} (10.51)

This condition is the essence of the Nash equilibrium in Game Theory (see chapter 19).
10.14 Metastable alternatives and adaptability

If a system has a single global minimum, or stable end point, it will seek out this point and never emerge from it. Few systems benefit from being this stable, since changes in their environments usually demand a greater flexibility.

A system that can adapt to new conditions needs to be able to get out of its state of stability, by perturbing it with a large enough perturbation of an appropriate type. We would then like it to settle into a new and ‘better’ stable state that is more appropriate to the new conditions.

Systems that have some kind of symmetry (a choice that doesn’t matter), will naturally have several equivalent stable states, each of which is equally good. The archetypal model for such a system is a system that has several internally stable regions (fig. 10.11), with barriers in between. If we provide a sufficient perturbation, we can end up in a different minimum that is as good. This is called tunnelling.

Figure 10.11: Multi-stability means that there are several alternative stable states that are equally good. These are said to be symmetrical. As environmental conditions change, one of these states might become preferable to all of the others. Then the symmetry is said to be broken. To tunnel from one state to the next, we must jump out of the region of internal stability, and back into it in a different location.

This kind of multi-stability will not help the system to adapt, unless the criterion separating them suddenly becomes important in making one state better than the other. In this case we say that the symmetry has been broken.

Example 107. Consider the set of states for access permissions to a computer password database. In fig. 10.11, the first minimum is the end state in which read-only permissions are granted to all; in the second minimum read-write permissions are granted to all; in the third minimum, privileged
access is given to the administrative user. Now suppose that, initially, only a single administrative user has physical access to the database so that the permissions on the database are irrelevant. If conditions change, so that several users are introduced to the system, then the symmetry is now broken under the criterion of ‘security’, where security means access to restricted data.

The idea of metastability or multi-stability is connected to the notion of the kernel of a graph. The kernel consists of stable end-points of a system’s policies, or preference criteria. If conditions change, then we want to be able to break down the internal stability or allow transitions out of it, backtracking out and then transferring into a new state. If we are interested in adaptability, then it is wise to arrange for several of these alternative states to exist, so that a rapid change of policy can select the new ‘best choice’ without major architectural change.

10.15 Final remarks

Stability is a powerful concept, with many interpretations. If we cannot make a system stable over a timescale of importance to us, we can’t generally make it work for us at all. Thus stability is the first step towards a functional system. We have considered a few possible interpretations in this chapter, but this by no means covers the full repertoire. The stability of mappings, for instance, are quite general — they are generic behaviours that one would expect to find in any directed mapping and thus we can expect these concepts to emerge in a variety of situations.

Equilibria of games, discussed in chapter 19, are a natural extension of the idea of stability to decision making. In collaborative networks, such as peer to peer networks, cooperation or conflict between rational members of the collective can be modelled as persistent coalitions between individuals pursuing selfish interests. The concept of imputations, or coalition that strictly increase a player’s benefit arises in this case (see [Rap70] for a lucid introduction); \(\psi\)-stability can then be used to describe the likelihood for lasting cooperation.

Stability provides a set of concepts that we can draw on to analyze the behaviour of human-computer systems — both stochastic and some to rule based systems. Concepts of stability prove to be particularly important when defining the meaning of a sustainable policy.
Applications and Further Study 10.

- Characterizing and quantifying stability.
- Relating stability to predictability.
- Using stability as a basis for choosing arbitrary policy in a system.
- Looking for controllable or maintainable pathways of change.
CHAPTER 11

RESOURCE NETWORKS

This chapter considers the relationship between a system and its critical dependencies: the description of basic resources that permit the system to function. It was commented earlier that any system can be thought of as an information system, because anything that happens to its resources must be described with the help of information; thus one is never more than one step of abstraction away from talking about the change of information about objects, rather than talking about the objects themselves.

11.1 WHAT IS A SYSTEM RESOURCE?

The word resource is often used to describe the assets of a system or enterprise, which is often used by businesses to describe what things are valuable to the business. Resources or assets are the objects that describe the make-up of a system; they are also the freedom to change in the system. The word resources is preferable to assets, since it does not imply something that is automatically good.

Processes manipulate a system’s resources, leading to a change in their organization or perhaps to an exchange of one resource for another. It is the book-keeping or accounting of resources that is described by the dynamics of a system. In short, all systems follow a kind of ‘economy’, where the accounting parameters have names and properties that go beyond money and goods.

A whole plethora of words has been introduced to describe assets and resources. Some of them refer to tangible, physical items and others are more abstract qualities. Here are some examples.

- Raw materials (stock)
- Processing power (CPU or human)
• Storage space (memory or archive)
• Movement in space
• Time
• ‘Potential’ for reward
• Personnel
• Property (real estate)
• Intellectual capital
• Respect, status
• Privilege

Resource availability is something that results in both freedoms and constraints. The availability of space allows a system’s expansion; conversely, the limited size of a space is a constraint on what can develop.

There are two issues with resources: how resources change and how they are organized, e.g. do we have access to them? Another way of saying this is that there are both local and global issues. Resources can be modelled as a number of nodes within a network topology, where the network indicates the pathway for interaction between the resources.

### 11.2 Representation of Resources

We want to be able to talk about resources, their distribution and their usage formally. In mathematical or formal terms, a resource is a quantity can be measured about the system. A measurable must be represented as a variable that depends on parameters that describe its distribution and change with respect to time.

Let us separate space and time as resources, and think of these rather as parameters that are perhaps restricted by boundary conditions.

**Definition 44** (Resource variables). *The resources of a system are functions of time, space and other parameters that describe their distribution and patterns of change. They are written as variables of state*

\[ q_i(t, x, \ldots) \quad \text{for } i = 0, 1, 2, \ldots \]  

(11.1)
The parameter space $X$ is sometimes called the configuration space of the system, because it describes how resources are arranged or configured.

Resources take values that are measured in some form of currency (see section 4.9), and they represent the valuables of the system, both actual, potential, material and social. A set of resource variables describes a partial state of the system. The amount or availability of a resource is described by a value, measured in its own form of currency, which represents the balance of payments which led to the current state of resources.

A description of resources involves a set of values at various locations $q(t, x)$. The set of all such values, $\{ q(t_0, x) \}$, at a given moment $t_0$ is called a configuration of the system. Managing system resources includes managing their configurations over time.

11.3 RESOURCE CURRENCY RELATIONSHIPS

Resources come in many flavours, and depend on various location parameters. A resource can also have a value measured on any number of different currency scales. If one wishes to express a statement saying that a resource has a value with respect to more than one system of values, then one must specify a relationship between those values, i.e. the value systems are not independent. This applies to any relationship between values in the system.

Example 108. ‘Time is money’ is a functional relationship between a parameter $t$ and a value $M$ belonging to a state space consisting of all the possible values that money can take. It is written $M(t)$ saying that time and money are related by a formula, e.g. in the simplest case,

$$ M(t) \propto t = kt. \quad (11.2) $$

This example is often quoted frivolously, but it expresses an economic truth about systems, in which the constant of proportionality $k$ is the the average sum of money that can be earned by the system per unit time. Since human-computer systems are frequently driven by economic interests, this kind of relationship will be used frequently, and will feed into models of the system.

Example 109. Consider social scales which drive the human components in a system. Intellectual capital is a potential for innovation within a human system. If intellect leads to respect and status, it could also lead to privilege or even money. Each of the emphasized words is a social value system. By identifying a relationship between them above, we are saying that they are all related. Thus, we might choose to measure intellect by IQ, but if intellect leads to respect, then respect must be...
measured on a scale which is a function of intellect (and other inputs), calibrated to the respect scale. Similarly, if this leads to status, then status must be a function of respect (and other inputs) calibrated to the status scale. This dependency chain continues.

In practice, particularly in the West, human social systems are organized around money, and all other scales can be measures in terms of money, according to some elaborate formula. While this is certainly a cynical view of things, and somewhat of an oversimplification, it is a pragmatic reality precisely because it leads to a concrete, measurable value for what is happening in society. A similar situation exists in computer systems, where most measurements can be associated with a certain amount of processing time, or memory consumed.

In making informed decisions about human-computer systems, we are often forced to make value judgements of the kind described above. The key to making rational, rather than ad hoc, decisions lies in quantifying that value system in relation to the other valuables in the equation.

**Example 110.** Suppose we are interested in implementing a security system for online banking (see section 19.10). A security consultant has determined the relative security of different available technologies and how much they will cost to implement, but how do we decide whether it is worth paying more money for a better solution or not? We can do this by defining the ‘payoff’ or utility of the solution, and define it by

\[ \text{Payoff (\(\Pi\))} = \text{Security evaluation (\(S\))} - \text{cost of implementation (\(C\)).} \]

At this stage, however, security and cost of implementation are measured using quite different scales, so we need to relate them to one another. Let us measure payoff on a scale from 0 to 100, and the security evaluation on the same scale. The cost of implementation can be measured in Euros, so we must relate Euros to security level. Let us suppose that the cost/security ratio is constant (“you get what you pay for”); then:

\[ \Pi = S - f(C) = S - kC, \]  \hspace{1cm} (11.3)

for some constant \(k\). There is no rational way of determining \(k\). It is a human value judgement: determining it must be a part of policy. Policy necessarily intrudes on the rational process through the need for certain ad hoc judgements.

### 11.4 RESOURCE ALLOCATION, CONSUMPTION AND CONSERVATION

Resources are both the machinery and the fuel in a system; thus they fall into two categories.
Chapter 11. Resource Networks

Definition 46 (Reusable resources). These are resources which are allocated temporarily to a particular task. Once they are no longer needed, they can be passed on to another task. Reusable resources include people, computers, storage, and communications lines.

Example 111. A car park (parking lot) consists of a number of parking spaces, which become occupied for a certain time and then are freed for re-use. A computer disk has a number of sectors which are used to store data for a certain time and are later freed for re-use.

Example 112. Shared infrastructure of all kinds is a reusable resource. If we view a state of continuous operation as a business objective, in a cloud infrastructure service (IaaS), then we can imagine container or virtual machine slots to be like reusable parking spaces for jobs. This is like air traffic control slots. Although one can never recover a particular moment in time, the progression of time itself becomes an irrelevant variable in continuous operation. As soon as a slot becomes empty, it can be used by a new job, continuing the renewal process.

Definition 47 (Consumed resources). These are resources which can only be used once. Once they are used, they disappear from the resource pool. The resource pool can be refilled, if a new quota of resources becomes available.

Example 113. Absolute time is a resource which cannot be re-used. Money, electricity, oil, write-once memory and damaged equipment are other examples.

Example 114. Permanent storage is a consumed resource. It suggests that we need to carry out some kind of garbage collection to prune useless data that unintentionally gets used up.

In the physical world, resources do not simply disappear or get used up; rather, they are converted from one form into another, or bound in some role, where they cannot be freed for re-use. In practice, however, it is convenient to think of resources as disappearing from the pool of resources. One says that the system is an open system. The mathematical expression of conservation, as expressed in the continuum approximation is given by the formula:

\[ \frac{\partial}{\partial x^i} J^i = - \frac{\partial \rho}{\partial t} \]

(11.4)

for some vector \( \vec{q}(t, x^i) = (\rho, J^i) \). It expresses the idea that, if there is regional conservation in a system, the density of a system property \( \rho(t, \vec{x}) \) cannot change except by spreading out into a current or flow \( J^i(t, \vec{x}) \) from that location (see for instance [Bur02a]).

Another consequence of open systems is that the distribution of resources is not always under the control of the system itself. Some resources are distributed at random, or by external agents.

Example 115. The number of passengers arriving at a bus stop is not determined by the bus transport system, but the arrival of buses is. The number of E-mail messages arriving at a computer

...
system is not determined by that system, but the availability of CPU and memory to handle the requests is.

The problem of utilizing available resources which are under local control involves two issues: allocation and reclamation.

**Definition 48.** Resource allocation is the association or assignment of a resource from a shared pool to a process within the system. The concept of ownership of a resource is achieved by attaching an extra parameter to the tuple $X$.

**Example 116.** When a process is defined, it is built up of an assembly of resources.

**Definition 49.** Resource reclamation is the freeing of a reusable resource back into a shared pool.

**Example 117.** When a process is terminated, the resources that were attached to it are returned to a state of disuse. When cars leave a car park (parking lot), the spaces become available for others to use, i.e. the spaces are not reserved from private use. Disk space that is allocated by quota to a given computer user does not become free for other users when it is no longer used by the private user.

### 11.5 Where to Attach Resources?

A system does not function without resources. Resources come in may forms, both literal (e.g. processor time, memory, floor space, raw materials) and abstract (e.g. goodwill, permission, credit). Materials, workforce, creativity, tools and equipment are all resources. We describe these resource by variables $q(t, A)$ that vary in time and address.

For a system architecture to work optimally, resources must be made available at the right places and times. Once we know where the resources are needed, this becomes a scheduling problem. Methods for approaching these matters have been considered in the previous chapters.

- Have the resource requirements been targeted correctly?
- Will resources reach their target or disperse before arriving?

In order for resources to be available to other parts of the system, there has to be a route connecting them to their point of consumption. Resources may be provided by direct injection, or by controlled delivery through a mediator (a metaphorical or literal valve or regulator). There are often administrative overheads involved with regulation.

Direct access to resources allows maximum efficiency, but sometimes resources have to be transported by an unreliable channel that is shared by several tasks.
Figure 11.1: A simplified functional dependency structure diagram for a university faculty department. The level of dependency increases with distance down the page. Note that, while the goal of the department is to furnish society with knowledge, the dependencies point to the staff as being the central element on which almost everything above depends. This indicates that investment in this resource is important to the system. The dependency chart continues downwards, of course, with food, power, and so on. A formal method for this analysis is given in section 6.5.

**Example 118.** The monitoring of resource usage (e.g. system accounting, or bureaucracy) can result in inefficiency, because it relies on sharing the same resources as the system itself. Thus it becomes overhead which takes resources away from the main purpose of the system. We must decide whether this overhead is worth the loss of resources.

**Example 119.** Network topology alone can offer guidance to where resources are likely to be needed by looking at how the probability of resource utilization has hotspots. Graph theoretical considerations may be of help here. See section 11.7.

The distance from resources supply to point of usage should be minimized in order to minimize delay and other losses. Food, power and basic freedoms are at the bottom of any system involving humans or machines. If access to these is restricted or constricted, the system performance will suffer. There are two extremes for injecting resources:

- **Bottom up:** Resources are inserted directly at the location within the functional tree, closest to the action (direct access, but no control or accountability)
- **Top down:** Resources are inserted at the top of the functional tree: disseminated by per-
colating down from above (has to pass through many junctures - this is inefficient, but accountable)

If two processes share a common dependency, resources can be inserted at the dependency for maximum efficiency, i.e. we can make the dependency a formal service that supplies resources to the processes. This is a form of *system normalization* (see chapter 14).

### 11.6 Access to Resources

The ability for information and resources to spread through a system depends on the availability of free channels of communication within the system. Percolation is the phenomenon that occurs when a fluid manages to penetrate a porous medium. For a network, the term is used in the following way.

**Definition 50** (Percolation transition). A network is said to percolate, if it is possible to reach any node from any other by at least one route.

This means that information or resources can be communicated to the parts of the system where they are needed; conversely, it might mean that an attacker can reach any part of the system from any other. Either way, the percolation transition in any network is important to understand. It tells us both about security and availability.

Since connectivity has such basic ramifications for a system, it is of considerable interest to be able to measure it. This is a particularly difficult task in large organizations where we might have only partial information about. In a dynamical system pathways of communication open and close at different times, perhaps for security reasons, and perhaps for efficiency reasons.

Since it is not always possible to measure a system completely or directly, we are interested in gauging the *probability* for percolation in two distinct cases:

- **Perfect information**: if we know precise details about the graph of a system, it is possible to work out whether there is a route connecting any two nodes. The All Pairs Shortest Distance Matrix, as defined by standard algorithms by Floyd or Dijkstra ([CLRS01, BG99]), for instance, is an example algorithm for evaluating the possibility for communication between nodes in a graph. Another measure is based on the connectivity comes from asking how many pairs of nodes, out of all possible pairs, can reach one another in a finite number of hops. We thus define the ratio \( R_C \) of connected pairs of nodes to the total number of pairs that could be connected ([BCE04a]):

\[
R_C = \sum_{i=\text{clusters}} \frac{\binom{n_i}{2}}{2N(N-1)} = 1. \tag{11.5}
\]
This is simply the criterion that the graph be connected. Normally it is only possible to evaluate this quantity for theoretical models, or for very small organizations under tight control, with perfect information, so we need other ways of estimating percolation with only partial information.

- **Partial information**: a system administrator is not always aware of every detail of the system or its users. Real systems are inherently probabilistic. In human-computer systems, especially, there is the possibility of covert channels of communication that link together parts of the system in a non-obvious fashion. For instance, a married couple working in different fragments of a system could easily leak information to one another. Conversely, an artificial barrier between nodes might be introduced by sickness or accidental disconnection of a node. In such cases, we must admit to possessing only incomplete or probabilistic information about a system and make do with an estimate of the likelihood for percolation.

In both cases, there are methods for determining the likelihood of complete connectivity within a part of the system.

In Appendix C, the results for percolation thresholds of approximately random graphs are derived. These are based on the work of [NSW01] for huge random graphs, and were adapted for small graphs in [BCE04a]. Random graphs are a useful measuring stick for actual graphs because they tend to percolate very easily, with only a small number of connections. They are very efficient at covering the nodes with available routes.

A connected part of a graph is called a *cluster*. We would like to find the sizes of clusters in a graph and see when they become large, i.e. of the same order of magnitude as the size of the graph. The giant component or cluster is thus defined to be a cluster that is of order \( N \) nodes. If such a cluster exists, then other smaller clusters of order \( \log N \) might also exist (see [MR98]). The large-graph condition for the existence of a giant cluster (of infinite size) is simply

\[
\sum_k k(k-2) p_k \geq 0. \tag{11.6}
\]

Here the sum is over \( k \), the degrees of nodes in the graph, and \( p_k \) is the probability of finding a node of degree \( k \) in the graph, i.e. it is the number of nodes of degree \( k \) divided by the total number of nodes \( n_k/N \) (see section 6.2).

This provides a simple test that can be applied to a human-computer system, in order to estimate the possibility of complete penetrability. If we determine only the \( p_k \), then we have an immediate machine-testable criterion for the possibility of a systemwide security breach, or efficient transmission.

The problem with the above expression is that it is derived under the assumption of there being a smooth differentiable structure to the average properties of the graphs. This is really only a good approximation in the infinite graph limit, so it is mainly of interest to huge systems like the entire
Internet. For a small graph with \( N \) nodes, the above criterion for a giant cluster is inaccurate. Clusters do not grow to infinity, they can only grow to size \( N \) at the most, hence we must be more precise and use a dimensionful scale rather than infinity as a reference point. For a small graph the size of a giant cluster is \( N \) and the size of of below-threshold clusters is \( \log(N) \) (see [MR98]). An improved criterion was found in [BCE04a], and is given by:

\[
\langle k \rangle^2 + \sum_k k(k-2)p_k > \log(N).
\] (11.7)

This can be understood as follows. If a graph contains a giant component, it is of order \( N \) and the size of the next largest component is typically \( O(\log N) \) (see [MR98]); thus, according to the theory of random graphs the margin for error in estimating a giant component is of order \( \pm \log N \). We thus add the magnitude of the uncertainty in order to reduce the likelihood of a false positive conclusion.

*Ad hoc networks* are dynamically *random graphs* in which connections are initiated and broken in a non-deterministic way. The above criteria are useful for estimating the penetrability of mobile and other ad hoc networks, on the fly.

## 11.7 Methods of Resource Allocation

Since resources are crucial to the ability for a system to function, resource allocation is a critical dependency. There are two sides to the problem: one is the organization of resources given to a single process, the other is how several processes can co-exist within shared resources. There has to be a pool of resources that acts as a constraint on the operation. Methods of assignment often need to make certain compromise.

**Example 120.** Suppose you need to draw a complicated map on a small piece of paper. How will you allocate the space on the page?

Since a dynamical system requires the flow of work or information, restrictions on the flow can limit the performance of the system. Time resources must be assigned to process incoming information to maintain the flow. Mismatches of flow rates between parts of the system can be a problem. If one part of a system has more resistance to the flow of work than another, it becomes a bottleneck that delays the entire collaborative network.

**Example 121.** Many systems employ resources to monitor the usage of other resources, e.g. bureaucratic controls or SNMP monitors. Too much monitoring or verification communication ties up resources which could be used to do useful work. In such a case, a resource becomes an overhead.
11.7.1 LOGICAL REGIONS OF SYSTEMS

Systems are well described by graphs, and graphs have properties that allow them to be classified into regions. A fitness landscape view of a graph can be created using eigenvalue centrality to define the ‘height’ of any node above an imaginary sea level or null point of zero centrality. Local maxima, the tops of the mountains, in this landscape provide a notion of maximum local importance. The nodes that are connected to these local maxima then define a region connected to that local centre. If we trace regions by travelling outwards from a local maximum and find nodes and meet a node in between two maxima, then it lies in a valley between the tops, and we can call it a bridge between regions. Thus we classify regions into maxima, bridges and regions. This will prove to be widely useful tool for identifying distinct regions in systems.

11.7.2 USING CENTRALITY TO IDENTIFY RESOURCE BOTTLENECKS

The graph theoretical technique above allows us to identify the places in a network where work flow resources are most crucial to the continued functioning of the system. Eigenvector centrality (see section 6.5) is a relatively easy and straightforward way of identifying the confluences of information flows in a network. It can be applied with various levels of approximation to take account of different aspects of the system. Let us consider how to identify the ‘hot-spots’ in a network, i.e. the places where resources will need to be invested in order to provide optimum fuel for the system.

Let the graph of the systems \((X, \Gamma)\) with configuration space \(X\) and arcs \(\Gamma\) be a representation of any aspect of a system involving association.

---

**Figure 11.2:** The classification of a graph into regions and bridges, using eigenvalue centrality as a sort criterion.
1. Draw a graph of the human-computer system, with all information flows or associations represented as arcs between the nodes.

2. For each connected fragment of the graph, construct the symmetrical adjacency matrix for the non-directed graph, setting a constant value (e.g.) 1 for a connection and a zero for no connection.

3. Calculate the principal eigenvector of the adjacency matrix, i.e. the eigenvector belonging to the highest eigenvalue.

4. Normalize the elements so that the maximum value is +1.

5. We now find the regions and bridges as described in section 11.7.1.

Each of the elements in the principal eigenvector now rank the importance of the nodes in the graph to workflow.

Example 122. Consider the human-computer system for Internet commerce depicted in fig. 11.3. This graph is a mixture of human and computer elements: departments and servers. We represent the outside world by a single outgoing or incoming link (node 5). The organization consists of a web server connected to a sales database, that collects orders which are then passed on to the order registration department. These collect money and pass on the orders to order processing...
who collect the orders and send them to dispatch for postal delivery to the customers. A marketing
department is linked to web server through the system administrator, and management sits on the
edge of the company, liaising with various staff members who run the departments.

Let us find the central resource sinks in this organization, first assuming that all of the arcs
are equally weighted, i.e. contribute about the same amount to the average flow through the
organization. We construct the adjacency matrix, shown in eqn. (11.8):

\[
A_{ij} = \begin{pmatrix}
0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
1 & 0 & 0 & 0 & 0 & 1 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
1 & 0 & 0 & 0 & 0 & 1 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 1 & 0 & 1 & 1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & 1 & 1 & 0 & 0 & 0 & 1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 & 0 & 1 & 1 & 1 & 1 & 1 & 0 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 \\
0 & 0 & 0 & 0 & 0 & 0 & 1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 1 & 1 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\end{pmatrix}
\]  

(11.8)

Some elements are marked bold here, for later removal. The eigenvalues of this matrix are now
computed and the transposed principal eigenvector is

\[
\vec{v}^T = (0.29, 0.49, 0.54, 0.75, 0.21, 0.89, 1.00, 0.58, 0.69, 0.69, 0.28, 0.28, 0.28, 0.55)
\]  

(11.9)

Node 7 is clearly the most central. This is the system administrator. This is perhaps a surprising
result for an organization, but it is a common situation where many parts of an organization rely
on basic support services to function, but at an unconscious level. This immediately suggests that
system administration services are important to the organization and that resources should be
given to this basic service. Node 6 is the next highest ranking node; this is the order registration
department. Again, this is not particularly obvious from the diagram alone: it does not seem to be
any more important than order processing. However, with hindsight, we can see that its importance
arises because it has to liaise closely with all other departments.

Using the definitions of regions and bridges from section 11.7.1, we can redraw the graph
using centrality to organize it. The result is shown in fig. 11.4. The structure revealed by graph
centrality accurately reflects the structure of the organization: it is composed largely of two
MARKETING ORDER PROCESSING

Figure 11.4: A centrality self-organized graph showing the structure of the graph centred around two local maxima or ‘most important’ nodes, that are the order registration department and the system administrator. There are also 4 bridge nodes and a bridging link between the regions.

separate enterprises: marketing and order processing. These departments are bound together by certain bridges that include management and staff that liaise with the departments. Surprisingly, system administration services fall at the centre of the staff/marketing part of the organization. Again, this occurs because it is a critical dependency of this region of the system. Finally the web server is a bridge that connects both departments to the outside world — the outside hanging on at the periphery of the systems.

The results of the example above could be enhanced by weighting the different connections in the adjacency matrix to reflect the work load. In a more realistic graph, one would measure the volumes of communication between different nodes and create a more detailed model. The inclusion of every person and computer as a separate node would then automatically provide the appropriate weighting.

Resource centrality is a powerful method for identifying stable regions of resource networks, as a guide to the appropriate investment of resources. Taking account of the directed nature of a graph can also affect the identification of regions (see section 6.6).
11.8 DIRECTED RESOURCES: FLOW ASYMMETRIES

As noted in section 6.6, the centrality of directed graphs is much harder to interpret; indeed, the concept falls apart for graphs that do not have enough return paths to make a self-consistent picture. As long as there is a well-defined principal eigenvector, we can use directedness in the centrality method to find important hot-spots in systems.

Consider the direction of the flows in fig. 11.3, and let us suppose that some of the flows are directional, as in fig. 11.5. The adjacency matrix is now made asymmetrical, by setting to zero the bold connections in eqn. (11.8).

Both $A$ and $A^T$ have a principal eigenvector, with the following ranking:
### Applications and Further Study 11.

- Organizing and planning the deployment of system resources.
- Formalizing the economic aspects of a system, both for humans and machines.
Chapter 12

Task Management and Services

Never fear the event
– Horatio Nelson (1801)

In previous chapters, we considered how resources fit into the scheme of a system. Now we consider how to optimize their allocation, according to quotas or allocation criteria. The resource we focus mostly on is time, since this is the fundamental currency underlying all dynamical systems, however this chapter is really an extension of the previous chapter on resource usage. Time allocation applies both to humans and computers and affects a strong dependency on the ability of the system to process information.

12.1 Task List Scheduling

Scheduling is a way of parsing a tree of tasks as efficiently as possible. The techniques for scheduling are well known from parallel processing and operating system design ([KN84, AD96]). An important aspect of configuration management is how management operations are scheduled within a system, both in response to specific events and as a general matter of the maintenance of the system.

Example 123. A service help-desk receives calls at a peak rate during the middle of the day, in its time zone, and at a slower rate at other times. The number of jobs, with a certain level of difficulty, follows a rough pattern. The number of available case handlers is also maximal during the day, but the numbers are not quite certain due to possible sickness. What is the optimum approach to completing all the incoming tasks?
Example 124. A list of routine maintenance operations, including system backup, security checks and software updates, must be completed each day to satisfy policy. Which ordering of tasks causes the least disruption to normal activities and leads to the most up-to-date, best maintained system.

Scheduling clearly encompasses precise comparisons and heuristic value judgements. Specifying a schedule naturally becomes a part of any policy.

Scheduling takes many forms, such as job-shop scheduling, production scheduling, multiprocessor scheduling, human time management and so on. It can take place within any extent of time, space or other suitable system parameter. The two main classes of scheduling are dynamic and static scheduling. Dynamical schedules can change their own execution pattern, while static ones are fully predetermined. In general, solving static scheduling problems belongs to the class of NP problems (it is presumed to be computationally intensive).

If we represent scheduling as a graph theoretical problem, it involves assigning the vertices (task nodes) of an acyclic, directed graph onto a set of resources, such that the total time to process all the tasks is minimised. The total time to process all the tasks is usually referred to as the makespan. An additional objective is often to achieve a short makespan while minimising the use of resources. Such multi-objective optimisation problems involve complex trade-offs and compromises, and good scheduling strategies are almost based on a detailed and deep understanding of the specific problem domain.

Most approaches belong to the family of priority-list scheduling algorithms, differentiated by the way in which task priorities are assigned to the set of resources. Traditionally, heuristic methods have been employed in the search for high-quality solutions ([KN84]). Over the last decade heuristics have been combined with modern search techniques such as simulated annealing and genetic algorithms (see [AD96]).

12.2 Deterministic and Non-deterministic Schedules

There is a distinction between the arrival of a random event (a stochastic task such as a telephone call or a server request) and a planned or structured task with long term predictability, (such as periodically executed tasks like daily cleaning, regular software updates, system backups). This is reflected in two possible ways of initiating a system task:

- A random starting point in the system graph.
- A random arrival time at some point in the graph.

Whichever method is used to parse the graph of tasks, the result of the process must end up with \( n \) ordered lists, to be carried out by \( n \) ‘servers’. A server might be a person, a computer or an enterprise.
CHAPTER 12. TASK MANAGEMENT AND SERVICES

EVENT HANDLING

System administration, employing agents or software robots, is an administrative method that potentially scales to large numbers of hosts in distributed systems, provided each host is responsible for its own state of configuration (see [BC03]). However, the inter-dependencies between networked computers makes cooperation essential, and the distributed nature of the system makes the timing of events, for all intents and purposes, random.

Example 125. Policy based configuration languages associate the occurrence of specified events or conditions, with responses to be carried out by an agent. Cfengine accomplishes this, for instance, by classifying the state of a host, at the time of invocation, into a number of classifiers. Some of these represent the time of invocation, others the nature of the environment, and so on. For example:

files:
(linux|solaris).Hr12::
/etc/passwd mode=0644 action=fixall inform=true

The class membership is described in the second line. In this instance, it specifies the class of all hosts which are of type Linux or Solaris, during the time interval from 12:00 hours to 12:59 (Hr12). Tasks to be scheduled are placed in classes which determine the host(s) on which they should be executed, or the time at which they should be executed. Actions are placed in such classes and are only performed if the agent executes the code in an environment which belongs to one of the relevant classes. Thus, by placing actions in judiciously chosen classes, one specifies actions to be carried out on either individual machines or on arbitrary groups of machines which have a common feature relating them. This is a scheduling procedure. We thus have:

- Scheduling in time.
- Scheduling by host attribute (location, type, etc).

TASK SCHEDULING

A set of ordered tasks or precedences can be represented by a directed graph, \((V, E)\), with vertices, \(V\), and directed edges, \(E\). The collection of vertices, \(V = \{v_1, v_2, ..., v_n\}\), represents the set of \(n\) tasks to be scheduled and the directed edges, \(E = \{e_{ij}\}\), define the precedence relations that exist between these tasks (\(e_{ij}\) denotes a directed edge from task \(v_i\) to \(v_j\)). The graph might contain loops or cycles. These can cause unwanted repetition of tasks by mistake. Edges can be removed and the graph to convert it into an acyclic graph, or spanning tree that avoids this problem.
The task management process can be understood as scheduling in several ways. A graphical representation allows modelling of task management (see fig. 12.1).

Within a single set of policy rules there is a set of schedulable tasks that is ordered by precedence relations (arrows in the graph). These relations constrain the order in which policies can be applied, and thus how the graph has to be parsed.

A second way in which scheduling enters, is through the response of the system to arriving tasks. Should the server agents activate once every day, hour or minute, in order to check for scheduled tasks, or immediately; should they start at random times, or at predictable times? Should the policies scheduled for specific times of day, occur always at the same times of day, or at variable times, perhaps random.

Finally, although scheduling is normally regarded as referring to extent over time, a distributed system also has two other degrees of ‘spatial’ extent: $h$ and $c$. Scheduling tasks over different hosts, or changing the details of software components is also a possibility. It is possible to confound the predictability of software component configuration to present a ‘moving target’ to would-be attackers (see [BS01, San01]).

### 12.3 Human-Computer Scheduling

Configuration management is a mixture of a dynamic and static scheduling. It is dynamic in the sense that it is an ongoing real-time process where policies are triggered as a result of the environment. It is static in the sense that all policies are known a priori during any work interval. Policies can be added, changed and removed arbitrarily in a dynamical fashion. However, this does not really interfere with a static model because such changes would typically be made during a time-interval in which the configuration tool were idle or offline (in a quiescent state).

The separation of time-scales is an issue that we harp on continually in this book. It is important
here too. The management of time can only be successfully prosecuted by separating independent
time scales from one another. Using the method of local averaging (see appendix B) can be used to
identify time scales over which average behaviour is only slowly varying. These are the independent
levels of scheduling.

Example 126. A simple analogy is helpful for understanding this. Suppose we are interested
planning the time of an enterprise that deals with selling furniture on the Internet (see fig. 11.3).
The timescale at which products are redesigned is on the order of a year, so we expect to have
to update web pages on this time scale. We must therefore allocate a number of man-hours per
year to this task. (It is not necessary to check this every few minutes.) However, the time scale on
which sales requests arrive is about one per hour, so we need to allocate sufficient order processing
power on the scale of an hour to check and process these orders. The number of web page accesses
is on the order of several hits per second, so a the web server must schedule resources to cope with
this demand.

Example 127. In modern shared infrastructure, known as cloud computing, the scheduling of jobs
and memory, network, and processing resources is one of the key problems to solve. Open source
schedulers[SKAEMW13] like Openstack, Kubernetes, Mesos and associated modules take on the
task of trying to place jobs in a network with specific topology to as to reduce the key variables of
latency, traffic intensity, and collisions. It never pays to be sending data over a shared network
if that can be avoided, so scheduling of unknown demand across a fixed supply, with redundancy
and sufficient overcapacity for handling peak demand is a problem one must revisit continually as
systems and workloads evolve.

12.4 Service provision and policy

In networks and systems alike, we are interested in maintaining predictable levels of service.
Service providers would, after all, like to sell these services to customers and thus need to be able
to offer guarantees about what will be delivered and what service will cost. This brings up two
issues:

- **Quality of Service (QoS):** Quality of Service a goal of all service providers, whether the
  service is a network transmission rate or a system up-time level. Although one often
  has the impression of service quality as being a solved problem, this is far from the case.
  Guaranteeing service levels requires one to address the uncertainties in service provision at
  all levels, and these are often complex.

- **Service Level Agreements (SLA):** A Service Level Agreement is a contract of service levels
  that is offered to a customer by a provider. The provision of service cannot be guaranteed
with complete certainty, as we shall see below, so Service Level Agreements are above determining acceptable margins of behaviour in the system, and what recompense will be offered to a customer if the levels are not met.

A strategy for meeting service level guarantees is over-provision to provide a margin that can absorb sudden demand. Clearly, no system can absorb any sudden change in demand, thus maximum limits are placed on expectancy.

12.5 QUEUE PROCESSING

Queues are an important model of load handling in any system, whether human or machine. Whenever there is a confluence of information at some point in a system, there is a serialization of processing that bottlenecks information into a queue. In stochastic systems and queueing theory, these processes are called birth-death processes. In physics, they are referred to as creation-annihilation processes or prey-predator models.

Although a vast amount of research has gone into studying queues [Kle76, GH98, SNO71, Coo90, Wal90, Gun93, Gun08, GPT15], we know that queues have basically two states. They are either lightly loaded and working well, or they are completely jammed and the system is ‘thrashing’ to keep its head above water. It is enough to gain an appreciation of the simplest models to get the basics of queueing behaviours.

Queueing networks are representations of resource consuming systems on graphs, such that information or work flows from node to node, often with cyclic repetition. A queue is a generic processing model: jobs arrive as events, or they are pre-allocated and some form of human-computer processing must be applied to eliminate them from the ‘to-do’ list.

Queueing models have been devised with all levels of complexity (see [Jai91] for an introduction); they are statistical models that describe the steady state properties of stochastic task systems. At the coarsest level of approximation, mean value theory can be used to obtain order of magnitude estimates of processing efficiencies. Queueing models consist of a number of choices or parameters:

- **Arrival time distribution process**
  
  If service requests occur at times \( t_1, t_2, \ldots \), the values of the random variable \( \tilde{t}_i = t_i - t_{i-1} \) are called the inter-arrival times of the process. It is commonly assumed that these form a Poisson (exponential) distribution; however, many network processes have long-tailed distributions (see section 10.10).

- **Process time distribution**
  
  The time that each client is engaged in requesting a service.
• **Number of servers**
  The number of humans, computers or other entities responsible for processing requests. The work-rate is a function of this number.

• **System processing limit**
  If the system has a maximum throughput, or processing rate, this limits the behaviour of the system as a whole.

• **Maximum population size**
  The maximum number of clients that can ask for services.

• **Scheduling policy**:
  Various queueing policies are used to try to empty the queue as quickly as possible. The most common is First Come First Served (FCFS) which is a first-in-first-out (FIFO) structure. Round-robin scheduling is a way of sharing time between multiple jobs. Shortest Job First picks the task whose estimated completion time is least. This potentially suffers from *starvation* of some tasks, i.e. long jobs never get executed because resources are saturated with an influx of new small jobs. Shortest Remaining Time First is a variation on the previous policy. Clearly, there are many possible strategies for processing requests. In a human context, a policy that is often used is Loudest Voice, First Served, i.e. those clients who make the biggest nuisance of themselves are disposed of quickly. The efficiency of these different policies is often difficult to evaluate, and depends on the nature of the system. In general, experimental analysis is required to determine an appropriate choice, or combination of choices.

### 12.6 Models

Queues are denoted in Kendall notation in the form \( A/S/c/(B/K/P) \), where \( A \) is the inter-arrival distribution that usually takes one of the following values:

- \( M \)  Memoryless (exponential/Poisson)
- \( E_k \)  Erlang with parameter \( k \)
- \( H_k \)  Hyper-exponential with parameter \( k \)
- \( D \)  Deterministic
- \( G \)  General

A deterministic distribution has constant inter-arrival times, with no variance, and ‘general’ means that the model’s results apply for any distribution. The term *memoryless* implies that the arrival process is a steady state process, in which the current state of the distribution does not depend on
what happened in the past: i.e. if the arrival time has a given form now, then it will have the same form for all subsequent arrivals—i.e. it is statistically stable.

$S$ is the processing time distribution, with the same values as for $A$. $c$ is the number of servers or service entities. $B$ is the number of buffers (the system processing limit), $K$ is the maximum population size and $P$ is the policy, if these are specified.

**Example 128.** The basic queue in section 12.7 is referred to as an $M/M/1$ model.

The Machine Repair-man Model ([Sch67]) is a simple queueing model that considers the problem of assigning the repair of machines to a repair queue. When a machine breaks, it is put into a queue until a repair-man can service it.

The Central Server Model ([Buz73, Buz76]) is an event driven model, with polling. A central server schedules a visit to a device (i.e. it polls a number of devices). If a request is pending, it is serviced; after polling is finished, the device returns to processing other tasks until the next event.

**Example 129.** The Simple Network Management Protocol (SNMP) uses essentially the Central Server Model. When an event or ‘trap’ occurs from a monitored device, a manager can poll the devices in the network and attend to any configuration changes in turn.

### 12.7 The Prototype Queue M/M/1

At a service centre (server), the incoming traffic has to balance with the outgoing traffic, or a queue will grow. In the worst case, incoming jobs will have to be dropped.

Suppose we have an average of $n - 1$ tasks already in a queue and tasks are arriving at a rate of $\lambda$ per second, then as soon as there are $n$ tasks in the queue some tasks must be forwarded at rate $\mu$, otherwise the average number of tasks in the queue will not stay constant. We can write this

$$\lambda p_{n-1} = \mu p_n,$$

$$p_n = \rho p_{n-1},$$

for any $n$, where $\rho = \lambda/\mu$. $\rho$ is called the traffic intensity. If $\rho > 1$ then the incoming rate is higher than the outgoing rate. Now since the recurrence relation above holds for all $n$, clearly

$$p_1 = \rho p_0$$

$$p_2 = \rho^2 p_0$$

$$p_n = \rho^n p_0.$$

The sum of probabilities is always 1, so

$$\sum_{n=0}^{\infty} p_n = 1.$$
This is a geometric series, so we can find $p_0$:

$$\sum_{n=0}^{\infty} p_n = \frac{p_0}{1 - \rho} = 1.$$ 

Thus we have, for any $n$,

$$p_n = (1 - \rho)\rho^n.$$ 

Although this is clearly an idealization (there is never an infinite number of tasks, even on the whole Internet), this gives us a distribution that we can use to estimate the average number of tasks in the queue. The expectation value of the number of tasks is:

$$E(n) = \langle n \rangle = \sum_{n=0}^{\infty} n p_n.$$ 

Substituting for $p_n$,

$$\langle n \rangle = \sum_{n=0}^{\infty} n(1 - \rho)\rho^n = \sum_{n=0}^{\infty} n\rho^n - \sum_{n=0}^{\infty} n\rho^{n+1}.$$ 

Relabelling $n \rightarrow n + 1$ in the second term gives us another geometric series:

$$\langle n \rangle = \sum_{n=0}^{\infty} n\rho^n - \sum_{n=1}^{\infty} (n - 1)\rho^n = \sum_{n=1}^{\infty} \rho^n = \frac{\rho}{1 - \rho}.$$ 

Thus the mean number of tasks is

$$\langle n \rangle = \frac{\rho}{1 - \rho} \rightarrow \infty \quad (\rho \rightarrow 1)$$

The variance can also be worked out

$$\langle (n - \langle n \rangle)^2 \rangle = \frac{\rho}{(1 - \rho)^2}.$$ 

This gives us an estimate of the size of queue that we need in order to cope with a normal traffic rate.

The system is busy whenever there is at least one job in the system. We can use this to characterize how busy a server is.
Definition 51 (Load average or Utilization). The probability of finding at least one job in the system is called the load average. It is ‘NOT’ the probability of finding no jobs in the system, i.e. $1 - P(n = 0)$.

For the M/M/1 queue, we thus write the load average, also called the Utilization, as

$$U = 1 - p_0 = 1 - (1 - \rho) = \rho = \frac{\lambda}{\mu},$$  \hspace{1cm} (12.1)

The assumptions above, that a fixed steady state (memoryless) rate equation holds, imply that the arrival times are Poisson distributed. However, research over the last ten years has revealed that Internet traffic does not satisfy this pattern. Voice traffic on the telephone system has always been well modelled in this way, but packet switched traffic is “bursty” – it has no scale that can be averaged out.

Example 130 (Sharing and contention). Some resources can only be used by one client at a time. If several clients try to use these simultaneously there is contention. Is the solution to put these requests in a queue, with some scheduling policy, or to make them try again later? A scheduling of requests requires a protocol or even a service to exist, to manage the queue. Without this, clients will compete and contend for the resource. To fully appreciate the nature of contention, we need to combine queueing theory with contention theory. See section 19.9 for a more complete method of analysis.

System loads which does not even out over reasonable time scales is said to be long-tailed and sometimes statistically self-similar (see section 10.10). This is often associated with a ‘burstiness’ or a power-law clustering of events. In self-similar traffic there is long-range dependence, and theoretically infinite (at least unpredictably large) fluctuations. This is means that queue size estimates need to be re-evaluated to avoid the queue length from growing uncontrollably.

Garbage Collection and Overflow

Not all systems have procedures in place to reclaim resources that are no longer in use. The reclamation of reusable resources, and the discarding of consumable byproducts is called garbage collection. In a queue, there is automatic reclamation as items are removed from the queue, but in memory or space allocation that is not necessarily true. Resource reclamation is crucial for the survival of systems in the long term\(^1\).

\(^1\)Humans eventually die because DNA does not perform (inverse) garbage collection of telomeres that are involved in DNA replication.
12.8  QUEUE RELATIONSHIPS OR BASIC “LAWS”

Basic dimensional analysis, or linear rate equations provide a number of basic “laws” about queues. These are rather simple relationships that barely deserve to be called something so elevated as laws, but they express basic truths about rate flow systems. They are nicely summarized in [Jai91]. We mention a few examples here by way of illustration. The laws express basic linear relationships about service flow rate. These form the basis of subsequent approximations in section 18.5, for instance. We use the following definitions for the $i$th server:

- **Arrival rate** $\lambda_i = \frac{\text{No. of arrivals}}{\text{Time}} = \frac{A_i}{T}$ (12.2)
- **Throughput** $\mu_i = \frac{\text{No. of completions}}{\text{Time}} = \frac{C_i}{T}$ (12.3)
- **Utilization** $U_i = \frac{\text{Busy time}}{\text{Total time}} = \frac{B_i}{T}$ (12.4)
- **Mean service time** $S_i = \frac{\text{Busy Time}}{\text{No. of completions}} = \frac{B}{C_i}$ (12.5)

### The Utilization Law

The utilization tells us the mean levels at which resources are being scheduled in the system. The law notes simply that:

$$U_i = \frac{B_i}{T} = \frac{C_i}{T} \times \frac{B_i}{C_i}$$ (12.6)

or

$$U_i = \mu_i S_i.$$ (12.7)

So utilization is proportional to the rate at which jobs are completed and the mean time to complete a job. It can be interpreted as the probability that there is at least one job in the system (see eqn. 12.1).

**Example 131.** Suppose a web server receives hits at a mean rate of 1.25 hits per second, and the server takes an average of 2 milliseconds to reply. The law tells us that the utilization of the server is

$$U = 1.25 \times 0.002 = 0.0025 = 0.25\%.$$ (12.8)

This indicates to us that the system could probably work at four hundred times this rate before saturation occurs, since $400 \times 0.25 = 100\%$.

Although the conclusion in this example is quite straightforward and only approximate, it is perhaps not immediately obvious from the initial numbers. Its value therefore lies in making a probable conclusion more obvious.
Example 132. A University teacher complains that most of the terminals or workstations in the terminal room of the University are idle, when he looks in the afternoon, so there must be too many terminals and money can be saved by at least halving the number. The students, on the other hand, complain that they can never find a free terminal when they need one. Who is right? The system administrator decides to look through the logs and apply the Utilization law. She finds that the time an average student spends at a terminal is $S = 1$ hour, and the average number of users using a given terminal per working day (of 8 hours) is $\mu = 4/8 = 0.5$ per hour. The utilization is thus $U = 0.5 \times 1 = 0.5$. In other words, the terminals are in use about half of the time.

The system administrator realizes that the peak load on the system is time-dependent. Around midday, many students arrive looking for a terminal and cannot find one. In the early mornings and late afternoons the students are all sleeping (or in lectures, or both), so there are many spare terminals. By reorganizing their time, the system administrator concludes that the students could make use of the machines that are there but that an overcapacity of double is acceptable for covering peak load.

**Series and Parallel Utilization**

When dealing with arrangements of queues, working in parallel (independently) or in series (waiting for each other), it is sometimes helpful to replace the array of queues with a single effective queue. We can rewrite the formulae to obtain effective formulae for these cases by assuming conservation of jobs: what goes in must come out. This leads us basically to the analogy with Ohm’s law in electrical circuits, or Kirchoff’s laws.

If a number of queue servers is arranged in series (see fig. 12.2), then the utilization law applies to each component. System components in series have the same average flow rate $\mu_i$ through each component, assuming that the system is running in a steady state – otherwise the pressure in the system would build up somewhere in one component, which would eventually cause a failure (like a burst pipe). Thus the utilization of each component is

\[ U_i = \mu S_i. \]  

(12.9)
The utilization for the whole series of queues is thus

\[ U_{\text{tot}} = \sum_i U_i = \mu \sum_i S_i = \mu S_{\text{serial}}. \tag{12.10} \]

Thus we see that the average service time for the whole series is

\[ S_{\text{serial}} = \sum_i S_i. \tag{12.11} \]

It is simply the sum of the service times of each sub-queue. If, on the other hand, we couple the queues in parallel (see fig. 12.3), then it is now the total utilization that is shared equally between the queues, i.e. \( U \) is common to each queue. Thus

\[ \frac{1}{S_{\text{parallel}}} = \sum_{i=1}^{N} \frac{1}{S_i}. \tag{12.13} \]

Figure 12.3: System components in parallel share the flow between the different server queues. Here it is the average utilization that is common to each server, since each experiences the same average incoming load.

\[ \mu = \sum_{i=1}^{N} \mu_i = \sum_{i=1}^{N} \frac{U}{S_i} = \frac{U}{S_{\text{parallel}}}, \tag{12.12} \]

and we can write

If we have \( N \) identical servers \( S_{\text{par}} \rightarrow \langle S_{\text{par}} \rangle / N \), i.e. the service time is reduced by a factor of \( N \) on average. Alternatively, throughput can be increased by a factor of \( N \). Note, however, that this does not tell us how long a client task will have to wait for completion. To determine that, we need a more advanced analysis (see section 12.9).

**Example 133.** Suppose a student terminal room has 100 computers to share between 500 students. If the number of students increases now by 60, how many extra computers do we require to maintain
the same level of utilization? The simplest answer to this is a naive ratio estimation, using the ratio of students to machines.

\[
\frac{\text{Students}}{\text{Terminals}} = \frac{500}{100} = \frac{560}{x}
\]

Thus \( x = 112 \) terminals. However, this result assumes that students actually arrive as a predictable flow process. In fact, they arrive as a random process, and things are not as bad as this. Utilization of overlapping random processes can be more efficient than for predictable processes the events of one process can slot into the gaps in another. See section 12.9.

**Little’s Law**

Another self-evident consequence of rates is Little’s law of queue size. It says that the mean number of jobs in a queue \( Q^n_i \), in queue \( i \), is equal to the product of the mean arrival rate \( \lambda_i \) (jobs per second) and the mean delay (seconds) \( R_i \) incurred by the queue:

\[
Q^n_i = \lambda_i R_i.
\]

(12.15)

If one assumes that the queue is balanced, as in eqn. (12.1), then \( \lambda_i \propto \mu_i \), and we may write

\[
Q^n_i \propto \mu_i R_i
\]

(12.16)

or

\[
Q^n_i = c \mu_i R_i.
\]

(12.17)

Another way of speaking this equation is to say that the amount of information in the queue is proportional to both the throughput rate and the amount of time each job takes to execute.

Before leaving this section, note that this equation has the generic form \( V = IR \), similar to Ohm’s law in electricity. This an analogy is a direct consequence of a simple balanced flow model; it is not so much as analogy as an isomorphism. We shall make use of this result again in chapter 18.

**Example 134.** In the M/M/1 queue of section 12.7, it is useful to characterize the expected response time of the service centre. In other words, what is the likely time a client will have to wait in order to have a task completed? From Little’s law, we know that the average number of tasks in a queue is the product of the average response time and the average arrival rate, so

\[
R = \frac{Q_n}{\lambda} = \frac{\langle n \rangle}{\lambda} = \frac{1}{\mu(1 - \rho)} = \frac{1}{(\mu - \lambda)}.
\]

(12.18)

Notice that this is finite as long as \( \lambda \ll \mu \), but as \( \lambda \to \mu \), the response time becomes unbounded.
RESPONSE TIME LAW

A situation that parallels the coupling of Ohmic resistances in series is to consider the coupling of a number of queues in series. This is common in systems of all kinds: once one part of a system is finished with a task, it is passed on to another part for further processing. What then is the response time of the whole system?

Let $Q_{\text{tot}}$ be the total number of jobs in the system. Then clearly,

$$Q_{\text{tot}} = Q_1 + Q_2 + \ldots + Q_n.$$  \hspace{1cm} (12.19)

Also, the total time can be written by Little’s law as a product of the average throughput and the total response time, as

$$Q_{\text{tot}} = \langle \mu \rangle R_{\text{tot}}.$$ \hspace{1cm} (12.20)

The law applies to each of the component queues also, so eqn. 12.19 becomes

$$\langle \mu \rangle R_{\text{tot}} = \sum_{i=1}^{n} \mu_i R_i,$$ \hspace{1cm} (12.21)

where $\langle \mu \rangle$ is the average throughput of the system. Thus, we have the response time

$$R_{\text{tot}} = \sum_{i=1}^{n} \frac{\mu_i}{\langle \mu \rangle} R_i = \sum_{i=1}^{n} x_i R_i,$$ \hspace{1cm} (12.22)

where $x_i$ is the capacity of the component, or the fraction of jobs that are held in flow on average in component $i$:

$$x_i = \frac{\mu_i}{\langle \mu \rangle} = \frac{\text{Jobs/s in component}}{\text{Average total jobs/s}} = \frac{C_i}{\langle C \rangle}.$$ \hspace{1cm} (12.23)

This tells us that the total response time is simply a weighted sum of the individual response times and any moment. It allows us to see where bottlenecks are likely to occur, so that processing resources can be reallocated.

12.9 Expediting Tasks with Multiple Servers $\text{M/M/}k$

How does the ability to process input change if we add more servers to accept jobs from the input queue? Intuition tells us that this must be more efficient, but as always the answer should be qualified and quantified to be certain about what we mean.

Suppose we have $k$ servers in parallel, removing jobs from a single input queue. We assume that the servers are identical components (computers, humans etc), so that the input rate is still $\lambda$
and each server has a processing rate of $\mu$. Clearly the maximum processing rate is now $k\mu$ when all of the servers are busy. If we assume that there is no overhead incurred in allocating tasks to servers, then this is also the rate at which work is expedited. If there are fewer than $n < k$ jobs waiting, then the average service rate will be $n\mu$. Thus our balance equations are now:

$$\lambda p_{n-1} = \begin{cases} n\mu p_n & (0 < n \leq k) \\ k\mu p_n & (n > k) \end{cases}$$  \hspace{1cm} (12.24)$$

These can be solved to give:

$$\lambda p_n = \begin{cases} p_0 \left( \frac{\lambda}{\mu} \right)^n \frac{1}{n!} & (0 < n \leq k) \\ p_k \left( \frac{\lambda}{\mu} \right)^{n-k} \frac{1}{k^{n-k}} & (n > k) \end{cases}$$  \hspace{1cm} (12.25)$$

and $p_0$ is found by normalizing:

$$\sum_{n=0}^{\infty} p_n = p_0 \left( \sum_{n=0}^{k-1} \left( \frac{\lambda}{\mu} \right)^n \frac{1}{n!} + \sum_{n=k}^{\infty} \left( \frac{\lambda}{\mu} \right)^n \frac{1}{k!(n-k)!} \right) = 1,$$  \hspace{1cm} (12.26)$$
i.e. if we now let the traffic intensity per server be $\rho = \lambda/\mu k$,

$$p_0 = \left( 1 + \sum_{n=1}^{k-1} (k\rho)^n \frac{1}{n!} + \frac{(k\rho)^k}{k!(1-\rho)} \right)^{-1}.$$  \hspace{1cm} (12.27)$$

The probability that a task will have to wait to be performed $\kappa$ is the probability that there are $k$ or more tasks already in the system,

$$\kappa \equiv P(n \geq k) = \sum_{n=k}^{\infty} p_n = \frac{(k\rho)^k}{k!(1-\rho)} p_0.$$  \hspace{1cm} (12.28)$$

Similarly, the average number of jobs in the system is the expectation value of $n$:

$$\langle n \rangle = \sum_{n=0}^{\infty} n p_n = k\rho + \frac{\kappa\rho}{1-\rho}.$$  \hspace{1cm} (12.29)$$

Little’s law again gives the average response time for the system in responding to a task as

$$R = \frac{\langle n \rangle}{\lambda} = \frac{1}{\mu} \left( 1 + \frac{\kappa}{k(1-\rho)} \right).$$  \hspace{1cm} (12.30)$$

Readers can verify that $\kappa(k=1) = \rho$ and that the expressions above agree with the single server queue for $k = 1$. The $k$ dependence of the above expression is rather complicated; in terms of power-counting $\kappa$ is approximately neutral to changes in $k$, but decreases slightly for increasing $k$. Expression
(12.30) therefore tells us that as the number of servers \( k \) increases, the response time for incoming jobs falls off slightly faster than \( 1/k \). Thus, the response time of a single queue with \( k \) servers is slightly better than \( k \) separate queues, each with a \( 1/k \)th of the tasks to complete. This assumes that all of the jobs and servers are identical, of course. Why should this be? The reason is that single queue servers make all jobs wait even when load is low, whereas a parallel server strategy can keep the level of incoming jobs below the queueing threshold a greater percentage of the time. The difference becomes most noticeable as the traffic intensity increases.

**Example 135.** A web hotel company has five customers who need their sites hosted. The company must decide whether to use a separate computer for each web server, or whether they should host all sites as virtual domains on a single site with a load balancer or multiple CPUs. Considering the performance aspect, they note that traffic arrives most of the time at a rate of 10 hits per second (\( \lambda = 0.01 \text{ per millisecond (ms)} \)) and that the web server has a service rate of 20ms (\( \mu = 0.05 \text{ per ms} \)). Let us model this as an \( M/M/5 \) queue.

The traffic intensity is \( \rho = \lambda/k\mu = 0.04 \), and thus the probability that the server is idle is:

\[
p_0 = \left[ 1 + \frac{(5 \times 0.04)^5}{5!(1 - 0.04)} + 5 \times 0.04 + \ldots \right]^{-1} = 0.82.
\]

(12.31)

Clearly normal traffic levels are low, and this is a high probability. The probability that all the terminals are busy is

\[
\kappa = \frac{(k\rho)^k}{k!(1 - \rho)} p_0 = 1.1 \times 10^{-5}.
\]

(12.32)

Hence the average response time is:

\[
R = \frac{1}{\mu} \left( 1 + \frac{\kappa}{k(1 - \rho)} \right) = 20 \text{ ms}.
\]

(12.33)

Now, suppose we have used 5 separate machines each with an \( M/M/1 \) queue. The effective arrival rate \( \lambda \) can be divided evenly between them, so that \( \lambda' = 0.01/5 \). The probability of a given queue being idle is

\[
p_0' = (1 - \rho') = 0.96,
\]

(12.34)

and the average response time is

\[
R' = \frac{1}{\mu - \lambda} = 20.8 \text{ ms}.
\]

(12.35)

This shows that the multiple queues lead to a slightly larger result, even at this low level of utilization.
Suppose now, we compare the behaviours of these two alternatives at peak times, where the number of arrivals is ten times as much. Substituting $\lambda = 0.1$ and $\lambda' = 0.1/5$, we find that the probability that the response time for the $M/M/5$ queue is largely unchanged

$$R = 20 \text{ ms.} \quad (12.36)$$

However, the response times of the multiple $M/M/1$ queues are

$$R' = 33.3 \text{ ms.} \quad (12.37)$$

We verify that a multiple server handling of a single queue is at least as good as multiple queues, because the probability of a multi-server handler being idle is much smaller. Multiple queues with separate servers force even the shortest jobs to wait unnecessarily, whereas the a multiple server scheduling would have cleared these jobs quickly, leaving a greater chance of being able to handle incoming tasks immediately. This phenomenon is related to the ‘folk theorems’ in section 18.3, eqn. 18.24.

12.10 LOAD BALANCING FROM QUEUEING THEORY

As more services move into high volume datacentres, we increasingly see architectures based around the $M/M/n$ model for handling bulk user traffic. Meeting the requirements set by Service Level Agreements (SLA), with variable demand and resources is essential to businesses[Sea, XZSW06, LSZ06]. A common strategy for keeping service promises is to opt for “over-capacity”, i.e. providing more resources than are strictly needed on average, by some acceptable margin. This is a straightforward way of dealing with uncertainty, but it can be a relatively expensive way and, without a model for capacity, it is not clear exactly what margin is required for meeting the peaks in demand. The cloud computing model grew precisely out of the desire to escape from this overhead by sharing high entropy workloads[Bur13a]. A simple approach, which remains popular, is to adapt the service scheduling strategy and resource pool to better adapt to changing requirements[XZSW06, LSZ06]. In each case, there is a role for parallelism in the provision of the services both for the purpose of load sharing and redundancy.

One may examine how the simple queueing models behave as estimators of local load balancing techniques, by seeing how response times vary as a function of traffic intensity and number of servers[BU06b, BU06a, BB06] (see figure 12.4). It is known that the limitations of bottlenecking traffic for splaying to multiple queue processors is a strategy that network device manufacturers have not been motivated to solve efficiently, as their business is about selling boxes to perform just this function.

**Example 136.** If one believes the manufacturers of network equipment, load balancing is a simple problem. One approaches the problem from the viewpoint that the network has no capacity
Figure 12.4: A typical load balancer configuration, with a single input stream splaying to a number of servers. The policy of the load balancing node tries to approximate the $M/M/n$ queue, with varying policies to scheduling limitations, and that servers all have somewhat similar tasks, capacities and behaviours. In fact, jobs have both semantics and dynamics. There are three main problems to load balancing:

1. Accumulating all requests in a single broadcast domain.
2. Forwarding requests to an array of server queues.
3. Maximizing the entropy (balance) of requests to maximize throughput.

What criterion should we use to try to balance the load? Simple sequential round robin? Least recently used server? Queue length? CPU utilization? Load average? Free memory of the server? Whatever choice we make, we base the choice on assumptions. Because networking equipment cannot have knowledge of the intended semantics of jobs and services, it can only offer purely dynamical measures on which to make a selection. However, this cannot allow for differences in job size, or knowledge of time to complete, or the resources needed to complete the request. Jobs handled by a single server can be in contention with one another, leading to highly unpredictable service times. The problem here is that load balancers have no cooperative relationship with the servers, and therefore ‘impose’ work leading to conflicts. An alternative is for servers to be in the driving seat of reasoning about their load capacity. They may ‘pull’ tasks [BB08] to maximize both dynamical and semantic information.

The results of experiments to test a commercial load balancer against the theoretical values, even in the mid 2000s, was quite interesting. The authors in [BU06b] found that for low loads, any sharing policy was equally fine; however, at high loads there was little to gain from trying to optimize the sharing according to different policies (round robin, least recently used, etc). While some performed better for a while, the additional cost quickly came to nothing the critical point. The best advice was simply to stay away from the critical region and not waste money on expensive hardware. Buying more of cheaper commodity switches was a better strategy, as they
Figure 12.5: The performance of a queue approaching the critical unstable traffic intensity. At around 100 requests per second, we see the response time for a server grow suddenly and rapidly as the queue becomes overloaded. This shows the two regimes of the queue: ‘all is well’ versus ‘thrashing’, with a sudden instability in between. At this point, all predictability falls apart. Thus, staying below this threshold is critical.

All failed in a similar way (see figure 12.6). It is worth noting how this study actually revealed the performance of the load balancer itself as a bottleneck when adding more servers. The problem of queue marshalling is not only dependent on traffic intensity but also number of destination queues. The study in [BU06b] called into question what still remain the most popular methods of load balancing in datacentres today.

The experiments were based on pushing into queues, with an unpredictable Markov process, but why would we expect this to give the best results for load sharing? Anycast methods have also grown in popularity for small networks, but they suffer from similar contention issues[EPB+98]. Allowing each server to throttle its own traffic using a ‘pull’ approach could also be considered. This would avoid the queueing instability all together on the server side, and push the uncertainty onto clients, where only single requests need to be handled. Pull based load balancing was studied in [BB08] and showed several benefits over the push approach, using voluntary cooperation as a paradigm for sharing. However, few systems are designed to scale in this way. One exception is
Figure 12.6: Showing how the M/M/5 performs compared to the theoretical $M/M/1 \times 5$ pre-shuffled. We see, as expected, that the $M/M/5$ outperforms 5 $M/M/1$ queues. The response time of a real switch tracks the $M/M/5$ well far from critical intensity; however, close to instability, it fares below the $M/M5$ theoretical value too, even for only 5 servers.

The so-called Content Delivery Networks (CDN) [CHC+11, VP03]. Regrettably, shared computing infrastructure or ‘clouds’ do not make currently such methods available to clients in a simple way. The theory suggests that this is a mistake. We’ll return to this issue in volume 2, where the notion of autonomous cooperation is studied at length.

### 12.11 Maximum Entropy Input Events in Periodic Systems

One of the themes that we return to repeatedly in this book is the idea of systems that respond to random events. When these events change measurables in an unpredictable way, we call them fluctuations. Characterizing fluctuations is important because it is a way of representing the spectrum of input.

The problem with random events is that they come from an environment that is complex and
that we know little about. By definition, the environment is that which is outside our control, so how should be describe the unknown? One way of approaching this problem is to assume the greatest level of randomness, given a constraint that says something about the interaction of the system with the environment.

If we characterize input as an alphabet of symbols labelled by \( i \) and with probability distribution \( p_i \), then the assumption of maximum randomness can be accomplished by using a maximum entropy principle (see section 9.11). This is a form of 'constrained fair weighting' that is sometimes used as a way of scheduling events between different service centres (servers).

**Example 137.** Consider the problem of modelling the input event stream of a server, with expected input current \( I \) symbols per second, formed from the weighted sum of individual symbols rates \( I_i \):

\[
\langle I \rangle = \sum_{i=1}^{N} p_i I_i. \tag{12.38}
\]

Thus, we maximize entropy given this 'constraint' on \( p_i \). The result is the well known Boltzmann distribution \( p_i \propto \exp(-\beta I_i) \).

The most common solutions of the maximum entropy hypothesis are the completely flat distribution (all input events equally likely) and the exponential Boltzmann distribution that follows from a constant sum constraint.

How good is this assumption of maximal entropy fluctuation? This was investigated partially in ref. [BHRS01], where it was found that the assumption is a good representation, but only if a modification is made to the maximum entropy signal. The modification is a periodic scale transformation that follows the daily and weekly rhythm of system behaviour.

If we fit a maximum entropy distribution to actual data, it will have constant variance, or the moments of the distribution will remain fixed. However, this would imply that the environment was a steady state process, which we know to be false. Observations show that there is a distinct periodicity, not only in the expectation values of environmental signals, but also in their variance over a periodic representation of several weeks (see fig. 2.1).

Some authors maintain that this is evidence for self-similar behaviour, and the modified distribution is in fact a power law spectrum rather than a maximum entropy distribution. One possible explanation for this is the similarity in form between the asymptotic form of the stable Lévy distributions and the form of the periodically corrected maximum entropy model ([Sat99]).

### 12.12 Miscellaneous Issues in Scheduling

Numerous additional factors about human-computer systems can be incorporated into time management: humans get bored, they have different specializations and preferences, different skill
levels etc. Finding the optimal use of time, given these additional constraints requires a more sophisticated model, with heuristic currencies.

Once model that describes human scheduling of interruptions or intervals of concentration is the game theoretical payoff model described in section 19.9. In this model, which concerns cooperation versus competition, we can think of time-cooperation as giving humans space to work in fixed schedules and competition for time as being continual interruptions at random. In other words, tasks cooperate in orderly scheduled quotas, or they demand resources at random. This model addresses many situations, and we can use it for human scheduling too. The basic result from game theory, although simplistic, tells us that random event competition works well as long as the number of jobs is small, i.e. as long as there is plenty of time and no one is taxed to the limit. However, as someone becomes very busy, it is better to schedule fixed quota time-slices for jobs, otherwise all jobs tend to suffer and nothing gets done. The overhead of swapping tasks at random can lead to great inefficiency if time is short.

The potential for harvesting the vast computing resources of the Internet has led to the notion of Grid computing, or massive distributed parallelism. To accomplish world-wide cooperation of computing resources, models extend their time-scales to include batch-style processing as part of a larger task. Examples of this include the screen-saver processing methods used by the Search For Extraterrestrial Intelligence (SETI) project. Each of these cases can be studied with the help of parallel and distributed scheduling models.

**Applications and Further Study 12.**

- Designing workflow patterns and schedules.
- Resource deployment.
- Measuring workflow and efficiency.
- Modelling of generalized queuing processes.
CHAPTER 13

SYSTEM ARCHITECTURES

In the foregoing chapters we considered mainly systems that could be described by simple scalar variables, with no larger structure. That was a useful simplification for considering the basic effects of change and dynamics, but few human-computer systems, worthy of investigation, are this simple. In technology, we normally want to design a system with a specific function in mind. System design requires a number of elements:

- Strategy.
- Policy.
- Procedure.
- Activity.

These fit together into flows of information that govern change within the system.

13.1 POLICY AND PROMISES OF ORGANIZATION

No system has to have to have an intended purpose in order to function, but observers might interpret a system as fulfilling a function independently. Biological organisms do not have a purpose *a priori*, they exist only to reproduce and start over; if they are lucky they find an ecological niche. Trees and plants may spring up and prevent the erosion of sand in a beach or desert. They were not planted on purpose (though this would be a possibility) but they nevertheless fulfil the function of preventing erosion. When a functional quality is emergent rather than intended, third parties may still make use of the outcomes, and even interpret them with intent\(^1\).

\(^1\)This is the crux of the confusion between Darwinian evolution and so-called intelligent design.
The concept of intent is subtle. Following the first 2003 edition of this volume, it became clear to me that the whole issue of intent was missing from the story of systems, and needed a description of its own. In computer programming, one could say that programs and algorithms represent intentional policies for behaviour, but only at a very details causative level. Intent spans more layers than flowchart algorithms. The outcome of that observation was the birth of Promise Theory [Bur05, BB14], which proposes to reformulate all aspects of interactions between system parts. In the spirit of overview, in this volume, we shall not describe it here. Rather, the application of Promise Theory to systems is covered in detail in volume 2.

A collection of promises made about the behaviour of a system is called a policy. Policy is thus used in describing the management of systems to mean a specification of a goal, together with decisions limiting the behaviour of processes within. Policy has potentially several kinds of goal:

- To ensure an outcome.
- To prescribe a constraint.
- To maximize production of a benefit.
- To minimize the extent of a problem.
- To identify and distinguish input (accountability).

These are ad hoc decisions about a system: they cannot be derived. Policy exists at the high level and the low level. High level policy might include ethical considerations and other human issues. Low level policy constrains the details of work processes, e.g. in the placement of regulating valves (administrative overseer) for control, convenience measures, security measures, consistency, and the overall flow of information.

### 13.2 INFORMATIVE AND PROCEDURAL FLOWS

Dynamical systems generate and manipulate information. They require input and they generate output. The information flows around the system as the system carries out its function, and the final result is usually a synthesis of products which has been accumulated and assembled from different parts of the cooperating system.

It is important to distinguish two types of flow, or development within systems:

---

**Definition 52** (Algorithmic flow). This is a causative map of the way the actions taken by the system interact, and how control flows from promise to promise, or from outcome to outcome, within the processes of the system. The instructions or promises within a system are often laid out from start to finish, in a causal list, or embedded in a control loop. This represents a flow of activity and authority in the system.
An example of this is a computer program.

**Example 138.** A computer program has a functional structure, with each function composed of a linear sequence of instructions. Some of these instructions require the evaluation of subroutines or subordinate functions. A university is a system which accepts students and churns them through a learning process, emitting them at the other end hopefully invigorated with new ideas and abilities. There is a definite flow within this system, from lecture to lecture, with Event driven systems, on the other hand, respond to specific happenings. A fire service is an example of an event driven system.

Another kind of flow is that taken by the resources that are manipulated by the processes.

**Definition 53 (Resource flow).** The produce of a system, or the work it actually carries out, also has a rate of change. This is the rate of resource and information flow.

The flow of information does not necessarily mirror the flow of algorithms and instructions which produces it.

**Example 139.** The command structure of an army might involve a flow of decision making by radio to and from a command centre, during a military operation. The movement of the troops (i.e. the resources) is not related to this information flow.

**Example 140.** In a user-support desk or help desk, a flow of control information passes between the user and those helping the user. The actors which help the user manipulate resources in order to solve the user’s problem. These two information flows are different.

The distinction between algorithmic and resource flow is important to the design of a system, because one should not be tempted to organize these two flows by the same standards. Both flows are important to the overall function.

### 13.3 Structured Systems and *ad hoc* Systems

In most cases, data and processes are arranged in components, objects or *entities* that allow specialization and collaboration. These cooperate in the execution of the system, by communicating via channels of communications. These might be ‘word of mouth’, written on paper or electronic; the means of communication is unimportant.

Consider the names in table 13.1. These names represent logically distinct elements within different kinds of system. The units are often arranged in a hierarchical fashion, e.g. a site contains several departments, which in turn contains several groups, composed of individuals; a database contains many records which, in turn, consist of many sub-records.

The structure of systems that perform a stable function is usually fixed, for the duration of that function. However, this limits their ability to adapt to slight changes in their environments. Change
can therefore be allowed to take place in a deterministic way by having a continual reevaluation of the system at regular intervals. An alternative procedure is to allow systems to perform randomly.

An *ad hoc* system has no predictable structure over long intervals of time. We can define it as follows:

| Definition 54 (Ad hoc system). A system is said to be ad hoc if its structure is periodically redetermined by random variables. |

**Example 141.** Mobile ad hoc networks are networks for radio communication that are formed by randomly moving mobile devices with transceivers. These devices link up in an opportunistic manner to form a relaying network. Messages can be routed and relayed through such networks if a sufficient number of active devices is in range of one another.

It is tempting to associate ad hoc systems with the concept of a random graph; however, an ad hoc system need not be a random graph — it could be based on a predetermined structural plan, but with only a finite probability of being connected at any given moment.

### 13.4 Dependence on External Agents

To discuss policy and intent, we need to describe agency, or simply *agents*. All components may be considered agents in their own right, which interact and behave according to the promises they make. This is the essence of Promise Theory (see volume 2). All systems of components depend on other components to keep their collective promises: that is implicit in the definition of what we mean by a system. This dependence might be a strong dependence, with dramatic consequences if a part fails, or a merely weak influence.

Some factors in a system enable the system to function, while others merely enhance or amplify its abilities. In the next chapter, we shall consider how to rationalize and analyze the effects of dependencies; for now, we introduce only the concepts.

| Definition 55 (Strong dependence). This is a dependence on another part of the system, in which the removal of the dependency (the tie between the dependent and the dependee) leaves the dependent (that which depends) unable to function. |
Strong dependence implies strong coupling, and non-linear behaviour, and so-called complex behaviours.

**Definition 56 (Weak dependence).** This is a dependence in which the removal of a dependency does not prevent the functioning of the dependent, but merely alters its possible behaviour.

Weak dependence implies weak coupling, and approximately linear behaviour, with decoupling of system scales.

**Example 142.** An aircraft depends strongly on its fuel; without fuel it cannot function. The same aircraft depends weakly on the weather; this can affect its performance, but does not prevent it from functioning.

A computer system depends strongly on its hardware and operating system; without this it cannot function. The same computer depends only weakly on its third party software.

**Example 143.** It is known from the study of cities (Metropolitan areas) that the scaling of economic output and certain measures, like the lengths of roads, etc, do not depend strongly on who or what happens in a city. Cities scale in a ‘universal’ manner, which is typical of weak coupling\[Bur16a\].

**Example 144.** River systems satisfy a universal empirical rule concerning the distance between riffles and pools (the riffle-pool sequence), which are found to be separated by 5-7 times the width of the river. In this form of scale-free universality, a dimensionless ratio of distance/width $\sim 5$ is characteristic of all rivers, in spite of the ground type or rock type, etc.

A system must be analyzed in terms of its dependencies to understand its behaviour at that scale. At scales much larger than this, weakly coupled systems may not reflect the low level dependencies.

Dependence on other parts of a system has many implications for a system, including its efficiency and its ultimate limitations. As implied above, a critical dependence on some component can affect the ability of the system to perform its function. If such a crucial part becomes damaged or unavailable, then the whole system can fail to function: the system is then said to have a single point of failure. Note this phrase does not necessarily mean that there is only one of them, but that one is enough to cause complete failure. In other words, the failure of the system at a single point is enough to halt it completely. This is clearly a precarious situation to be in, and is usually a sign of poor design. A strategy for avoiding such problems in system design is redundancy, or the use of backup systems.

**Definition 57 (Redundancy).** The duplication of resources in such a way that there are no common dependencies between duplicates, i.e. so that the failure of one does not lead to the failure of the others.
Here is an example of lack of redundancy.

**Example 145.** A clustered structure of computers centred around a single server forms a hub-centric structure. If the computer at the centre of the hub should fail, all of the dependent computers will cease to function properly. There is no backup.

**Example 146.** Developed societies are almost completely dependent on electricity for their functioning. If a bad storm, or flood were to take out power lines to a major community, it would cease to function. This is the reason why military operations usually target an enemy’s critical infra-structure first.

Redundancy is an important strategy, but too much redundancy can lead to inefficiency.

**Example 147.** Systems depend on the flow of information through channels of communication. Unstructured communication often results in too much communication (repetition and little progress). Regulation of dependencies is a strategy for minimizing unnecessary uncertainty.

An alternative to redundancy is rapid repair or maintenance.

**Example 148.** If we can repair a system faster than users sample its services, then that system looks a lot like a Shannon error correction channel. Rather than failing over to a redundant component, a failed component can be repaired quickly to make the system available again, before anyone notices. If repair is cheap and fast, this may be preferable to having multiple redundant resources that sit idle (underutilized).

### 13.5 System Design Strategy

There are many ways to attempt the design of a system. One looks for a strategy for breaking a complex task down into manageable pieces in order to execute it. Experienced practitioners rely on their experience to suggest solutions, but sometimes intuition can fail and a more rational enquiry is needed to make a decision. We shall deal with the requirements for rational decision-making in the remainder of the book. Some common alternatives are described here.

Note that the graphical view of systems expounded in this book implies no particular need for hierarchy. There is a long standing tendency of imposing hierarchies on systems, even where they might be detrimental to functioning, because hierarchies are ingrained into most societal structures. One of the hardest lessons to learn in system analysis is that decision making does not necessarily imply a directed branching tree-like structure: it can also arise by competition, by voting, by ad hoc timing or by random events. Our main concern is whether systems are appropriately stable in their decision making.
MODULAR DESIGN AND SCALE ASSUMPTIONS

Different authors use this expression in different ways. Modular design is, first and foremost, a strategy of breaking a system up into smaller parts, in order to promote comprehensibility, adaptability and extensibility. Is a single, closed function a module? The answer is clearly yes, but one normally reserves this expression for larger entities than this.

Modularity is a form of reductionism[Bur15]. It is important to remember that the correct functioning of all modules in a system does not guarantee the correct functioning of the whole. At each scale of modularity, where components are connected together and interact, there is new information that is not present in the components themselves. This is crucial to the functionality, or promises made by the system as a whole.

Example 149. In the testing of systems (e.g. software unit testing, the checklist validation of factory processes, etc), a fully compliant checklist of all promises does not guarantee a properly working system. A television can be made up of perfectly working components, but fail to work because a connection is missing or because it is not configured to receive the appropriate channels.

What is modular at one scale may seem integrated at a larger scale. In any system, modularity is a perception rather than a reality. It makes most sense to speak of modules when they are weakly coupled.

Example 150. At the scale of a person, a car appear as a collection of components (wheels, seats, engine, etc). At the scale of a region or country, cars appear as a formless gas of ‘traffic’ whose behaviour is not at all dependent on any modules within any car.

TOP DOWN VERSUS BOTTOM UP

Two design strategies for building systems have emerged, and have developed with increasing refinements and compromises between the two viewpoints. Although it is too much to ask for a consensus of naming, broadly speaking, these are represented by the following:

- Top down: The name ‘top down’ is motivated by the traditional way of drawing hierarchical structure, with high level (low detail) at the top, and increasing low-level detail at the bottom. A top down analysis is goal driven: one proceeds by describing the goals of the system, and by systematically breaking these up into components, by a process of functional decomposition or normalization.

- Bottom up: In a ‘bottom up’ design, one begins by building a ‘library’ of the components that are probably required in order to build the system. This approach is thus driven by specialization. One then tries to assemble the components into larger structures, like building blocks, and fit them to the solution of the problem. This approach is useful for building a solution from ‘off the shelf’, existing tools.
The difference between these two strategies is often one of pragmatism. A top down design is usually only possible if one is starting with a bank slate. A ‘bottom up’ design is a design based on existing constraints, namely the components or resources that are to hand. An advantage of building from the bottom up is that one solves each problem only once. In a top-down strategy one could conceivably encounter the same problem in different branches of the structure, and attempt to solve these instances independently. This could lead to inconsistent behaviour. The process of ‘normalization’ (see chapter 14) of a system is about eliminating such inconsistencies.

**Example 151.** (Computer system - ‘top down’) In a the design of a new computer system, one examines the problem to be solved for its users (banking system, accounts), then one finds software packages which solve these problems, then one chooses a platform on which to run the software (Windows, Macintosh, Unix), and finally one buys the and deploys the hardware that will run those systems.

(Computer system - ‘bottom up’) First one buys reliable hardware, often with operating system already installed, and installs it for all the users; then one looks for a software package that will run on that system and installs that. Finally, the users are taught to use the system.

**Example 152.** (Enterprise - ‘top down’) In the organization of a maintenance crew, one looks at the problems which exist and breaks these down into a number of independent tasks (plumbing, electrical, ventilation). Each of these tasks is broken down into independent tasks (diagnosis, repair) and finally individuals are assigned to these tasks.

(Enterprise - ‘bottom up’) The enterprise bosses look at everyone they have working for them and catalogues their skills. These are the basic components of the organization. They are then grouped into teams which can cooperate to solve problems like diagnosis and repair. Finally these teams are assigned tasks from the list of plumbing, electrical work and ventilation.

In system administration, especially configuration management, these two strategies are both widely used in different ways. One may either implement primitive tools (bottom up) that are designed to automatically satisfy the constraints of a system, or one can use trial and error to find a top down approach that satisfies the constraints.

**FUNCTIONAL DESIGN AND OBJECT DESIGN**

Similar to the ideas of top down and bottom up design, are the design strategies widely used for building computer software, based on functional and object decomposition. Computer software is simply a system of algorithms and data structures, which operate on the resources of a computer. The principles of software design are the same as those of any system design.

A functional design is an algorithmic (top down) design, in which the goals of the system motivate the architecture of the components within the system. The system is geared around the evaluation of its function, i.e. performing its task. This approach is illustrated in figs 13.1, 13.2 and
11.1. The system state is usually centralized in a monolithic data structure, whereas operators are distributed throughout the different functional entities within the system.

![Figure 13.1: A functional structure diagram for a computer system, somewhat simplified, shows how each level of the computer system depends on other parts.](image)

In a object oriented design (bottom up), one begins by isolating logically independent objects or tasks within the whole system, i.e. tasks which are able to carry out a specific function without outside help (see fig 13.3). One then constructs the system around these objects by sending messages between them. Any details which only apply to the particular object are contained within it, i.e. hidden from public view. Whereas a functional model reflects a flow of activity and information between the levels of the boxes, an object model requires a separate plan for action, at each level of its operation. Private algorithms are usually referred to as the methods of the object. An object model requires at least one public ‘method’, or control function, which binds the objects together. The state of an object is private to that object, and all of the operators which change that state are private to the object. In order to function in concert, one often requires the guidance of a simple functional system. The total state is decentralized, and is the sum of the private states of each of the objects.

It is sometimes said that ‘objects’ communicate by sending messages (like a peer to peer network). While this is the idealized viewpoint taken in pure object oriented design, it is often an exaggeration of the truth. In practice, there is some guiding functional super-structure which manages the objects; seldom are they able to work independently. We can imagine this as a principle.

**Principle 6 (Separation of management and work structures).** A higher level, weakly coupled management framework may be helpful to bind low level operations together, and guide them towards the larger goal of policy, both in human and machine parts of any system. The management framework is formally separable from the low level operations only if weakly coupled.
Figure 13.2: A functional structure diagram for the meta-system which manages (administers) a computer system. The diagram, somewhat simplified, shows how each task level of the system depends on sub-tasks. This diagram is easily drawn and easily used to over-simplify the issues in system management. For instance, it gives the impression that management is a one-off process and moves from left to right in the diagram whereas, in fact, it is a dynamical process in which all of the parts are called upon repeatedly.

Each object has a private notion of state, and its own internal methods for changing that state. The object can communicate with the outside world, through authorized channels, but is otherwise closed. From a security or privacy perspective, the object model is based on the Clark-Wilson security model, or role-based security.

Object strategy is conventionally mixed together with the idea of classification also. Classification has to do with object types or name conventions. For example, if one defines a object which is a bank, then there might be several instances of a bank system in use, at different locations. If all of those instances share the same design, then they are then all said to be members of a class of banking systems.

In a rough sense, a pure, functional decomposition is rather like the dreams of the communist states: in which every part of the system is steered from above by the orders propagating down the hierarchy, and in which every part contributes in its own unique way to the larger goal of executing the function of the system. An pure object model, on the other hand, is somewhat like a system of decentralized control, in which different objects work independently, but cooperate in order to achieve larger goals. These are two different strategies for building systems. The relative efficiencies of the two strategies are hotly disputed, but there are no unilateral conclusions. At best we can say that the ability to control requires the system to be well behaved, or sufficiently stable. Then, ‘control’ implies change by external influence.

**Principle 7 (Control).** *The perception of control may be illusory. Objects must either compete freely with one another, or be guided by a super-structure in order to be ‘controlled’, i.e. to comply with intent. This does not necessarily require centralization or hierarchy, only persuasive interaction.*
Figure 13.3: In a object design, one bases the structure around the objects which need to be visible at the functional level of the system. Any dependencies which belong only to the object are concealed within the object concerned.

The meaning of control will be discussed at length in volume 2.

The advantage of functional decomposition is that it has a structure which directly reflects the task it is designed to perform. It is therefore easy to understand in a causal sense. The advantage of the object model is a more rigorous separation of logically independent tasks and resources, which makes the reuse of those separate items in other contexts easy. The lack of an obvious structure of algorithmic flow makes understanding the causality of an object system harder, however. Current trends tend to favour the object model because, if nothing else, it provides a tidy way of house-keeping the parts of the system and their dependencies.

In practice, it is rare to find either of these strategies in a pure form. Both strategies need each other in order to succeed: an object model has no implied algorithmic flow, so it needs a functional model along side in order to guide the flow of production. Similarly, any function within a functional design is a rudimentary object, and benefits from a disciplined logical decomposition. Thus one must view these two philosophies are being complementary.

CLIENT MODEL AND PEER MODEL

Another way of looking at the centralized versus de-centralized debate, is through the paradigm of services. In a functional decomposition, one could say that each function requests services of its subordinate functions. Similarly, in an object decomposition, each object contains internal services (methods), but can also opt to send messages to other objects in order to request information, by delegation: this request is a service performed by one object for the other.

Example 153. A building contractor might hire a sub-contractor to perform part of a job, in which it lacks skills. The sub-contractor does not work 'under' the main contractor in actuality; it works
along side it. Thus while the conceptual flow was initiated by one contractor, there is nothing intrinsically subordinate about the sub-contractor.

Rather than focusing on issues that are subordinate to others, in the manner of a hierarchy, one can also view a system from the perspective of its actors. This is particularly appropriate to systems which are already distributed in space, either by geographical considerations, or by network.

- **Client-service model:** In this viewpoint, a system is broken up in an orderly fashion into a service providing part and a service-requiring part. This is the traditional model of shop and customer.

- **Peer model:** This viewpoint is a more anarchistic viewpoint of the actors, as skilled individuals. A peer model has no ‘policy centre’. Everyone can do something for everyone else. No one is intrinsically a client or a service provider; anyone can play any role. In this way, one hopes to maximize access to individual resources, without shielding those resources from view, by layers of organization. This model lacks a clear organization and is vulnerable to loss of focus. On the other hand, it is highly robust to total failure.

These models tend to apply in different contexts.

**Example 154.** Computers that are ‘policy independent’, i.e. which have independent policies can interact but often wish to protect themselves. There is no automatic trust. A client-server type of service request, opens up one party (the server) to a situation of vulnerability in which clients might actually attack. In [BB05b], the idea of voluntary cooperation is used, in which each individual device in a network collaborates by prior agreement, and then only from behind a ‘hands-off’ firewall system, like the one-way drawers that are used to pass money over the counter in banks and post offices.

Open source software development is an example of a peer model, for instance. A peer model also applies in the larger landscape of commerce, for instance, where many companies and individuals require each others services, and they are all equally important to the running of the ‘total system’. This is the way that society works, as a whole. The peer model is also very much an object viewpoint: one defines objects, based on their particular attributes, without attempting to evaluate which objects are subordinate to others (at least to a first approximation). All objects are equally important, but there is cooperation between them. The peer model can be very efficient, in local groups, but it presents new problems in scaling to large groups.

The client-service model is a more focused strategy for a single part of a larger process. It is more restricted, but more orderly. It is more susceptible to bottlenecks and efficiency problems, because one focuses the execution of all work at the service provider. Many clients to a single service provider can easily overwhelm an poorly designed service provider.
13.6 Capabilities and Limitations

Not all system designs can make or keep promises of arbitrary kinds. A completely centralized system could not promise resilience, nor could a totally distributed system promise total consistency, without disclaimers. In approaching system design, we have to be ready to confront limitations that are inherent within them. Some are basic laws of physics: we cannot exceed the speed of light in transmitting data, for example.

- Dependencies on technology (human, computer, etc).
- Dependencies on procedures and algorithms.
- Dependence on memory and storage.
- Dependence on availability.
- Dependence on consistency.

In the theory of distributed systems, in Computer Science, much effort is invested in proving theorems about consistency of data under different conditions. Some of these are at the stricter end of the spectrum (e.g. Paxos consensus discussion [Lam01]), while others are loose and handwaving (e.g. to so-called ‘CAP theorem’[Bre00], which is not a theorem at all in the mathematical sense).

Timescales are always a critical aspect of system behaviour. We can never expect a system to keep any promise without specifying the timescale over which we expect the promise to be kept. This follows from the fact that any realization of a promise requires a process, which necessarily involves changes and hence the passage of time.

Example 155. We may have innovative ideas quickly, but it takes a long time to develop them into a result or a product. We may be able to sample data quickly from a sensor, but it takes a long time to build up a calibrated picture of what is normal or abnormal. We may be able to access the domain name address of a website many times per second, but if it only changes once per month, such a process would be inefficient.

13.7 Event Driven Systems and Functional Systems

Systems can be characterized by whether they are designed only to respond to the requests of clients (i.e. they are services), or whether they have a pre-programmed procedure that evaluates some function (they are standalone programs).

- A system that exists only to service others, needs to be able to schedule work flow to cope with demand at the behest of others.
A system that produces, manufactures or evaluates something is freer to organize its time and resources according to its own scheduling plan.

In large organizations and computer systems, there is often a mixture of these two types of system and the resource requirements for the two processes can conflict.

Example 156. A help desk that is constantly being interrupted by telephone calls cannot produce information, courses or solve real problems that require more functional work.

Example 157. A computer that is acting as a database server, and as a numerical analysis engine must share resources between these tasks. The server requires intensive access to disk and interrupt driven resources, whereas the numerical calculation requires constant periods of CPU concentration to advance. These two requirements are not strongly compatible.

There are two ways to cope with this kind of conflict:

- Separate the two parts of the system into independent units.
- Interleave the scheduling of resources in a single unit.

The latter strategy is more difficult for humans to accept than for computers, as humans are slower to switch context than machines. The advantage of the second strategy for human organization, if it is performed as a long term job rotation, is a broader experience. Humans are, after all, learning systems and can benefit from broader insight into a system’s operation. However, the time scale of the interleaving must be appropriate for humans.

13.8 THE ORGANIZATION OF HUMAN RESOURCES

Organization includes many issues:

- Geographical organization,
- Psychological organization,
- Process decomposition: how the organization is split up into different activities (whether these overlap, leading to consistency and integrity issues).

The geographical organization of human-computer systems has become less important with the arrival of information networks and telephone systems, but there are still issues where geography can play a role, such as the need for face to face communication. To analyze resource organization, we need a value system or currency of organizational structures.

1. Efficiency (cost/speed).
2. Convenience.

3. Comprehensibility.

This is a task for observational verification over time.

Humans are known to be good at decision making and creative thought, but poor at discipline and consistency. It seems sensible to assign humans to creative work and machines to repetitive, precision work\(^2\). The role of human beings in systems, be they human-computer systems or other man-machine liaisons, has been studied in a variety of frameworks, using fuzzy logic, symbolic interpretation and other cognitive hooks. The extensive literature can be navigated starting from a few bases, see [McR80, Ras83, Rou89, End95, She96], for starters.

Socio-anthropological research suggests that these two human faculties require different organizations. Decisions are made quickly by individuals, but individuals do not always have impartial interests or complete information, thus one generally involves several individuals in decision making. Peer review of decisions is performed by committees. For effective committee work, where decisions need to be made quickly, a group size of no more than six is found to be a limit beyond which ordered decision becomes chaos. Conversely, for brainstorming and creative thought, a larger group size is an advantage ([Dun96]).

Research into humans’ abilities to collaborate is based on the brain size to group-size hypothesis. Human social group sizes are observed to be limited to 150 people in organized society; this is about the number of friends and acquaintances that we can relate to. For animals with smaller neo-cortices, the number is smaller. This suggests to anthropologists the hypothesis that organizations that grow beyond about a hundred individuals are likely to become unmanageable.

Another datum is of interest here. Hunter-gathering tribes of humans have evolved to work in groups of more than about 30 before they become unwieldy and break apart, even though they might regroup in a social context to numbers of up to a hundred and fifty, so it is thought that busy groups of workers are limited by this order of magnitude ([Dun96]). In other words, when we are preoccupied with work, we have less aptitude for dealing with other persons. Thus, if we take our evolutionary heritage seriously, it makes sense to pay attention to any hints provided by our genetic heritage, and perhaps limit the sizes of working groups and enterprises with these numbers in mind.

\(^2\)It has been suggested to the author, informally, that skill based management is a waste of time for companies. In terms of economics, one has more to lose today from the consequences of expensive license agreements and contract clauses than from learning particular technical skills. The losses due to incompetence are negligible compared to the cost of expensive license agreements for a company’s software base. Readers may calculate for themselves whether this might be true in their own enterprise.
13.9 **Principle of Minimal Strong Dependency**

If strong dependencies can lead to cascade failures, then it makes sense to try to minimize their use in system design. The minimization of dependencies is a principle that attempts to reduce the possibility for failure and inefficiency, including the collateral cost and damage associated with failure. In any system there may be a conflict of interest between tidiness of structure and efficiency of operation.

**Example 158.** *A traditional hierarchical structure, stemming from a military past leads to great orderliness, but is based on strong dependency of layer upon layer.*

Dependency carries with it a *functional inertia or resistance*, due to cooperative communication, that makes systems potentially inefficient. It also makes dependencies into points of failure, since it increases the number of parts that have to be connected for a functioning system.

**Definition 58 (Point of failure).** *In a graph* $(X, \Gamma)$, *any node or edge of the graph is a potential point of failure.*

The principle of minimal dependency sometimes conflicts with the need for system normalization (see chapter 14). The balance between order and efficiency may be formulated as a *game* of strategy, in the Game Theoretical sense (see chapter 19).

13.10 **Decision Making within a System**

Systems have to make decisions in many contexts. Decisions affect changes of process, strategy or even policy, based on information which is available to a part of the system. The question of where decisions should be made is a controversial one, both in human-centric and computer-centric systems. The decision requires both access to the information and a knowledge of the goals, i.e. the policy of the system.

13.10.1 **Layered Systems: Managers and Workers**

All decisions are ultimately made by humans at some level (if only the decision to delegate to an prescribed algorithm or formal process). Sometimes decisions are cached or assigned to proxies to implement. Traditionally, system decision making has been a task assigned to ‘managers’ or ‘commanding officers’. This stems from two reasons: first, autonomous machine decision has not always been technologically possible, so it has fallen to humans to guide every detail as a matter of
history; second, is the prejudice that managers or commanders have superior intelligence to lower level workers or components\[^3\] . Consider these strategies:

- Move decision-making as close as possible to the part of the system which is affected by the decision. Input from higher levels is applied by globally available policy constraints, to avoid the need for directed message passing, since message passing would consume resources.

- High level decisions act as constraints on low level decision making. Authority to act is delegated as far as possible to the part of the system which is affected.

- By definition, high level processes depend on low level ones. Thus high level processes need to search for strategies that are implementable using the low level components.

Too much policy constraint and monitoring from higher levels stifles the freedom of the lower levels to perform their function. Too much communication back and forth leads to an inefficiency.

**Example 159.** A symphony orchestra is a system executed by humans, following mechanical instructions, with a human interpretation. Here the players are technically specialized in their individual instruments, whereas the conductor's job is to look at the broader picture. The conductor does not have detailed skills in the individual instruments, nor does he have the resources to follow every detail of each instrument's part, but he has a unique perspective on the broader picture, which is not available to the individual players, because they are shielded from the full sound by their neighbours. The conductor's role is therefore to support the individual players, and orchestrate their collaboration. Decisions about how to play the instruments and interpret the music are made by the players, with high-level hints from the leadership.

The next example takes a more questionable approach to its task:

**Example 160.** A security monitoring system is a usually collection of cameras and alarms, linked to a control centre, which is monitored by a human. The human can respond to alarms and data from the camera, by locking doors, and by going out to investigate. This reliance on a human manager introduces a communication channel and a dependency for each alarm. If one of these fails, the system could be delayed or fail to function.

\[^3\]This latter assumption, has its roots in history, where the upper classes placed themselves at the helm, and were usually better educated than the working classes who carried out their orders. In our present technological society, the association between education and position has been dismembered and replaced by an association of personalities and interests with position. This tends to invalidate the assumption that high level means more qualified to decide on every issue. What emerges, in a skill based system, is a layered approach to decision making.
13.10.2 Efficiency

Decision making is an example of dependency management. Systems are dependent on decisions being made, and thus the decision making process is an area which needs to be crafted carefully.

From the viewpoint of efficiency, a brief inspection of the figures 6.17 or 6.19 is clear that autonomous decision making, in the individual parts, is the optimal, because it avoids communication and thus introduces no further dependency, or delay. If information has to be passed up or down through the system in order to be analyzed, then significant overhead can be incurred. In human-centric systems, however, it is a matter of policy that decisions are made by ‘management’. Management can have a bird’s-eye view that lower level agents cannot.

Many companies in the technology sector have reorganized themselves from a hierarchical model to an object model, because object models are built around specialization, rather than control (see [TM95]).

Specialized knowledge about the system is found within specialized components at a low level, but the command decisions are usually made from the top. How does the information get from top to bottom to top again? i.e. how does decision making circulate around the system?

A global decision is the most resource consuming kind of decision, usually only required in cases of global policy review.

1. Data collection from all levels.
2. Data interpretation by all levels.
3. Re-examination of goals.
4. Strategy options.
5. Policy adjustment, system changes.

13.11 Prediction, Verification and their Limitations

Maximizing predictability is a key aim of system design. Similarly, one would like to verify that a system design is in accordance with expectation and policy. The idea that systems have well defined states with deterministic behaviour is common in many branches of computer science and has a limited validity. Only simple systems are deterministic. In complex systems, particularly those immersed in an environment, information is being injected into the system from an unpredictable source all the time. This means that absolute determinism is unrealistic. Verifiability is thus not only useful when designing a system, but during its operation, in order to evaluate its behaviour.

There are several levels of verification. To verify correctness in a system that generates non-ordered behaviour, one can use a checksum or hash function to map results to a single scalar value that can easily be compared.
Example 161. MD5 and SHA checksums are used to verify the correctness data transmitted through routed network systems, compared to original control-values, after being reformatted and encapsulated many times to ensure correct transmission.

To compare more structured operations, in which dependent ordering is involved, we use the notion of language and grammar.

Example 162. OSI model network packets that have multi-layer encapsulation have a simple grammatical structure. Each layer of the TCP/IP system has a header, followed by a payload that include the previous layer. The IEEE 802 data-link layer protocols, such as Ethernet, have both a header and a trailer. These encapsulations have a simple grammatical structure that is verified by the unpacking process.

Example 163. The configuration rules of a routing policy protocol, such as BGP, form a structured language of relationships with dependencies. This forms a recursively enumerable graph (for instance, see [GW02, QN03]).

In chapter 5, we reviewed grammars as a way of describing structural complexity in strings of operations. If the syntax of a system, i.e. the list of all legal operation strings, is described by a known grammar then its correctness can easily be verified by an automated procedure or automaton that attempts to parse it. Grammar works as a sophisticated operational checklist for system correctness. Errors in system functioning can easily be identified by comparing the actual behaviour of the system to the legal strings of the language. Correctness can then be evaluated as true or false.

This is the idea behind software engineering tools such as the Unified Modelling Language (UML), which attempts to apply the methods of algorithmic rigour to the complexities of human-computer system interaction. The problem with grammatical structure is that it is only a guide to structure in many systems. The rule based part of a system is often at a low level. The part that resist such formalization is the high level behaviour. An obvious example of this is in biology: at a low level, simple rules of chemistry tell us how molecules fit together, but as we put molecules together to form cells, and cells together to form tissue, and tissue to form organisms, the idea of simple structural rules becomes absurd.

13.12 GRAPHICAL METHODS

Graph theory is a mode of structural expression that is closely related to grammatical methods. It can be used to describe and discuss various aspects of architectural and structural efficiency. For example:

\footnote{In the opinion of the author, UML has been rather unsuccessful in this task. Although it offers a formal framework, it provides no way of dealing with the scales of complexity that are characteristic of actual systems of substance.}
• Connectivity and robustness.

• Optimal constructions
  – Minimum weight spanning trees.
  – Maximum weight branchings.

• Shortest path problems.

These topics find a natural language of expression in graph theory, and there are plenty of theorems and results that can be used to assist in the design and understanding of systems. For an excellent introduction to the graph theory of these issues see [Bal97].

**Applications and Further Study 13.**

• Combining the stuff of the previous chapters into a plan for connecting components.

• Collaboration and partitioning.

• Understanding hierarchy versus flat web-like structures.
CHAPTER 14

SYSTEM NORMALIZATION

Normalization of a system is a process by which one verifies the uniformity or compliance of its goals and promises with respect to guiding principles (see [Dat99]). Normalization is a form of system symmetrization. By making systems from similar normalized units, one hopes to enhance predictability and simplify expectation.

Normalization principles are meant to avoid certain problems of design and operation by reducing complexity, and limit the semantics of system operation as far as possible. This has consequences for maintenance and long term consistency. For instance, systems should not contain elements which actively oppose one another in the performance of a task; such a system would be wasteful of resources, and possibly even harmful.

Data structures form the foundations of any system. The purpose of a data-structure is to catalogue and organize variables, i.e. the changing configurations and records which form the substance of a system. These are then arranged in a pattern which which can be navigated in a manner most conducive to the efficient functioning of the system.

In any system, the law of causality applies. For every effect there must be one or more causes. Determining the causes of an effect becomes increasingly difficult with the increasing complexity of the system. One says that the effect is dependent on the cause.

14.1 DEPENDENCY AND THE SCALING OF AGENCY

In a dynamical process, it is not only the information that has relational patterns, but also the functional components, or ‘subroutines’ of the total process. In either case, guidelines exist for breaking up larger data-structures and processes into smaller, optimal types of parts, the purpose of which is to isolate ‘repeatedly used information’ for optimal re-use and maintenance. In this
way, one avoids unnecessary repetition which can lead to inconsistency and dependency conflicts. Seeking the optimal decomposition into layers is the process referred to as normalization, and is commonly discussed in relational databases (see, for instance, [Dat99]).

**Example 164.** This book is an example of a passive data structure that is not fully normalized. Complete normalization would lead to an obnoxious use of sub-sections and references of the form “see section XX” to avoid repetition and multiple dependency. A fully normalized data structure does not make for easy reading, but it does allow great precision and economy of representation, with potentially no inconsistency.

![Functional structure diagram for a university faculty](image)

Figure 14.1: An excerpt of a functional structure diagram for a university faculty. This shows that, within the different departmental groups, there is a common body of functionality. This repetition suggests that one might rationalize the system by separating this aspect of the groups from the departmental groups’ area of responsibility. This process of rationalizing a structure diagram is called normalization.

A functional dependency is expressed by the mapping of several parameters $k_i$, (usually called keys in database theory), into a single object $O$ of some information space:

$$O = f(k_1, k_2, \ldots, k_n),$$

i.e. we have a relationship between the result $f$ and the combination of keys or parameters which produce that result. We say that $f$ depends on $k_1, k_2, \ldots$, since each combination can yield a
Table 14.1: Comparison of numerical and associative values, through a functional dependency

<table>
<thead>
<tr>
<th>Numerical</th>
<th>Abstract (associative) keys</th>
</tr>
</thead>
<tbody>
<tr>
<td>( y = f(k_1, k_2, k_3) )</td>
<td>lecture = f (teacher, time, subject)</td>
</tr>
<tr>
<td>12 = ( f(14, 3, 6) )</td>
<td>addition = f (Mark, Tuesday, Algebra)</td>
</tr>
</tbody>
</table>

different result for \( f \). The function \( f \), whatever it is, defines a relation. For example, see table 14.1. In continuous functions, parameters can vary along simple lines of numbers, with infinite variety, e.g., a position or a time (see chapter 7). In a discrete structure, such as a database, the parameters are discrete lists, or sets (see chapter 5).

**Definition 59** (Cascade failure). The failure of a strong dependency leads to a succession of determined failures in all agents that rely on that dependency.

Dependencies lead to cascade failures, so it makes sense to try to try to minimize them, or bolster them with redundancy.

**Principle 8** (Weakening dependency (autonomy)). The presence of strong dependencies exposes a system to cascade failure. System designers may weaken dependency relationships to avoid these failure modes by:

- Always having semantic alternatives available for failover redundancy.
- Dynamically repairing dependencies that do not keep their promised function, on a timescale that is faster than the dependency is sampled

Alleviation of dependency is a semantic and a dynamic problem.

### 14.2 THE DATABASE SCHEMA MODEL

The typing of objects is the beginning of addressing the idea of semantics, or interpretation of significance, formally. A relational database (or diagram) is a functional mapping from a set of parameters (lookup keys), to data objects (usually tables or vectors) of the form:

\[
\text{Table} = \tilde{D}(k).
\]  

(14.2)

The vector arrow, denotes the fact that the value returned by the association is assumed to be a table or vector, in the general case, i.e., a bundle of values that are related in some way.

**Example 165.** Many systems have the form of a rudimentary relational database. For example, a building can be thought of as a set of rooms associated with room-numbers. A table of information for each room number (key) might consist of an inventory of the room, who sits there and so on.
A database of computers, with a primary key which is the serial number of the computer, might contain tables of information describing the location, operating system type, name, address and so on. In this latter example, it might make sense to have a sub-database (a set of tables within a set of tables), documenting information about each type of operating system. The key would be the operating system name, and the sub-table might contain the version number, the manufacturer, their service telephone number and so on.

![Figure 14.2: Notations for a functional mapping from k to a table, represented as a table with a key and as a vector with subscript.](image)

If we look at this from the viewpoint of a database, then we envisage the system as a set of entities that are related to one another by functional dependencies. To decompose a system, we ask:

- What are the entities?
  These include computers, services, humans, departments, job positions, etc.
- What are the primary keys that label and classify data in system administration?
  These include things like host names, group names, project names, departments, etc.
- What are the data and other attributes?
  These are specific instances of computers, disks, data, persons and other resources etc.
- Is the system database structure hierarchical?
  We often impose a hierarchical structure on systems, out of fear for loss of control. It is a common myth that the hierarchy is the only reliable model of control.

### 14.3 Normalized Forms

Database normalization seeks to ensure: consistency of data, no hidden dependencies, and the avoidance of internal conflict. These are desirable properties for any system, and we shall see
how the guidelines can be used to learn something about the organization of general systems, by
mapping general systems onto relational databases.

A database is a set of tables (also called vectors), organized into a list, so that each distinct
element is labelled by a unique key, or combination of keys. The keys are the ‘coordinates’ of the
objects in the database. For example, a single key database is simply a list of tables:

$$ Database = \{ T(k) \} = \{ T(1), T(2), \ldots, T(n) \} $$

$$ = \{ \left( \ldots \right)_1, \left( \ldots \right)_2, \ldots, \left( \ldots \right)_n \}.$$  

The point of the normal form is to extract structurally similar components, and repeating
patterns, and place them in separate abstractions with their own labels. This means that the
association is one to one, and that data or entities are not repeated unnecessarily. There are
both practical and aesthetic reasons for this. A practical reason is that, one should not duplicate
information or effort in competing locations, since this would lead to contention, competition and
thus inconsistency. The aesthetic reason is the same as that one uses in programming: repeatedly
useful sub-routines or data are best separated and called up by reference to a new abstract entity,
rather than copying the same code or information in different locations. This makes maintenance
easier and it makes explicit the logical structure of the task.

**FIRST NORMAL FORM**

The first normal form is really a definition about the type of objects one chooses to call a database.
It restricts tables to being simple vectors of a fixed size and shape. The purpose of this definition is
to ensure that all of the objects in the database are comparable. i.e. in order to be able to compare
any two tables in a database, one must be able to compare their contents meaningfully. This is only
possible if the tables are constructed in a similar fashion.

For example, the simplest case a that each vector or table-instance consists only of scalar
values:

$$ T(k) = \begin{pmatrix} s_1(k) \\ s_2(k) \\ s_3(k) \end{pmatrix} = \begin{pmatrix} \text{Algebra} \\ \text{Tuesday} \\ \text{Mark} \end{pmatrix}.$$

Sub-vectors (sub-tables) are allowed as long as they have a fixed, predictable size: for instance, the
following is acceptable because it can easily be rewritten as a larger vector of scalars:

$$ T(k) = \begin{pmatrix} s(k) \\ v(k)[6] \\ v(k)[2] \end{pmatrix}. $$
However, one could not have the following objects in the same database, because they are not comparable objects:

\[
\begin{pmatrix}
  s(k) \\
  v(k)[6] \\
  v(k)[2]
\end{pmatrix}
\neq
\begin{pmatrix}
  s(k) \\
  v(k)[5] \\
  v(k)[9]
\end{pmatrix}
\] (14.7)

The comparison of these objects is meaningless.

What does this have to do with systems in general? We understand that consistency of form is important for stacking data in rows, or for stacking boxes in a warehouse, but what does this mean for system administration? The main thing it tells us is that objects which have a similar structure can be handled by the same system, but objects which are structurally dissimilar should be dealt with by separate systems.

**Example 166.** In spite of the fact that computers are produced by very different manufacturers and use a variety of software, the information about computers within an enterprise is structurally similar: each computer has a serial number, a name, a location and an operating system. The means that a system which deals with these aspects of computers can be handled by a single system. If we also go deeper and consider the details of different software (operating system, for example), we find that Windows and Unix are structurally dissimilar in a number of ways. The normalization rule tells us that it is therefore unnatural to try to combine these into a single system (see fig 14.3).

![Figure 14.3](image-url)

**Figure 14.3:** How normalization by the first normal form suggests that a system should be organized. The common parts belong in a single system of tables, while the structurally dissimilar items are separated into independent sub-systems.
SECOND NORMAL FORM

The second normal form says that, given any 1NF association $\vec{T}(k)$, which has a common repeating pattern, one should extract that pattern and parameterize it with a new label (or key). This is the introduction of ‘database subroutines’ in the data-structure. For example, suppose one has the set of the tables:

$$\left\{\vec{T}(k)\right\} = \left\{ \left( \begin{array}{c} s_1 \\ 1 \\ 1 \end{array} \right), \left( \begin{array}{c} s_2 \\ 1 \\ 1 \end{array} \right), \left( \begin{array}{c} s_3 \\ 1 \\ 1 \end{array} \right), \ldots, \left( \begin{array}{c} s_4 \\ 1 \\ 0 \end{array} \right), \left( \begin{array}{c} s_5 \\ 1 \\ 0 \end{array} \right), \left( \begin{array}{c} s_6 \\ 1 \\ 0 \end{array} \right), \ldots \right\}.$$  \hfill (14.8)

Here we see a repeating pattern. The vectors

$$\left( \begin{array}{c} 1 \\ 1 \\ 1 \end{array} \right), \left( \begin{array}{c} 1 \\ 0 \\ 1 \end{array} \right)$$  \hfill (14.9)

are common to several of the elements. In geometry this is called an invariant sub-space. The second normal form demands that we recognize the importance of this structure, and parameterize it as a vector $\vec{v}[k_s]$ with a new sub-key $k_s$:

$$\vec{v}[1] = \left( \begin{array}{c} 1 \\ 1 \\ 1 \end{array} \right), \vec{v}[2] = \left( \begin{array}{c} 1 \\ 0 \\ 1 \end{array} \right).$$  \hfill (14.10)

The normal form thus transforms elements

$$\vec{T}(k) \rightarrow \left\{ \vec{T}(k; k_s), \vec{v}(k_s) \right\}$$

$$\left( \begin{array}{c} s_k \\ s_k \\ s_k \end{array} \right) \rightarrow \left\{ \left( \begin{array}{c} s_k \\ \vec{v}[k_s] \\ v_{k_s} \end{array} \right), \left( \begin{array}{c} s'_k \\ \vec{v}'[k_s] \\ v'_{k_s} \end{array} \right) \right\}.$$  \hfill (14.11)

In the example above, we note that the pattern is simple, and $k_s$ is varying a third as fast as $k$. This means that the keys are simply related. One could therefore argue that there is really only one key. We could, in principle, repeat this procedure again for sub-tables in sub-tables, and so on, until all of the parameterizations were separated. This is just like the problem of recursively breaking up functions into sub-functions in programming.

The implication of this rule for general systems, is that where-ever we see repeated sub-structures in formally separate systems that are associated with one another, these should be be removed from the separate systems and be replaced by a single, independent instance which can serve both of the systems instead.
Example 167. *The science, engineering and arts faculties of a university all have separate student registration, printing and accounting departments. The task of registering students is the same in all three faculties; the same is true of the financial accounting and printing services. The second normal form therefore suggests that these functions should be removed from the three faculties and be replaced by three independent services: a common registration department, accounting department and printer.*

Example 168. *The science and engineering faculties of the same university all have laboratory engineers who maintain and manage the laboratories. Although these engineers have analogous functions in their respective faculties, their tasks are quite dissimilar, so the normalization rule does not apply here. It would not make sense to group dissimilar laboratory engineers into a common service, because they do not have enough in common.*

**Third normal form**

The third normal form is about parameterizing a structure in a non-redundant way. In geometry, it helps to avoid ambiguity and inefficiency, when traversing vector spaces, by marking out labels using *linearly independent* or *orthogonal* coordinates. Similarly, the third normal form is about securing this kind of independence in discrete, tabular structures.

Inter-dependence of the elements within a vector can occur in two ways. The first is a simple linear dependence, in which two of the scalar values contain common information, or depend on a common value. The second is to avoid making convoluted structures which can feed information back through a chain of relationships, which lead to a cyclic dependency. Consider the table

\[
\vec{T}(k) = \begin{pmatrix} g_1(k) \\ g_2(k) \\ g_3(k) \end{pmatrix}.
\] (14.12)

The third normal form seeks to avoid *transitive dependence*, i.e. a relation of two non-key elements through a third party.

\[
g_1(k) = g_2(g_3(k))
\] (14.13)

Here, two elements \( g_1 \) and \( g_2 \) are not related directly — they do not contain common information directly, but rather they are both functions, derived from a common item of data, thus they are related because one of them depends recursively. This is a non-linear relation.

Another way of putting this is that one attempts to eliminate items of data in any type of object which do not depend on the key directly and explicitly. The presence of a ‘behind the scenes’ relationship, represents an unnecessary symmetry, or a hidden/covert channel from one part of the system to another.
There are other normal forms for even more complex structures, and these can be applied in kind to the simplification of a relational system, but the main points are covered in these three cases.

**Example 169.** In a combined Windows and Unix environment, user data are collected both in the SAM database and in the Unix files `/etc/passwd`. The same data are registered on each and every machine. The user registration data thus form a distributed database, and consist of two different types of data records (tables), containing similar information. One way to normalize this database would be to use an LDAP directory service for both.

**Example 170.** Datacentres and factory processes often try to make infrastructure uniform or ‘normalized’ to bring a predictable symmetry to machinery. Computer racks are organized in regular arrays, with the same numbers and types of hardware in each. Factory machinery is duplicated identically, and desk space is generic facilitating a redundancy to increase the flexibility of the infrastructure to reorganization.

**Denormalized data**

The insistence of normalization of structure, in data and modular design, leads to an essential fragility in systems, which can become problematic at large scale. If we require no deviations from regularity, this becomes a critical dependency. It imposes the responsibility of correctness onto data, which are static, rather than onto software or humans which can adapt. Normalized data are effectively a rigid protocol. This, in turn, makes data corruption, error and other forms of interference into critical dependencies, leading to the possibility of frequent and avoidable failures.

Perhaps for this reason, the popularity of entity relation modelling has waned in modern software systems, and has been replaced by *schemaless* approaches to data structure, in many applications. Schemless key value pairs promise semantic completeness on an *ad hoc* basis.

**Example 171.** NoSQL databases, and interface schemas (so called APIs or Application Programmer Interfaces) allow designers to create data structures with partial occupancy. Data may or may not be regular in form. What matters is that key-value pairs have unique names so that the semantics of data may be determined from the key, rather than from assumptions about the schema. This is one strategy in making systems of data dependency fault tolerant. New data formats like JSON and YAML have made this ad hoc approach to data semantics easy to adopt, through standard interchange formats, that contrast with more rigid schema formats like XML and YANG.
14.4 Promise Theory, Semantic Spacetime, and Uniformity

The description of data and structural normalization have been addressed by a model called semantic spacetime, using the language of promise theory [Bur14, Bur15, Bur16b], in which agents collectively form a ‘space’. In this approach, one can discuss and evaluate the value of different representations, without requiring normalization by convention. The normalization of agents depends on scale, because some agents may be inside clusters that act as ‘superagents’. This model has been used as a unifying approach not only to describe human-computer systems, but also knowledge representations (see chapter 17). The topic is too extensive to describe here. Readers are referred to the references above. The semantic spacetime model offers a formulation of database schemas and other semantic structures in a simple graph theoretic form, eliminating the need for a specialized theory.

Applications and Further Study 14.

- Rationalizing the entities and players in a system for planning and design. Rational decision criteria for segregating processes and information conduits.
CHAPTER 15

SYSTEM INTEGRITY

Integrity is about the preservation of a system’s policy, its resources, and its ability to function consistently. Integrity can be dealt with easily in the context of promises. A system that makes clear promises can be checked for compliance over time, and an absence of change in its compliance may be taken as a measure of its integrity. System integrity can also be couched in terms of the communication of those assets from place to place, or from time to time, using information theory; however, transfer of information is not always reliable, and promises are not always kept.

The theory of communication, and its generalization ‘information theory’, seek to answer questions such as how rapidly or reliably the information from the source can be transmitted over a channel. Insofar as we can define the code by which the work in a human-computer system can be transmitted, we can use the tools of information theory to describe the possibility of flow or corruption of those assets. Shannon’s work is significant, because it addresses fundamental limits on the ability to resolve and preserve information. He addressed the problem of coding of symbolic information, in such as way as to maximize the rate and fidelity of transmission, given that the representation might change. He also achieved considerable success with his technique of random coding, in which he showed that a random, rather than a sequential, encoding of data can, with high probability, give essentially optimal performance.

15.1 SYSTEM CONVERGENCE TO A DESIRED STATE

The process of communication is essential in any information system. We find the promises of outcome in a variety of forms:

- Between computer programs and their data,
• Between computers and devices,
• Between collaborating humans (in teams),
• Between clients and servers,
• Between computer users and computer systems,
• Between policy decision makers and policy enforcers,
• Between computers and the environment (spilled coffee).

The intent of these communications is constantly being intruded upon by an environmental noise. Errors in this communication process occur in two ways:

• Information is distorted, symbols are changed, inserted or omitted, by faulty communication, or by external interference,

• Information in interpreted incorrectly; symbols are incorrectly identified, due to imprecision or external interference.

For example, suppose one begins with the simplest case of a standalone computer, with no users, executing a program in isolation. The computer is not communicating with any external agents, but internally there is a fetch-execute cycle, causing data to be read from and written to memory, with a CPU performing manipulations along the way. The transmission of data, to and from the memory, is subject to errors, which are caused by electrical spikes, cosmic rays, thermal noise and all kinds of other effects. These errors are normally corrected by error-correction mechanisms, originating from Shannon’s work. The computer program itself, manipulates the data in the memory and rewrites it to the memory with a new coding.

From this point of view, one may think of the memory of the computer itself as being both transmitter and a receiver, and passing through the ‘CPU plus computer program’ communication channel. This communication channel does not transmit the data unaltered; rather, it transforms the data according to specific rules, laid down in the computer program (see fig. 15.1). In other words, the very operation of a computer fits the paradigm of communication over a coded channel. The fact that the channel is also noisy, is a result of the physical environment. Computer operation, at this level, is largely immune to environmental noise, because it employs error correction methods. At higher levels, there are no standardized error correction mechanisms in common usage.

Suppose now that an administrator sends a configuration message to a host, or even to a single computer program. Such a message takes place by some agreed form of coding: a protocol of some kind, e.g. a user interface, or a message format. Such a configuration message might be distorted by errors in communication, by software errors, by random typing errors. The system itself might change, during the implementation of the instructions, due to the actions of unknown
parties, working covertly. These are all issues which contribute uncertainty into to the configuration process and, unless corrected, lead to a ‘sickness’ of the system, i.e. a deviation from its intended function.

15.2 The Shannon Error Channel Interpretation of Integrity and Convergence

The idea of convergence is introduced in section 10.4, to describe the behaviour of such error-correction. It suggests a process of continual regulation, in order to correct deviations from system policy. In other words, it does not simply ensure that a configuration message is transmitted correctly once, it views the entire time-development of the system as the on-going transmission of the message, and seeks to correct it at every stage. This is also our definition of a reliable system (see section 4.11).

![Figure 15.1: An instruction loop, showing the development of a computer system in time, according to a set of rules. The efficacy of the rules may be distorted by users from local and remote domains, who change the conditions under which the message was applicable. This change may be viewed as an intentional change, or as a stochastic error.](image)

At each level of computer operation, one finds messages being communicated between different entities. System administration is a meta-program, executed by a mixture of humans and machines, which concerns the evolution and maintenance of distributed computer systems. It involves:

- Configuring systems within policy guidelines,
- Keeping machines running within policy guidelines,
• Keeping user activity within policy guidelines.

System administration requires computer-computer interaction, human-computer interaction, and human-human interaction (fig. 15.2). Each of these communication channels is subject to error, or misinterpretation.

Figure 15.2: The human-computer interaction is also a form of communication. Collaboration between system administrators, or users is the least reliable form of communication, since one can not be sure that the two parties even use the same symbolic alphabets. Thus, there is the danger not only of noise, but also of misunderstanding, i.e. perversion of meaning.

• **Instruction**: system administration is about developing a policy (instruction manual for machine and humans) for use of the system, including the choice of programs which are executed by the system. A complete policy can therefore be identified with the sum of all programs and ad hoc rules which lead to any planned change of the system. This includes normal usage of the system.

• **Propagation**: as the system evolves in time, according to policy (or deviating), it propagates information from the past into the future, modifying the system. If the system is stable, this iterative mapping will not lead to any divergent behaviour; if it is unstable, then even a small error might cause a runaway breakdown of the system.

• **Collaboration**: programs and humans exchange information in the course of this loop of instruction and propagation. If collaboration is interrupted, or errors occur, then the
CHAPTER 15. SYSTEM INTEGRITY

Noise
Sender Receiver

Figure 15.3: The transmission of information over a channel takes place from a sender (channel input) to a receiver (channel output). Along the way, the signal might become distorted by noise.

enactment of policy-correct behaviour is spoiled, with possibly dangerous consequences for the system. Humans also frequently misunderstand one another’s commands.

• Automation: automatic processes which monitor systems, even perform routine maintenance, are not immune to errors, because they depend, for input, on data which are influenced by the external environment.

• Repair: if some influence causes an error, then the error must be corrected in order to uphold policy, else the correct propagation of policy over time is corrupted. The same thing applies to human policy and purely automated policy. This process of maintenance, reparation, or regulation is central to the stability of information systems.

To summarize, one may view system administration as communication over a communications channel at several levels.

• Input: policy, instruction.

• Noise: stochastic user activity, illegal behaviour, random error, systematic error.

• Output: the system as we see it.

Having made this identification, the question becomes: does this help us to build computer systems which behave in a stable and predictable fashion?

15.3 EXTERIOR INFLUENCE AND STRATEGIC INSTRUCTION

The transmission of configuration information requires a language which, in turn, requires an abstract alphabet or set of codes to encode it. This ‘alphabet’ might be one of strings or of shortest length symbols. The information content will tell us how compressible the actual transmitted configuration is and therefore how concisely we can express system policy.

Each word of symbol in the language must represent a basic operation of the system like “create file” or “insert string” etc. These operations can be represented as single letter codes, with
accompanying data (like opcodes in machine language), or they could be packed into lengthy XML wrappers to name two extremes.

**Example 172.** Suppose the operation to change the permissions of a Unix file (chmod) is coded in a data stream by the letter “A”, and the operation to change the owner of a file (chown) is “B”. The operations need both data parameters and operands. A configuration policy can be written in any language that the system understands, e.g.

<table>
<thead>
<tr>
<th>Human symbol</th>
<th>Compressed code</th>
</tr>
</thead>
<tbody>
<tr>
<td>chmod</td>
<td>A</td>
</tr>
<tr>
<td>chown</td>
<td>B</td>
</tr>
<tr>
<td>700</td>
<td>a</td>
</tr>
<tr>
<td>770</td>
<td>b</td>
</tr>
<tr>
<td>755</td>
<td>c</td>
</tr>
<tr>
<td>644</td>
<td>d</td>
</tr>
<tr>
<td>600</td>
<td>e</td>
</tr>
<tr>
<td>555</td>
<td>f</td>
</tr>
</tbody>
</table>

So, if we number files according to their filesystem entries (e.g. index node number), the command to set permissions on file 12 would be A d (12) or A (12) d, where (12) represents a suitable shortest representation of a number 12. This set of symbols will suffice for a limited number of operations.

The amount of information that has to be specified in order to express a configuration depends on the expert knowledge of the system receiving the instruction. There are two approaches to this:

- If we build expert operators that know how to do their jobs and only require a simple signal to activate them, then configuration policy can be written in a very short and compact form, e.g. as in the single letter codes above. This reflects the fact that the detailed procedures are coded into operators and therefore do not need to be reiterated in every configuration message. (This strategy allows maximal compression and optimal normalization of information, since there is only one copy of the expertise in the operators. Thus a single symbol can represent an expert operation.)

- If the operations contain no internal expertise, then each precise sequence of primitive operations must be expressed in the policy message. This, in principle, involves a precise specification with redundant information.

**Example 173.** A compressed message to an expert operator providing maintenance might take the form

CheckAndRepair(routine_1)
An extensive form of the instruction could take the form

Locate panel screws
Rotate screws anti-clockwise
Remove screws
Lift lid
Locate memory slot 12
Insert new memory into slot 12

In the latter form, the detailed procedure is described in the message; in the form case the procedure is coded into the operators themselves, and only a short message needs to be passed on to start the operators executing their policy instruction.

An example of the former is found in refs. [Bur95, CD01] and examples of the latter include the Simple Network Management Protocol (SNMP), Arusha ([HP01]), etc.

Example 174. The human representation symbol alphabet used by cfengine, which describes information in a plain text file, consists of all of the printable ASCII symbols; the set of symbols required to implement a policy decision about file permissions is the set of all rules (one for each item), which for Unix might look like this:

files:

# symbol 1

file1 mode=0644 owner=mark group=users action=fix

# symbol 2

directory1 mode=a+rX owner=root recurse=true action=fix

In this case, each entire rule can be a single symbol of the higher level policy alphabet, and when it is coded in this fashion, since the number of variations is finite. This short symbolic coding of policy, is robust to accidental or random error, and is easy to reapply (retransmit), should external factors alter its result.

The symbol objects that represent new effective entities of the system. They form new and higher alphabets of preferably non-overlapping objects. To perform configuration management, we need to reiterate this configuration message over time, correcting for any random error. We now examine how to characterize the error in a stream of these symbols.
CHAPTER 15. SYSTEM INTEGRITY

GENERATIVE CONFIGURATION

Conventional wisdom suggests that, when an instruction for building configuration state is specified, the order of operations is important to the outcome of the configuration (see arguments in [Tra02], for instance). However, an alternative prescription is for configuration is based on the idea of expert operators and convergence (see section 5.8). We refer to these two alternative forms of instruction as extensive and strategic respectively, and keep the property of convergence towards an ideal state separate, since it can in principle be implemented by either approach.

- **Extensive configuration instruction**

  In the extensive approach, each individual decision in the configuration state is represented as part of a tree with exponential complexity. If the number of symbols is $N$ and the alphabet size is $m$, then the amount of information that must be maintained is of the order $m^N$.

- **Strategic configuration instruction**

  In a strategic approach, the decision trees are built into the properties of the operators that carry out the maintenance (see [Bur04, CD01, CS03]). Here the complexity of the configuration is at most of order $N^2$, and the information represented is no larger than $N$. The ordering of these operators is not essential, provided the configuration message is repeated over and over again as regular maintenance, since the operators measure their activities relative to both desired policy and current environment. If any operation depends on another having preceded it, it simply waits until the necessary conditions exist for it to proceed. in order for this to work, the operations must be quite primitive.

ORDERING OF PREREQUISITE DEPENDENCIES

There is a connection between the ordering of operations and the uniqueness of the task completed by a schedule (see section 9.8, for instance). The information required to perform a schedule depends on whether task precedence matters (i.e. whether the graph is directed or not). This, in turn, depends on whether the task alphabet commutes or not:

**Definition 60** (Commuting operations). If two system operation codes commute then

$$[\hat{O}_1, \hat{O}_2] \equiv \hat{O}_1 \hat{O}_2 - \hat{O}_2 \hat{O}_1 = 0. \quad (15.1)$$

If operators commute, it implies that the order of their execution does not matter to the system. This requires special properties and is seldom true in every instance, since ordering reflects the structure that distinguishes a system from a random assembly of components. However, there is a possible solution to this using the concept of orthogonality.
Contrary to many expectations, most simple configuration tasks can be performed convergently, by randomly or cyclically scheduling non-ordered, commuting operations. This can be done by making operation orthogonal (see [Bur04, CS03]).

**Definition 61 (Orthogonal operators).** An operation is said to be orthogonal to all other operations, if it is inequivalent to any combination of other operations, i.e. any representation of the operation is linearly independent of all others:

\[
\hat{O}_i \neq \sum_{j \neq i} c_j \hat{O}_j,
\]  

for some constants \(c_j\). Orthogonal operations are automatically commutative and ordering does not matter.¹

¹This definition, as given, describes strictly linear independence. Orthogonality implies that an inner product of representative basis vectors would vanish. It is possible to find a representation in which this is the case, but we shall not go into it here. See ref. [Bur04] for a discussion of an explicit matrix representation.

Consider how configurations are built up. Let the state \(|0\rangle\) denote the base-state of a host after installation. This is a reference state to which any host may be returned by re-installation. From this state, one may build up an arbitrary new state \(|a, b, c, \ldots\rangle\) through the action of sequences of the configuration operations. The set \(a, b, c\) may be regarded as the system policy specification. Once a desirable state has been reached, one may renormalize these definitions to allow \(|0\rangle\) to be the new base-state. Using this representation, one can now define the meaning of convergence in terms of these operations.

**Definition 62 (Convergent and idempotent operators).** Let \(|s\rangle\) be an arbitrary state of the system. An operator \(\hat{O}\) is idempotent if \(\hat{O}^2 = \hat{O}\), i.e. its potency to operate is always the same. A convergent operator \(\hat{C}_\alpha\) has the more general property

\[
(\hat{C}_\alpha)^n |s\rangle = |0\rangle, \\
\hat{C}_\alpha |0\rangle = |0\rangle,
\]  

for some integer \(n > 1\), i.e. the \(n\)-th power of the operation is null-potent, and the base state is a fixed point of the operator.

In other words, a convergent operator has the property that its repeated application will eventually lead to the base state, and no further activity will be registered thereafter.

The use of orthogonal, convergent operations implies that only one type of prerequisite dependency can occur. For example, let \(\hat{C}_C\) mean ‘create object’ and let \(\hat{C}_A\) mean ‘set object
attribute'. The following operations do not commute, because the setting of an attribute on an object requires the object to exist. On an arbitrary state $|s\rangle$, we have

$$[\hat{C}_C, \hat{C}_A]|s\rangle \neq 0.$$  

(15.4)

Thus the ordering does indeed matter for the first iteration of the configuration tool. This error will, however, be automatically corrected on the next iteration, owing to the property of convergence. To see that the ordering will be resolved, one simply squares any ordering of the above operations.

**Theorem 4.** The square of a create-modify pair is order independent.

$$([\hat{C}_C, \hat{C}_A]|s\rangle)^2 = 0.$$  

(15.5)

This result is true because the square of these two operators will automatically result in one term with the correct ordering. Orderings of the operators in the incorrect order are ignored due to the convergent semantics.

To prove this, suppose that the correct ordering (create then set attribute) leads to the desired state $|0\rangle$:

$$\hat{C}_A\hat{C}_C|s\rangle = |0\rangle;$$  

(15.6)

performing the incorrect ordering twice yields the following sequence:

$$\hat{C}_C\hat{C}_A\hat{C}_C\hat{C}_A|s\rangle = |0\rangle.$$  

(15.7)

The action of $\hat{C}_A$ has no effect, since the object does not exist. The under-brace is the correct sequence, leading to the correct state, and the final $\hat{C}_C$ acting on the final state has no effect, because the system has already converged.

The same property is true of sequences of any length, as shown in ref. [CD01]; in that case, convergence of $n$ operations is assured by a number of iterations less than or equal to $n$.

**Theorem 5.** A sequence of $n$ self-ordering operations is convergent in $n$ iterations, i.e. is of order $n^2$ in the primitive processes.

The proof may be found by extending the example above, by induction (see [CD01]).

The important thing about this construction is its predictability. We might not know the exact path required to bring a system into a policy conformant state; indeed, a given specification might meet obstacles and fail to work. However, any policy expressed entirely in terms of convergent, commuting operators is guaranteed to do something, indeed it will always have the same result$^1$.

---

$^1$This does not mean that every possible state is necessarily reachable by convergent, commuting operators, but we are suggesting that any states not reachable by this approach represent inappropriate policies.
Commutation becomes not only a desirable property, but an essential one in ensuring predictability. An extensive approach is not guaranteed to be implementable or stable once implemented, but a convergent strategic approach is (see [Bur04, CD01, CS03] for a proof).

One of Shannon’s conclusions was that, if one splits up a signal into a coding scheme which is randomized in ordering, then many this is a good defence against random noise, because any systematic error will be reduced to maximum entropy random error, rather than concentrated in one area of greater damage. The advantage of commuting operations is that they do not have to rely on a particular sequence being fulfilled in order to produce a result. A string of commuting operators can thus often be compressed in communication, because the intelligence lies in the operator rather than in the sequence of codes: a short code for a complex task, rather than a detailing of the internals of the task.

15.4 STOCHASTIC SEMI-GROUPS AND MARTINGALES

The discussion of convergent operations has an analogue in the mathematics of semi-groups, or transformations that make transitions in only one parameter direction. If there are \( n \) states \( \{ q \} \), then a transition \( T(t; q, q') \) from a state \( q \) to a state \( q' \), at time \( t \), is an \( n \times n \) matrix that acts on a state vector \( \vec{q}(t) \).

**Definition 63** (Semi-group of transformations). The set of transformations \( \hat{T} \in \{ T(t; q, q') \} \) is said to form a semi-group (for \( t \geq 0 \)) if:

1. \( \hat{T}_0 = I \) is the identity transformation that leaves a state vector invariant: \( \hat{T}_0 \vec{q} = \vec{q} \).
2. \( \hat{T}_t \) satisfies the Chapman-Kolmogorov equation:

\[
\hat{T}_{s+t} = \hat{T}_s \hat{T}_t, \quad s, t \geq 0.
\] (15.8)

A stochastic semi-group is a group of stochastic (probabilistic) transitions (see [GS01]). These groups are clearly of central interest to configuration management. For a discussion ‘couched’ in these terms see [CS03].

Another probabilistically convergent process is a **martingale**. The study of martingales is the study of sequences that converge in the sense of their total value. The term martingale comes from gambling.

Suppose a gambler has a fortune. She wagers 1 Euro on an evens bet. If she loses, she wagers 2 Euros on the next play, and \( 2^n \) on the \( n \)th play. Each amount is calculated to cover her previous losses. This strategy is called a martingale.
**Definition 64** (Martingale). A sequence $S_n$ with $n \geq 1$, is a martingale with respect to another (reference) sequence $t_n$, $n \geq 1$ if, for all $n \geq 1$:

1. $\langle |S_n| \rangle < \infty$,
2. $\langle S_{n+1}|t_1, t_2, \ldots, t_n \rangle = S_n$.

Note that, we define a reference sequence that is usually the progression of time (ticks of a clock), but any synchronizing pulse sequence would do.

Martingales always converge, in the following sense. If $S_n$ is a martingale with $\langle S_n \rangle < M < \infty$, for some $M, n$, then there exists a random variable $S$ such that $S_n \to S$, as $n \to \infty$. This is a form of statistical convergence, as discussed in section 10.9. The convergence time of a martingale is a topic of particular interest, that is beyond the current text.

### 15.5 CHARACTERIZING PROBABLE OR AVERAGE ERROR

The measures of entropy introduced in chapter 9 provide a precise characterization of how much uncertainty there is in the processes of propagating any message over any channel. We now have a configuration alphabet that needs to be propagated into the future, or reproduced for repair or backup purposes. We apply the measures of informational uncertainty from chapter 15 to measure the amount of information that must be transmitted.

**Example 175.** You decide that, in addition to backing up your user data, you should also backup your users by copying their DNA. This is rather straightforward, since DNA is written with only a four symbol alphabet and can be performed by almost any cell. On observing the result of cell division, you find the following data for the probability of successful copying. There seems to be a problem with the fidelity of the copying:

<table>
<thead>
<tr>
<th>Trans/Recv</th>
<th>A</th>
<th>C</th>
<th>T</th>
<th>G</th>
<th>Marginal</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>$\frac{1}{8}$</td>
<td>$\frac{1}{16}$</td>
<td>$\frac{1}{32}$</td>
<td>$\frac{1}{32}$</td>
<td>$\frac{1}{4}$</td>
</tr>
<tr>
<td>C</td>
<td>$\frac{1}{16}$</td>
<td>$\frac{1}{8}$</td>
<td>$\frac{1}{32}$</td>
<td>$\frac{1}{32}$</td>
<td>$\frac{1}{4}$</td>
</tr>
<tr>
<td>T</td>
<td>$\frac{1}{16}$</td>
<td>$\frac{1}{16}$</td>
<td>$\frac{1}{16}$</td>
<td>$\frac{1}{16}$</td>
<td>$\frac{1}{4}$</td>
</tr>
<tr>
<td>G</td>
<td>$\frac{1}{4}$</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>$\frac{1}{4}$</td>
</tr>
<tr>
<td>Marginal</td>
<td>$\frac{1}{2}$</td>
<td>$\frac{1}{2}$</td>
<td>$\frac{1}{8}$</td>
<td>$\frac{1}{8}$</td>
<td>$\frac{1}{2}$</td>
</tr>
</tbody>
</table>

Note that we also calculate the marginal distributions here. The marginal distributions are found
by summing the rows or columns. The dual distribution is thus

\[ p(T, R) = \begin{pmatrix}
\frac{1}{8} & \frac{1}{16} & \frac{1}{32} & \frac{1}{32} \\
\frac{1}{16} & \frac{1}{8} & \frac{1}{32} & \frac{1}{32} \\
\frac{1}{16} & \frac{1}{16} & \frac{1}{16} & \frac{1}{16} \\
\frac{1}{4} & 0 & 0 & 0
\end{pmatrix} \]  

(15.9)

Had the communication been perfect, this would have been:

\[ p(T, R)_{\text{perfect}} = \begin{pmatrix}
\frac{1}{4} & 0 & 0 & 0 \\
0 & \frac{1}{4} & 0 & 0 \\
0 & 0 & \frac{1}{4} & 0 \\
0 & 0 & 0 & \frac{1}{4}
\end{pmatrix} \]  

(15.10)

As it is, however, this diagonal perfection is smeared out in both directions. We note that there is a persistent error in the channel that causes a “G” to be received as an “A”.

From the marginal distributions, we have

\[ H(R) = -\frac{1}{2} \log_2 \frac{1}{2} - \frac{1}{4} \log_2 \frac{1}{4} - \frac{1}{8} \log_2 \frac{1}{8} - \frac{1}{8} \log_2 \frac{1}{8} \]

\[ = \frac{1}{2} + \frac{1}{2} + \frac{3}{8} + \frac{3}{8} \]

\[ = \frac{7}{4} \text{ bits.} \]  

(15.11)

\[ H(T) = 2 \text{ bits.} \]  

(15.12)

The conditional distributions are

\[ H(R|T) = \sum_{i=1}^{4} p(T=i)H(R|T=i) \]

\[ = \frac{1}{2} H\left(\frac{1}{8}, \frac{1}{16}, \frac{1}{32}, \frac{1}{32}\right) + \frac{1}{2} H\left(\frac{1}{8}, \frac{1}{16}, \frac{1}{32}, \frac{1}{32}\right) \]

\[ + \frac{1}{4} H\left(\frac{1}{16}, \frac{1}{16}, \frac{1}{16}, \frac{1}{16}\right) + \frac{1}{4} H\left(\frac{1}{4}, 0, 0, 0\right) \]

\[ = \frac{11}{8} \text{ bits.} \]  

(15.13)

\[ H(T|R) = \frac{13}{8} \text{ bits.} \]  

(15.14)

\[ H(R, T) = \frac{27}{8} \text{ bits.} \]  

(15.15)

The interpretation of the numbers is as follows. \( H(T) \) is the uncertainty per symbol in our ability to describe the message that is transmitted made available for copying, as what the next symbol
is likely to be. $H(R)$ is the uncertainty per symbol in our ability to describe the message that is received, i.e. to predict the occurrence of each symbol in a stream. $H(R|T)$ is the uncertainty per symbol in the received message’s integrity, given that the transmitted message is known. This is the main quantity of interest to us: it characterizes the likely integrity of the copy, given that the original is completely known.

To get some idea of how high this uncertainty is, there are $\log_2 4$ bits per symbol i.e. 2 bits per symbol \{A, C, G, T\}. Thus, the uncertainty in the original transmitted message $H(T)$ is maximal; no one symbol occurs more frequently than any other, so we cannot say anything about the original copy to compress it or to simplify it. We have to know the exact message in order to copy it.

The uncertainty in the received message is actually less than this. Why? Because the copying is imperfect and it biases the message in a systematic error (see $p(R, T)$). The uncertainty in the copy, given that the original is known exactly $H(R|T)$ is about half a symbol per symbol! This is a very high probability of error, far from appropriate for backing up our users\(^2\).

### 15.6 Correcting Errors of Propagation

One of Shannon’s accomplishments was to prove that any communications channel has a limited capacity, and that it is possible to find a coding scheme which achieves reliable transmission of symbols with an efficiency which is arbitrarily close to that channel capacity. This is known as Shannon’s theorem. Shannon imagined a generic model for transmission over a noisy channel, shown in fig. 15.4. This model can also be applied to the transmission of policy, or system integrity.

![Figure 15.4: Shannon’s view of the correction of data on a coded channel, can also be applied to the correction of policy propagation, or system ‘health’. It makes clear the need for a correctional system.](image)

\(^2\)It has come to the author’s attention that not all reader possess a sense of humour. For the record, it should be pointed out that backing up users is not an ethical procedure. Please do not try this at home.
Errors creep into the transmission of rules and actions, even with digital channels. One of the reasons for introducing graphical user interfaces to computers, for instance, was to reduce the possibility of error, by condensing difficult operations into a few simple symbols for inexperienced users. In spite of the effort, users still hit the wrong icon, or menu item, and the simplifications which are assumed do not always apply, because external conditions invalidate the assumptions that were made in selecting the symbols.

Shannon’s theorem, however, tells us that a suitable coding scheme (protocol) can assure the transmission of a system policy within arbitrary accuracy. Windowing systems use error correction protocols, such as ‘Are you sure?’ dialogue boxes, in order to catch random errors — not usually because they believe that users will change their minds very often.

The signal we wish to propagate is this:

\[
\text{Ideal signal} = p(\text{usage} | \text{policy})
\]  

(15.16)

Policy can be communicated by declaration (see, for instance, [Bur95, DDL00]), or by simulated dialogue ([Lib90, Cou00]). The latter is becoming more common, with technologies like SNMP ([OBC99]) and XML based grid services ([XR]). In either case, its transmission or implementation through fallible humans and technologies is subject to the incursion of ‘noise’.

The issue now, at this high level, is more subtle than for simple bits however. Do we really want to achieve such a level of precision as to lead to no errors at all? As long as humans are part of the equation, there is the question of user-comfort and human welfare.

Thus, in order to apply error correction to larger social and ecological systems, we must choose the symbols rather carefully. Unlike Shannon, we must pay attention to the semantic content of symbols when formulating policies which one wishes to have enforced (see, for instance, [Zad73]).

Proactive error correction is one way of dealing with this issue: by requiring confirmation of a policy breach, one can avoid spurious over-reactions to acceptable transgressions. For example, the double keys used in nuclear missile launches, or the double signals used in co-stimulation of the immune response, are simple security features which prevent potentially damaging responses to innocent errors. Hamming codes and checksum confirmation are other examples of this type of protocol coding.

### 15.7 Gaussian Continuum Approximation Formula

It is not always convenient or appropriate to provide a complete description of transmission joint probabilities in the form of a matrix. If the number of symbols is effectively infinite, i.e. of the signal varies as an arbitrary real number, rather than as a digital signal, then the characterization of probable error must be performed in terms of functions or distributions rather than matrices. If we believe that a transmission channel is basically reliable, but with a quantifiable source of
random error, then it is useful to use a simple continuum approximation model for the expected error. The expression for the capacity of a channel with Gaussian noise is one of the classic results in information theory, and has many applications. Consider the probability distribution

\[ p(q) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left(-\frac{q^2}{2\sigma^2}\right). \]  

(15.17)

We have:

\[ -\ln p(q) = \ln \sqrt{2\pi\sigma^2} + \frac{q^2}{2\sigma^2}, \]

(15.18)

and thus entropy

\[ H(q) = -\int p(q) \ln p(q) dq \]

\[ = \ln(\sqrt{2\pi\sigma^2}) \int p(q) dq + \int p(q) \frac{q^2}{2\sigma^2} dq, \]

\[ = \ln(\sqrt{2\pi\sigma^2}) + \frac{\sigma^2}{2\sigma^2} \]

\[ = \frac{1}{2} \ln(2\pi e\sigma^2). \]  

(15.19)

Now consider a time series \( q(t) \) to be a series of real numbers measured at arbitrary times, and let us consider the total system to be a mixture of an average signal \( \langle q(t) \rangle_s \) and a noise term \( \delta q(t) \); i.e.

\[ q(t) = \langle q(t) \rangle_s + \delta q(t). \]  

(15.20)

and \( \langle \delta q \rangle = 0 \). We shall assume Gaussian noise so that eqn. (15.17) applies, and obtain the classic result due to Shannon ([SW49, CT91]). The information that can be transmitted by this continuous channel is infinite, in principle, since we can transmit any real number at any time with no error. However, this is not a realizable situation, since there is a physical limit to the information that can be inserted electrically or optically into a physical channel. It is thus normal to calculate the mutual information that can be transmitted given that there is an upper bound on the average signal power \( P = S + N \), where \( S \) is the signal power and \( N = \sigma^2 \) is the noise power. The power varies like the signal squared, so we apply the constraint:

\[ \frac{1}{\Delta T} \int_0^{\Delta T} q^2(t) dt \leq P. \]  

(15.21)

If we exceed the maximum power, the channel could melt or be destroyed.

The channel capacity is defined to be the maximum value of the mutual information in the average signal, given this power constraint:

\[ C(1) = \max_{\langle q \rangle} H(\langle q \rangle; q). \]  

(15.22)
This tells us how many digits per sample we are certain are being transmitted, since it is the fractional number of digits of information required to distinguish the average string that was transmitted over the communications channel. In this case we feed the locally averaged signal $\langle q \rangle$ (the smooth part of the noisy signal that we are responsible for), and we extract the full noisy signal $q$ at the output.

$$H(\langle q \rangle; q) = H(q) - H(q|\langle q \rangle)$$
$$= H(q) - H(\langle q \rangle + \delta q|\langle q \rangle)$$
$$= H(q) - H(\delta q|\langle q \rangle)$$
$$= H(q) - H(\delta q). \quad (15.23)$$

The last line follows from the independence of $\langle q \rangle$ and $\delta q$. Now, for the Gaussian channel, we have

$$H(\delta q) = \frac{1}{2} \ln(2\pi e\sigma^2) = \frac{1}{2} \ln(2\pi eN). \quad (15.24)$$

Thus

$$H(\langle q \rangle; q) = H(q) - H(\delta q),$$
$$\leq \frac{1}{2} \ln(2\pi e(S + N)) - \frac{1}{2} \ln(2\pi eN).$$
$$= \frac{1}{2} \ln \left(1 + \frac{S}{N}\right). \quad (15.25)$$

To reconstruct the signal we must sample it at twice its maximum frequency, by the Shannon-Nyquist sampling theorem; thus the channel capacity of a channel of parallel bands of width $B$ cycles per second is,

$$C(B) = \sum_{n=1}^{2B} C(1)$$
$$C(B) = B \log_2 \left(1 + \frac{P_S}{P_N}\right) \quad (15.26)$$

where $P_S$ and $P_N$ are the total power in all the band frequencies.

**Example 176.** An Bangalore Internet Service Provider has a critical copper Internet cable with bandwidth 2GHz to share amongst customers. This cable is carried on pylons through the city. Calculate the reduction in usable channel capacity $C$ due to thermal noise (power proportional to $kT$), if the temperature changes from 15 degrees Celsius to 35 degrees Celsius, given that the signal to noise ratio is 50dB at 15 degrees Celsius.

The signal to noise ratio is defined by

$$10 \log_{10} \left(\frac{P_S}{P_N}\right) = 50,$$  \quad (15.27)
hence

\[
\left. \frac{P_S}{P_N} \right|_{T=(273+15)} = 10^5. \tag{15.28}
\]

Given that the noise power is proportional to absolute temperature, at 35 degrees Celsius the signal to noise ratio is thus

\[
\left. \frac{P_S}{P_N} \right|_{T=(273+35)} = 10^5 \times \frac{273 + 15}{273 + 35}. \tag{15.29}
\]

The capacity of the cable is thus

\[
C(T = 15) = 2 \times 10^9 \log \left(1 + 10^5 \right) \tag{15.30}
\]

\[
C(T = 35) = 2 \times 10^9 \log \left(1 + \frac{10^5 (273 + 15)}{(273 + 35)} \right) \tag{15.31}
\]

\[
C(T = 15) - C(T = 35) = 2.9 \times 10^7 \tag{15.32}
\]

Thus there is a loss of channel capacity of about 30 Megabits per second.

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**Applications and Further Study 15.**

- Relating expectation about a system to observed behaviour.
- A method of describing errors (incompatible events) that occur in system processes and information flows.
- A method of quantifying losses.
CHAPTER 16

POLICY AND MAINTENANCE

When we speak of policy, we really mean a way of defining and constraining the behaviour of a system to keep certain promises. A policy description must include the configuration of a system and offer guidelines as to how it should be allowed to evolve. Policy and configuration management are thus aspects of the same thing. We know that random fluctuations will always lead to some changes that do not agree with policy, so fluctuations themselves cannot be made policy conformant. This implies that we need something to keep fluctuations from permanently altering a system.

16.1 WHAT IS MAINTENANCE?

When does a process become a maintenance process? The notion of system administration is closely allied with that of maintenance. We need some general notion of maintenance that can be described quantitatively. Maintenance is a process that tends to oppose fluctuation — i.e. minimize short term change and provide medium term stability. We are not interested in managing systems that cannot achieve a basic level of stability, since these cannot perform any reliable function.

In this chapter we shall think of maintenance as a response to a stochastic process. There is a parallel here to Shannon’s discussion of communication theory ([SW49]) discussed in chapter 9. To overlay the language of stochastic systems onto the maintenance process, we need to make a separation into what is policy conformant and what is anomalous. The meaning of conformant and anomalous is not automatically clear, but if fluctuations have finite variance, then it is self-consistent to associate these concepts with slowly and rapidly varying changes, respectively, measured in relation to user-behaviour (see [BHRS01]).
Chapter 16. Policy and Maintenance

The separation of slow and rapid changes to configurations can be made precise by observing the system through a local averaging procedure. We shall refer to the schematic diagram in fig. 16.1.

Suppose that the total configuration $C = \{q(\vec{x}, t)\}, \forall \vec{x}$, that is the sum of all objects where $\vec{x}$ is the address of each high level object representation and $t$ is time, is written as a sum of two parts:

$$C(t) \equiv \langle C(t) \rangle + \delta C(t),$$ (16.1)

where $\langle C \rangle$ refers to a slowly-varying, local average value of $C$, and $\delta C$ refers to a rapid fluctuating, stochastic remainder. This decomposition isolates those parts of the environment that lead to a stable (smooth) average configuration and which parts tend to be rough and unpredictable. In systems that are manageable of interest, one expects $|\delta C| \ll |\langle C \rangle|$, else we are doomed to unpredictability.

Note also that, by definition, $\langle \delta C \rangle = 0$, thus the fluctuations are evenly (though not necessarily symmetrically) distributed about the local mean value. This means that, if fluctuations tend in one particular direction, they will drag the mean value with them, preserving their zero mean. If one wishes to avoid a change in the mean value, then one must either offer dynamical resistance to this kind of monotonic drift, or respond to it with a counter-change, which balances it on average. This concept of preserving the mean behaviour provides us with a notion of maintenance.

Figure 16.1: An schematic picture of the separation of scales in an open dynamical system, which satisfies eqn. 16.1. The jagged line represents the highest resolution view of what $q(t)$ is doing. The solid curve is a short-interval local average value of this behaviour, and the solid blocks (dotted line) are a much coarser local average. The order of magnitude of the system’s approximate oscillations is $\omega^{-1}$. 

16.2 Average Changes in Configuration
CHAPTER 16. POLICY AND MAINTENANCE

TASKS

The concept of a task is needed to discuss a part of a system which operates autonomously for some purpose, such as maintenance, ([Bur03]).

**Definition 65 (Task).** Let a task \( \tau(t) \) be a system contained within a subset \( s \) of the total system \( S \):

\[
\tau(\vec{x}, t) = q(\vec{x}, t) : x \in s,
\]

where the restricted coordinates \( x \) ranges only over the subsystem.

**Example 177.** A task is an autonomous sub-part of a system, like a computer program or sequence of external changes made by a user. If a task is closed, it does not affect anything outside of its own resources; if it is open it can affect the state of the rest of the system also. In a distributed environment a program on one host can affect the state of a program on another host. The actions of a human interacting with the system can also lead to a task.

We now have a representation of programs running on the system as well as processes carried out by external agents (other computers and humans). One can now define maintenance in terms of the effect of sub-systems on the total system.

**Definition 66 (Maintenance Task).** Let \( \tau_M(\vec{x}, \vec{t}) \) be a task in a system \( S \) with configuration spanning \( s \), and \( \tau_{M_c}(\vec{y}, \vec{t}) \) be the complement to the subset, i.e. the remainder of the configuration of \( S \) spanning \( s_c \); then \( \tau_M(\vec{x}, \vec{t}) \) is said to be a maintenance task if \( \{ \tau_M(x \in s, \vec{t}) \} \) is an open system and

\[
\frac{d}{d\vec{t}} \left( \sum_{x \in s} \log \tau_M(\vec{x}, \vec{t}) + \sum_{y \in s_c} \log \tau_{M_c}(y, \vec{t}) \right) < \frac{d}{d\vec{t}} \left( \sum_{y \in s_c} \log \tau_{M_c}(y, \vec{t}) \right). \tag{16.3}
\]

In other words, the presence of a maintenance task \( \tau_M \) reduces the total rate of change of the average configuration state \( q(\vec{x}, t) \) in \( S \); i.e. it counterbalances the information in the fluctuations \( \delta q \) within any smoothed time interval \( \Delta t \). If the rate of maintenance is less than the rate of fluctuation, it will lead to a window of uncertainty in the value of \( \langle q \rangle \), which can result in a real change of average state. The logarithms in these formulae make the ordering and overall scale of the changes unimportant. This is a characterization of the change of information in the configuration, where the spatial ordering is unimportant.

**Example 178.** In terms of discrete information coding, a fluctuation composed of operators

\[
\hat{O}_1 \hat{O}_2 \hat{O}_6 \hat{O}_1 \hat{O}_3
\]
Stochastic open system

<table>
<thead>
<tr>
<th>Fluctuations, system operations $\delta q$ environmental changes</th>
<th>$T_c \sim T_e &lt; T$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cycles of persistent behaviour</td>
<td>$T \equiv 2\pi \omega^{-1}$</td>
</tr>
<tr>
<td>A coarse grain of $N$ cycles</td>
<td>$\Delta t = NT \gg T$, i.e. $(N \gg 1)$</td>
</tr>
<tr>
<td>User/policy time scale</td>
<td>$T_p \gg T$</td>
</tr>
<tr>
<td>Long term behavioural trends</td>
<td>$T_b \gg T_p$</td>
</tr>
</tbody>
</table>

Table 16.1: The separable time-scales for changes in a computer system interacting with an environment.

As the string, with entropy $H$ grows longer, the likelihood $2^{-H}$ of being able to find the precise counter-string becomes exponentially smaller, if the exact sequence is required. If the operations commute, then there is an average chance $1/NH(c)$ of being able to counter the string, since order no longer matters.

The definition of maintenance allows for gradual evolution of the idealized persistent state (e.g. a slow variation in the average length of a queue), since the average value can be slowly modified by persistent fluctuations. This change of the persistent state is said to be adiabatic in statistical mechanics, meaning slow compared to the fluctuations themselves. A summary of time scales is shown in table 16.1.

In order to describe and implement a system policy, for managing the behaviour of a computer system, it must be possible to relate the notion of policy to rules and constraints for time-evolution which are programmed into $q(\vec{x}, t)$. Such rules and constraints are coded as software in $q(\vec{x}, t)$, or are issued verbally to users in the environment of the system. The behaviour of the configuration state is not completely deterministic and is therefore unpredictable. By separating slowly and rapidly varying parts, using a local averaging procedure, we find an average part that is approximately predictable.

We note, as a commentary, that while this shows that the rate of change in the system can be arranged to maintain a particular state over a consistent set of time-scales, it does not specify a unique route to such a state through the state space (including space and time scheduling) of the Human-Computer system (see [Tra02, CK96]). The existence inequivalent different routes must be handled by a framework in which they can be compared in some system of returned value. The theory of games, as presented in the final sections of the paper, is suitable for selecting such a route. The existence of a unique path has been addressed in [Bur04].
16.3 The Reason for Random Fluctuations

In the study of dynamical systems, the environment is not normally modelled as a detailed entity owing to its complexity; rather one considers the projected image of the environment in the main system of interest. The essence of the definition is that the environment leads to a projected component in $S$ which appears to be partially random (stochastic), because the information about cause and effect is not available. This causes $S$ to behave as an open dynamical system.

**Definition 67.** An open dynamical system is the projection of an ensemble of interacting systems $E = \{S_1, S_2, \ldots, S_N\}$, onto $S_1$. The time development, $\hat{D}$, of the open system, may be considered as operating over a noisy channel, since information from the rest of the ensemble affects the total configuration of the host connected to the ensemble $C_1(q_1(\vec{x}, t))$. The closed rule for development of all system is intertwined:

\[
C_1(t + dt) = (1, 0, \ldots, 0) \begin{pmatrix} \hat{D}_{11} & \hat{D}_{12} & \cdots & \hat{D}_{1N} \\ \hat{D}_{21} & \hat{D}_{22} & \cdots & \hat{D}_{2N} \\ \vdots & \vdots & \ddots & \vdots \\ \hat{D}_{N1} & \hat{D}_{N2} & \cdots & \hat{D}_{NN} \end{pmatrix} \begin{pmatrix} C_1(t) \\ C_2(t) \\ \vdots \\ C_N(t) \end{pmatrix}
\]

This definition is an admission of unpredictability in a system that is open to outside influence. Indeed, this unpredictability can be stated more precisely:

**Lemma 1.** The configuration state of an open system $S$ is unpredictable over any interval $dt \sim T_e$.

(See table 16.1)

**Proof.** This follows trivially from eqn. (16.4). There is no equation for the evolution of any part of the system in isolation from the others:

\[
C_1(t + dt) \neq \hat{D}C_1(t),
\]

for any $\hat{D}$, since $C_1(t + dt)$ is determined by information unavailable within all of $S$, iff $D_{ij} \neq 0$ for $i \neq j$, which defines the open system.

The only way that a system can become exactly predictable is by isolating itself and becoming closed.

**Example 179.** The idea of closed systems can be turned around and made into a requirement. See for instance the approach advocated in [CHIK03]. By forcing the closure of a sub-system and placing restrictions or constraints on channels of communication, one maximizes consistency and predictability.
16.4 HUGE FLUCTUATIONS

In section 10.10, it is remarked that there exist stable stochastic distributions that have large fluctuations with formally infinite variance. These have been observed in network traffic, for instance (see [LTWW94, WP98]). How do these distributions fit into the picture of maintenance described here? In short, these statistical states cannot be maintained without infinite resources. This is not an acceptable maintenance cost.

Statistical stability is thus not enough to ensure maintainability. Fluctuations must be finite in order for us to have a chance. There are two ways to regard this: we can say that the fluctuations are beyond our ability to maintain and resign ourselves to that fact that systems with such behaviour are not maintainable, or we must reinterpret policy to incorporate this environment: stable distributions supercede maintenance—they are already stable. Policy should be based on the appropriate definition of stability for the system. If that includes allowing for power law fluctuations, then that is the best we can do.

16.5 EQUIVALENT CONFIGURATIONS AND POLICY

A high level partitioning of the configuration, which evolves according to rules for time-development at the same level, leads to the appearance of symmetries, with respect to the dynamical evolution of a computer system. A symmetry may be identified, whenever a change in a configuration does not affect the further evolution of the system except for the order of its elements. The configurations of the system which are symmetrical, in this sense, form a group.

\textbf{Definition 68.} A group $\mathcal{G}$ of transformations is a symmetry of the high level configuration $q(\vec{x}, t)$, if for some $x$ and time $t$, the transformation of the configuration domain

\begin{equation}
q(\vec{x}, t) = q(g(x), t),
\end{equation}

is an identity, and $g \in \mathcal{G}$.

Thus a relabelling of process addresses is unimportant to the configuration, as is any change in $\hat{D}_t$ which leads to a relabelling in the future. These are just arbitrary labels, unimportant to the functioning of the system. Since the deterministic part of the mapping $\hat{D}_t$ is coded in $q(\vec{x}, t)$, this includes changes in the way the system evolves with time.
Definition 69. A group $\Gamma$ of transformations

\[ \gamma : Q_\ell \to Q_\ell. \]  \hspace{1cm} (16.7)

is a symmetry of the state space $Q_\ell$, if

\[ q(\vec{x}, t) = \gamma(q(\vec{x}, t)), \]  \hspace{1cm} (16.8)

is an identity, and $\gamma \in \Gamma$.

Thus, if two states are equivalent by association, the system is unchanged if we substitute one for
the other.

Symmetries are hard to describe formally (they include issues such as the presence of comments
in computer code, irrelevant orderings of objects, and so on), but they have a well-defined meaning
in real systems.

Example 180. Renaming every reference to a given file would have no effect on the behaviour of
the system. Another example would be to intersperse instructions with comments, which have no
systemic function. Another an important symmetry of systems is independence of the system to
changes in parts of the configuration space $R^\ell$ which are unused by any of the programs running
on the system.

The presence of symmetries is of mainly formal interest to a mathematical description of
systems, but their inclusion is necessary for completeness. In particular, the notion of equivalence
motivates the definition of a factor set of inequivalent configurations

\[ P(t) \equiv \frac{C(t)}{G \otimes \Gamma}. \]  \hspace{1cm} (16.9)

which allows us to use one representative configuration from the set of all equivalent configurations.
In just a moment we shall claim that this quantity is intimately associated with the idea of policy.
This factored system is now uniquely prescribed by an initial configuration, rules for time develop-
ment and the environment. It is scarcely practical to construct this factor set, but its existence is
clear in a pedantic sense.

16.6 IDENTIFICATION OF A POLICY WITH A STATE OF PROMISES KEPT

Up to stochastic noise, the development of the open system is completely described by a configura-
tion of the form of eqn. (16.9), which includes the programs and data which drive it. Conversely,
the behaviour at level $\ell$ is completely determined by the specification of a $P(t)$. With a bottom-up
argument about dynamically stable configurations, we have therefore found a set of objects, one for each inequivalent configuration chain, that can be deemed stable and has the potential to be unique in some sense, yet to be clarified. This is therefore a natural object to identify with *system policy* ([Bur03]).

In practice, only a part of the configuration will directly impact on the evolution of the system at any time. If a constant part of $P(t)$ can be identified, or if $P(t)$ is sufficiently slowly varying, then this quantity plays the role of a *stable policy* for the system. If no such stability arises, then the policy and configuration must be deemed unstable.

How does this definition of policy fit in with conventional, heuristic notions of policy? A heuristic definition is i) a system configuration, ii) rules for behaviour of the system (programmed), iii) rules for human users (requested), and iv) a schedule of operations. Of these, i) and ii) may be coded into the configuration space without obstacle. iii) needs to be coded into the environment, however the environment is not a reliable channel, and can only be expected to obey policy partially, thus there will be an unpredictable component. iv) is also programmed into the computer, but there is also a schedule of random events which belongs to the environment; this also leads to an unpredictability. The resulting ‘error’ or tendency towards deviation from steady behaviour must be one of two things: a slow drift $\Delta P = P(t) - P(t')$ (systematic error) or a rapid random error $\delta P(t)$ (noise). In order to use a definition of policy such as that above, one is therefore motivated to identify the systematic part of system change.

### 16.7 CONVERGENT MAINTENANCE

The notion of convergence was introduced conceptually in [Bur98a] and explicitly in [Bur98b]. Some authors later seized upon the word homeostasis to describe this, appealing to a biological analogy ([SF00]). It is related to the idea of the fixed point of a mapping (see [Mye91] for an introduction). If $q' = U(q)$ is any mapping, then a fixed point $q^*$ is defined by,

$$q^* = U(q^*). \quad (16.10)$$

This definition is too strict in a dynamical system, rather we need a limiting process that allows for some fuzziness:

$$q^* - U(q^*) < \epsilon. \quad (16.11)$$

As defined, a policy is neither a force for good nor for evil, neither for stability nor for chaos; it is simply an average specification of equivalent system behaviours. Clearly, only a certain class of policies has a practical value in real systems. This refers to policies that lead to short term stability, thus allowing a stable function or purpose to be identified with the system. A system which modifies itself more rapidly than a characteristic human time-scale $T_p$, will not have a stable utility for humans.
The notion of convergence is especially useful ([Bur98b, CG99, CD01]) for regulating systems. A system which possesses a cycle that persists over a given interval of time can be defined as having predictable behaviour over that interval.

**Definition 70 (Convergent policy).** A convergent policy $P(t)$, of order $n$, is one whose chain of time transitions ends in a fixed point configuration $q(\vec{x}, t_f)$, for all values $x$ and times $t_i > t_f, f \leq n$. i.e.

$$(\hat{D}_t)^n q(\vec{x}, t_i) = q(\vec{x}, t_f), \text{ for some } n \geq 0, t_i < t_f. \tag{16.12}$$

The fixed configuration on which the time development ends is sometimes said to be ‘absorbing’, since once the system has entered that state, it does not change again. In the language of system administration, one says that the system has converged. In a stochastic, interacting system, this finality cannot be guaranteed precisely. Within a short time period a change away from the final state can occur at random, thus it is useful to define the notion of average convergence.

**Definition 71 (Convergent average policy).** A convergent average policy $P(t)$, of order $n$, is one whose average behaviour in time ends in an average state $\langle q(\vec{x}, t_f) \rangle$ between any two times $t_i$ and $t_f$, such that $t_f - t_i > \Delta t$.

$$\left\langle (\hat{D}_t)^n q(\vec{x}, t_i) \right\rangle = \langle q(\vec{x}, t_f) \rangle, \text{ for some } n \geq 0, t_i < t_f, \tag{16.13}$$

where $\langle \ldots \rangle$ is any local averaging procedure.

This condition is weaker, because it allows the final state of exhibit fluctuations that are balanced within the time of the averaging interval.

A discrete chain interpretation of periodicity may be found in [GS01]; it is convenient here to use the continuum approximation. Over the time interval, it can thus have the general form:

$$\langle q(\vec{x}, t) \rangle = \left\langle Q_0(x) + A(t) \text{ Re } \exp \left( \frac{i\omega t}{n} \right) \right\rangle = Q_0(x), \tag{16.14}$$

i.e. it has an average value and oscillations whose average effect is zero. Since $Q$ is positive, $A < Q_0/2$. Notice that a process that has converged becomes memory-less, i.e. its dependence on previous states becomes irrelevant.

A policy in which the average resource usage is constant over the policy timescale $T_p$ is a convergent average policy; e.g. a policy of deleting all old temporary files, killing old processes and so on, or by adding new resources, so that that fraction of used resources is constant on a average of a few cycles.

Another example of convergence would be one in which errors in a configuration file, made by human error, were corrected by an automatic process, within a short time interval, by regular
checkups, thus preserving the average condition. This has already become a common practice by
many system administrators, so convergence is a commonly used strategy for achieving stability.

**Persistence**

Implicit in the foregoing discussion of averages are two notions of stability which now crave
definition, at the level of the continuum description. These form the basis for a self-consistent
definition of convergent system policy, which show that system administration is a soluble problem,
within clear limits.

The coarse graining procedure is a redigitization of the time-line. Local averaging procedures
are used to separate structures in the time evolution of systems at different levels. One begins by
digitizing a details function of time into coarser blocks (like a pixelized image). As one zooms out,
the behaviour of a local average looks smooth and continuous again.

**Definition 72 (Persistent state).** A persistent state $\Psi(\vec{x}, t) = q(\vec{x}, t)$ is a configuration for which
the probability of returning to a configuration $\Psi(\vec{x}, t_0)$ at a later time $\Psi(\vec{x}, t_0 + \Delta t)$, for $\Delta t > 0$
is 1. In the continuum description, persistence is reflected in the property that the rate of change of
the average state $\langle \Psi \rangle$ be much slower than the rate $\omega$ of $\delta \Psi$:

$$\left| \frac{1}{\langle \Psi \rangle} \frac{d\langle \Psi \rangle}{dt} \right| = \left| \frac{d}{dt} \log \langle \Psi \rangle \right| \ll \omega \quad (16.15)$$

i.e. the fast variation extends over several complete cycles, of frequency $\omega$ (see table 16.1), before
any appreciable variation in the average is seen.

**Example 181.** A system job queue has a fluctuating queue size, whose average length can be
determined as a matter of policy, based on observed behaviour, by choice of a scheduling. Since the
arrival of jobs in the queue cannot be accurately predicted, the average length will vary slowly, as
long as jobs are expedited at only approximately the same rate as they arrive. There is thus a short
term cycle; add job, expedite job, that increases then decreases the queue size. A persistent state is
much larger than this cycle. It means that the cycle is locally stable. If the system is characterized
by a convergent policy (incoming jobs are indeed expedited at an appropriate rate), then any
fluctuations occurring at the rate $\omega$ will be counteracted at the same rate, leading to a persistent
(slowly varying average) state. See fig. 16.2.

Thus the meaning of a convergent policy is its resulting persistence. Thus, policy itself must be
identified with that average behaviour; this is the only self-consistent, sustainable definition, as
long as there are stochastic variables in the system, due to environmental interaction.

The development of an open system is stochastic and this indicates the need for a local
averaging procedure to describe it. The split one makes in eqn. (16.1), therefore ensures that
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Figure 16.2: A persistent state is one in which the cycle does not vary appreciably over many cycles. Here one sees small variations repeated many times, on a slowly varying background.

the fluctuations are zero on average, distributed about the average behaviour, so by blurring out these fluctuations, one is left with a unique description of the average behaviour. The normalized, coarse-grained policy may now be written:

\[
\langle P(\vec{x}, t) \rangle = \frac{\int_{t-\tilde{t}/2}^{t+\tilde{t}/2} dt P(t) \rho_E(t)}{\int_{t-\tilde{t}/2}^{t+\tilde{t}/2} dt \rho_E(t)} = \langle C(\tilde{t})/(G \otimes \Gamma) \rangle, \quad (16.16)
\]

In other words, we have show that the short term evolution of policy can only be identified with a local average configuration in time; i.e. a set of locally average variables, at an appropriate coding level for the system.

16.8 THE MAINTENANCE THEOREM

With the meaning of the local averaged mean-field established, it is now a straightforward step to show that local averaging leads to persistence, and hence that this measure of stability applies only to locally averaged states. We thus approach the end of the lengthy argument of this section, which shows that policy can only be an agent for average system state. The theorem suggests that a strategy for maintaining stability in computer systems is to strive for convergence.

Theorem 6. In any open system \( S \), a policy \( P(\tilde{t}) \) specifies a class of persistent, locally average states \( \langle q(\tilde{t}) \rangle \) equivalent under symmetry groups \( G \) and \( \Gamma \), if and only if \( P(\tilde{t}) \) exhibits average convergence.

The proof of this is found in [Bur03], and follows basically from the fluctuation rates. From lemma 1, in an open system \( S \), a configuration is unpredictable over a time scale \( T_e \sim \omega^{-1} \) (see table 16.1), hence a configuration can only be guaranteed persistent on average. We thus need only
to show that a convergent average policy $\langle P(t) \rangle$, of order $n$, is persistent for a time $\Delta t \gg T$, since, by definition, this implies a set of equivalent persistent average configurations, under the available symmetries. From the definition of the maintainable fluctuations, one has:

$$\left| d\frac{\langle P \rangle}{dt} \right| \ll \omega \left| \langle P \rangle(t) \right|,$$

hence

$$\left| \frac{1}{\langle P \rangle} \frac{d\langle P \rangle}{dt} \right| \ll \omega,$$

and $\langle P \rangle$ is persistent. $P(t)$ is associated with a class of states, equivalent under a symmetry group $G$, which can vary no faster than policy, since it is a part of the policy, hence a locally average state, resulting from a non-divergent policy specification is persistent. This completes the proof.

The maintenance theorem provides a self-consistent definition of what a stable state is, and hence what a stable policy is, for a computer interacting with external agents (users, clients etc). The implication is thus that system administration can be pursued as a regulation technique (see [HL93, SS97, GK96, HZS99, DHP02]), for maintaining the integrity of policy, provided one can find a convergent average policy. It sets limits on what can be expected from a policy in a dynamical environment. Finally, the argument makes no reference to the semantic content of policy; it is based purely on information and timing.

It is interesting to note another theorem which is better known but also applicable (and very similar) to the stochastic and semantic views of policy as a propagating influence: it is simply a transcription of Shannon’s channel capacity theorem for a noisy channel ([SW49]).

**Theorem 7.** There exists a policy $P(t)$ which can evolve in time with arbitrarily few errors, i.e. the system can be forced to obey policy to within arbitrary accuracy.

Shannon’s original theorem stated that “there exists a message coding which can be transmitted with arbitrarily few errors”; i.e. by creating a policy which is so strictly enforced as to police the activities of users in every detail, one could prevent users from doing anything which might influence the strict, predictable development of the system. Such a policy is possible if the average configuration of the host that it represents has sufficiently low entropy that it can be compressed into a part of the system dedicated to maintenance (error correction). It exists because of the finiteness of the digital representation.

### 16.9 Theory of Backup and Error Correction

As an application of many ideas thus far in this book, we consider now a basic example of system maintenance: a theory of system backup. This is both an issue of great practical importance and it
is an interesting analytical problem. Unfortunately, this single topic could easily fill a half a book of this size, so we can only outline the analysis and refer to details in [BR06].

The theory of backup is a theory of random change in systems and the effort to catch up by making a response to each change. It is the study of trying to hit a moving target. This situation is not unique to backup, of course; here are some analogous examples:

- Random change of files leads to the need for renewed backup.
- Configuration error leads to the need for renewed maintenance.
- Accumulation of garbage leads to the need for garbage collection (tidying).
- Arrival of tasks in a queue leads to need for server or human action.

There are several parts to this model, all of which address the competing random processes in the problem. We must address:

- Change detection (digital or continuous; i.e. symbolic or probabilistic).
- Rates of change or event arrivals (clustered or independent arrivals).
- Rate of measurement (including scheduling intervals or detection ‘dead times’).
- Rate of transmission of response (capacity of communication channel).

The arrival of events can be modelled in a number of ways. The events might be faults, intrusions, accidents, arrival of users, cleanup after departure of users etc.

In traditional statistical theory, arrival events are always assumed to follow the pattern of a Poisson arrival process, i.e. as a stream of random, independent arrivals. This assumption is made because it is simple and has special analytical properties. The study of this kind of system models well the detection of particles from a radioactive source, e.g. by a Geiger counter. These are truly independent events. Such processes are called renewal processes (see [GS01]). However, it is known from observation that many arrival processes are not Poisson processes: arrivals of events are often clustered or come in ‘bursts’. The only way to determine this is to observe actual systems and collect data.

The most basic observation we can make about user behaviour is that it follows a basic daily rhythm. Taking data from the measurements at the author’s site, shown in fig. 2.1, we see a distinct maximum in user processes around midday and a lull at around 5:00 or 6:00 in the morning. Clearly this user behaviour must be correlated with changes in disk files, since there is a direct causation. It is a necessary but not sufficient condition for change of disk data. We thus expect that most changes will occur during the day time, and few changes during the night. This is all very well, but what about the fluctuation distribution of the arrival process? There are various ways of characterizing this: by deviation from mean or by inter-arrival time.
ARRIVAL FLUCTUATION PROCESS

Suppose we consider inter-arrival time distributions. What is the likelihood of being able to predict the next file system change? By taking direct measurements of user disks at the author’s College, one finds a highly clustered and noisy distribution of data, with a long tail. There is clearly no theoretical maximum inter-arrival time for file changes, but there is a practical maximum. During holiday periods, inter-arrival times are longest because no one is using the system.

Most changes occur closely together, within seconds, between related files (we are unable to measure multiple changes to the same file simply by scanning the current state as a backup system does). This short time part of the spectrum dominates by several orders of magnitude over the longer times. However, there is a non-zero probability of measuring inter-arrival times that are minutes and hours and even days. Clearly the breadth of scales is large, and this tells us that a simple independent Poisson type of distribution, such as we used in queueing analysis, is unlikely to be an adequate explanation of this total process.

In the measurements made while writing this book, a rather dirty power law type distribution or low order was found to approximate the arrival process:

$$p(\Delta t) \sim \Delta t^{-\alpha},$$

(16.19)

where $1 < \alpha < 2^1$. We can note that many phenomena that are driven by social networks are described by power laws (see, for instance, [Bar02] for a discussion of this). The fit is not very precise, but it is adequate to sketch a functional form$^2$.

What about the magnitude of each fluctuation? This is harder to gauge, since it would require a memory of what has gone on during the entire history of the system, and since most backup systems have to transfer an entire file at a time, this is not immediately interesting (though note that the rsync program can transfer differential content, see [TM96]).

How should we deal with these changes? There are several questions:

1. When should a detector start scanning the filesystem for change, over the daily cycle?

2. What is the risk associated with waiting once a change is made, or having a dead time between backup processes?

3. What is the cost associated with the backup process?

---

$^1$One could try to model this process by splitting up the process into a superposition of well-defined processes, and thus take account of causation.

$^2$With a noisy distribution, the value of $\alpha$ is uncertain, so we choose it to be a normalizable distribution $\alpha > 1$ to define a probability.
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DETECTION PROCESS

Clearly, in an ideal world one could make an immediate backup of every file as soon as it were changed.

Example 182. This method is used in copy-on-write mirroring technologies, such as the Distributed Computing Environment filesystem DFS. But this is expensive, both in terms of machine resources and in terms of storage. Moreover, this choice involves a policy decision. By simply mirroring the file, we choose not to keep multiple archive versions. Thus if a mistake is added to a file, it will immediately be propagated to the backup also. This causes us to lose one reason for making a backup.

Thus the decision is no longer just one about efficiency, but policy. Let us ignore the copy-on-write policy here and focus on an intermittent backup.

In order to detect changes without notification by the filesystem directly, we need a detection process. This is a process that must scan through the system, either using its hierarchical structure, or using its disk block structure to detect change. There are two strategies one might use here:

- Backups parse file tree as quickly as possible (spans shortest possible time)
- Backups parse file tree slowly (spans a large interval of time, several runs do not overlap).

In the first case, the backup process presents the shortest interruption to system resource availability but with high load (backup can be a disk and CPU intensive process that disrupts system performance for users). In the latter case, one presents a low load to the system over a longer interval of time. The extreme case here would be to have a continuous scan of the file tree, picking up modified files continuously and backup them up as a low priority process.

In the first case, one takes a rapid sharply focused snapshot of the filesystem. In the latter case, we take a blurred snapshot capturing more changes over a longer time. It is not obvious which is these strategies can capture most changes over a shortest ‘risk interval’. We would like to minimize this risk interval, i.e. the time between a file change and its backup.

In either case, we have a process of file change arrivals overlapping with a detection process. If both processes are random in time, these are two superposed random processes. As we know from section 10.9, the superposition of random processes does not lead to the same kind of random process except in a few special cases. If the detection process is regular in time, we might suppose that detection is not a random process, but this is not the case. In detection one must parse the file structures in some regular manner in order to cover the file space evenly; one cannot predict when the actual change will be registered during this parsing. With partially ordered events, for instance, a dependency structure that can be represented graphically, results of experiments indicate that a random sampling can be an efficient (see [San01]) way of parsing the task list. However, random access makes it difficult to cover the file-tree in an even handed manner, and could demand large
disk head movements. We must always remember the physical limitations of disk backup: it is
the mechanical movement of the disk head that retards the system performance the most, and the
thus increases the performance cost. Since interleaving a continuous scan of the disk with random
access must lead to large movements at least some of the time. This suggests that copy on write
would be a most efficient way of making an immediate copy, since it requires no additional disk
head movement.

In the maintenance theorem, we avoided mentioning the form of fluctuation distributions
explicitly, dealing only with rates and finiteness since computer systems are finite. Here, observation
directs us towards an approximate power law distribution.

**POLICY CHOICES**

The mechanisms behind the optimization of backup are rather complex and a full understanding of
them goes beyond the scope of this book; however, suppose we have satisfied ourselves as to the
causes and have a satisfactory model of file arrivals that describes a file time distribution of the
form

\[ P(\Delta t) = A \Delta t^{-\frac{3}{2}} e^{-\gamma \Delta t}, \]  

where \( A \) is a constant amplitude and \( \gamma \) is the decay time of the distribution that truncates the infinite
power-law tail to no longer than a day (since the process is pseudo-periodic, with daily period,
there is not point in looking to times longer than this to describe a maintenance procedure that will
also periodic). We now consider what this means for the best approach to capturing change with
minimal risk of loss.

Most changes are for small \( \Delta t < 10 \) seconds (see fig. 16.3), so a frequent search is likely to
reduce the risk time between change and backup more effectively than a regular daily backup. A
social network model should also be used to look at the locations of these files, and relate it to the
scanning policy used by the backup program to see whether changes are likely to be clustered in
the file tree as well as in time.

How is risk evaluated for a backup strategy? The pathways for failure are various:

- Accidental file deletion.
- Malicious file deletion.
- Disk failure.
- Natural disaster.

In defence against these “gremlins” we have choose:

- Copy on write.
Figure 16.3: A sketch of the distribution of inter-arrival times for file changes. A log-log plot has a gradient of about 3/2, hence the power law in eqn. (16.20)

- Backup daily (over short bursts).
- Backup daily (over long times).
- Continuous search and copy.

We can set these up in opposition to one another, by creating a matrix of cases, either knowing the relative probabilities of the loss mechanisms, or starting from the worst case scenario. In chapter 19, a method for calculating and optimizing these strategies is described. The risk or payoff associated with each pair of strategies can then be modelled, for instance, by the probability that the copy is up to date and divided or subtracted by the relative disruption cost of the backup method to system productivity. We can make various assumptions about systems by observing their nature.

Example 183. Taking a backup at night is not obviously the best strategy. It reduces the overhead, but if one accepts that most accidents happen while the users are busy, then this also maximizes the chance of changes being lost due to some catastrophe.

Example 184. Using a program that backs up data very quickly provides only a momentary snapshot of one time of day. From the data here, we should can minimize risk by choosing the backup time to be somewhere close to the maximum of the graph of $R(t)$.

This vagueness underlines the point that, in science, there is virtually never only one right answer to a question. In system administration, it is about making an arbitrary compromise.

Let us examine slightly more closely how we might evaluate these strategies in terms of data capture rates. If a backup is to success, then its average performance must be such that its
average capture rate is equal to the average rate at which changes are made to the file system. The cumulative change to the file system at rate $R(t)$, over an interval $\Delta t$ is (see fig. 16.4):

$$C(\Delta t) = \int_0^{\Delta t} R(t) \, dt. \quad (16.21)$$

Note the regions of differing rates during on and off-peak times, $m_1$ and $m_2$, where $m_1 > m_2$. In region $m_1$ there is high risk of losing changes: there is $m_1/m_2$ times the risk in region $m_2$. The likelihood of disruption by taking backup, on the other hand is greater in region $m_1$, so we might try to even out this disruption by capturing at a rate not exceeding $m_2/m_1$ of the total rate of change. The simplest solution, when these are in balance, is thus when $m_1 = m_2$ and we have the constant rate of capture with dotted line, in fig. 16.4.

Now, if we want to perform a quick non-disruptive backup, then this diagram tells us to perform the backup at the positions marked by the dark blobs, since these lie at the end of the period of maximum change. This assumes that the rate of failure leading to need for backup is much slower than the rate at which data change on the disk, and can therefore happen with more or less equal likelihood at any moment. By placing the backup after the maximum rate period, we assume capture of the largest number of changes since the last backup, while at the same time minimizing the disruption.
It is almost obvious from the diagram that, in this simplified picture, there is no advantage to choosing any special time for a short-burst daily backup except to minimize disruption. We do not capture any greater number of changes in this way, on average. However, if we consider making more than one backup per day, or using a continuous scan technique, there are advantages to picking the time more carefully and a more detailed analysis is required (see [BR06]).

If we change any of these simplifying assumptions to make the model more realistic, some of the details will change, but the essence will most likely remain somewhere between continuous transfer and short burst rates.

**Applications and Further Study 16.**

- Determining whether a system design is maintainable (sustainable) or not.
- Basis for investigating generalized renewable processes.
- Foundation for the very existence of systems.
CHAPTER 17

KNOWLEDGE, LEARNING AND TRAINING

The great tragedy of science –
the slaying of a beautiful hypothesis by an ugly fact.

– Thomas Huxley

Human-computer systems are often called knowledge-based systems, because they rely on a ‘database’ of procedural or factual expertise to carry out their function. Knowledge has to be acquired somehow, and then summarized and stored. There are two forms of stored knowledge:

- Statistically acquired data that document experience.
- Rules or relationships matched to observation, then summarized in algebraic form.

In the first case, we observe and check and let experience be our guide; in the second case we believe that we have understood and so summarize what the data seem to show as a formula, in the context of a theory. Similarly, knowledge is acquired in one of two ways:

- Supervised learning or training: we are certain of the value of information (e.g. is a fact true or false) and all that remains is to incorporate this classification into decision making procedures of the system.
- Unsupervised learning or just learning: we are not certain of the precise classification of information, and have to rely on the statistical significance of information to formulate a hypothesis.
The scientific process itself is about continually questioning and verifying our beliefs about such knowledge until we are satisfied with their accuracy. It is unquestionably a process of unsupervised learning: no-one knows the right answers; we have to make do with dribs and drabs of evidence to make sense of the world.

Experts tend to overestimate their competence in making judgements, so it is important to have impartial methods that can mitigate this overconfidence with a reality check. Probability theory is a key tool in estimating the uncertainties in the learning process, but even absolute probabilistic descriptions cannot always be made with sufficient accuracy to make knowledge reliable. We therefore need to consider how sufficient knowledge is built up and refined in a quantitative fashion 1. Note that, while ordinary statistics have no ‘direction’ (the correlation of two variables implies no ordering or asymmetry between them), the Bayesian viewpoint is directional and can therefore be used to discuss cause and effect, or the arrow of development.

17.1 INFORMATION AND KNOWLEDGE

A system or organization’s knowledge is built up gradually from experience, by revising facts and procedures until the process converges on some corpus of expertise that enables it to carry out its function. Uncertainty remains throughout this process, but can be quantified and allowances can be made for the imperfection. This chapter brings us back to the importance of empirical observation.

Rather than giving up on a model, if we do not have sufficient data to reasonably estimate the parameters, we can use hints and guesses to ‘bootstrap’ the investigative procedure, and then revise estimates based on testing out the early assumptions. Bayesian statistics are widely used as a tool for modelling learning (machine learning, human learning, behavioural adaptation etc.). The idea is to gradually refine knowledge and move from a situation with imperfect information to one with perfect information. When we learn, by gathering new evidence of phenomena, it is equally important to forget outdated knowledge, so that contradictions do not arise.

The idea of expertise and knowledge really brings us back to the philosophical issues about science and its interpretation of the world (see section 2.3). Without getting embroiled in this tarpit, we can refer to the problem of expertise as being a human one, that relates to policy. The roles of humans in making such interpretations are central.

1 Many is written on the subject of learning and statistical inference that goes way beyond the scope of this book. Readers are referred to books on causal statistics, e.g. [Bre70, Pea88] or on pattern recognition, e.g. [DHS01] for more information on this vast and subtle topic. Bayesian methods are frequently cited, and often used to throw a veil of philosophical subtlety over the subject. Here be dragons!
17.2 **Knowledge as Classification**

In chapter 9, we defined information by the number of symbols that must be transmitted in order to exactly reproduce an object or procedure independently of the original. The uncertainty about the original was measured by the informational entropy. Both of these definitions hinged on the need to classify observations into *symbol classes*. Even time series seek to identify from a list of basic features. The more focused a classification, the lower the uncertainty (entropy). We say that the acquisition of information about something decreases out uncertainty about it, so we need to receive classified symbols to achieve this. This tells us that classification is central to identifying data with knowledge.

**Definition 73 (Knowledge).** Knowledge is a systematic classification of facts and algorithms, i.e. it is about identifying events with a correct hypothesis (class) for their cause.

We shall now return to the idea of classification to define what we mean by learning. Learning is closely related to the acquisition of information, but it is not identical because information might tell us all kinds of contradictory things about a system. It is our ability to classify those pieces of information into a consistent picture of *probable cause* that is knowledge.

Pattern recognition is central to ability to classify (see [DHS01]). As with the classifications of knowledge above, this falls largely into two types: deterministic matching and probabilistic matching of patterns. Biological pattern recognition, as exercised by the immune system, it believed to be a statistical hybrid of a symbolic (deterministic) model of string matching ([PW97]). This is an approach that has been used to look at integrity checking and anomaly detection (see [PFH96]).

What is impressive about the biological immune system is that it recognizes patterns (antigens) which the body has never even seen before. It does not have to know about a threat in order to manufacture antibody to counter it. Recognition works by jigsaw pattern-identification of cell surface molecules out of a generic library of possibilities. A similar mechanism in a computer would have to recognize the ‘shapes’ of unhealthy code or behaviour [Mic, ea]. If we think of each situation as begin designated by strings of bytes, then it might be necessary to identify patterns over many hundreds of bytes in order to achieve identify a threat. A scaled approach is more useful. Code can be analyzed on the small scale of a few bytes in order to find sequences of machine instructions (analogous to dangerous DNA) which are recognizable programming blunders or methods of attack. One could also analyze on the larger scale of linker connectivity or procedural entities in order to find out the topology of a program.

**Example 185.** To see why a single scale of patterns is not practical we can gauge an order of magnitude estimate as follows [PW97]. Suppose the sum of all dangerous patterns of code is $S$ bytes and that all the patterns have the same average size. Next suppose that a single defensive spot-check has the ability to recognize a subset of the patterns in some fuzzy region $\Delta S$. i.e. a given
agent recognizes more than one pattern, but some more strongly than others and each with a certain probability. Assume the agents are made to recognize random shapes (epitopes) that are dangerous, then a large number of such recognition agents will completely cover the possible patterns. The worst case is that in which the patterns are randomly occurring (a Poisson distribution). This is the case in biology since molecular complexes cannot process complex algorithms, they can only identify affinities. With this scenario, a single receptor or identifier would have a probability of $\frac{\Delta S}{S}$ of making an identification, and there would be a probability $1 - \frac{\Delta S}{S}$ of not making an identification, so that a dangerous item could slip through the defenses. If we have a large number $n$ of such pattern-detectors then the probability that we fail to make an identification can be simply written,

$$P_n = \left(1 - \frac{\Delta S}{S}\right)^n \sim e^{-n \frac{\Delta S}{S}}.$$  \hspace{1cm} (17.1)

Suppose we would like 50% of threats to be identified with $n$ pattern fragments, then we require

$$-n \frac{\Delta S}{S} \sim -\ln P_n \sim 0.7.$$  \hspace{1cm} (17.2)

Suppose that the totality of patterns is of the order of thousands of average sized identifier patterns, then $\Delta S/S \sim 0.001$ and $n \sim 7000$. This means that we would need several thousand tests per suspicious object in order to obtain a fifty percent chance of identifying it as malignant. Obviously this is a very large number, and it is derived using a standard argument for biological immune systems, but the estimate is too simplistic.

Testing for patterns at random places in random ways does not seem efficient, and while it might work with huge numbers in a three dimensional environment in the body, it is not likely to be a useful idea in the one-dimensional world of computer memory (though see [San01]). Computers cannot play the numbers game with the same odds as biological systems. Even the smallest functioning immune system (in young tadpoles) consists of $10^6$ lymphocytes, which is several orders of magnitude greater than any computer system. What one lacks in numbers must therefore be made up in specificity or intelligence. The search problem is made more efficient by making identifications at many scales. Indeed, even in the body, proteins are complicated folded structures with a hierarchy of folds which exhibit a structure at several different scales. These make a lock and key fit with receptors which amount to keys with sub-keys and sub-sub-keys and so on. By breaking up a program structurally over the scale of procedure calls, loops and high level statements one stands a much greater chance of finding a pattern combination which signals danger.

17.3 BAYES THEOREM

The basic formula normally used in learning is Bayes theorem for conditional probability. This prescribes a well defined method for a learning procedure, but it is not the only method (see section
We have already seen how conditional probability allows us to attach a causal arrow to the development of system information (see section 9.7). We now take advantage of this to develop a method of increasing certainty, or refined approximation by including the effect of observed evidence.

Bayes formula is an expression of conditional probability. The probability of two separate events, $A$ and $B$, occurring together may be written

$$P(A \text{ AND } B) = P(A \cap B) = P(A|B)P(B) = P(B|A)P(A).$$

(17.3)

If the events are independent of the order in which they occur, i.e. they occur simultaneously by coincidence, then this simplifies to

$$P(A \text{ AND } B) \rightarrow P(A)P(B).$$

(17.4)

The symmetry between $A$ and $B$ in eqn. (17.3) tells us that

$$P(A|B) = \frac{P(B|A)P(A)}{P(B)}.$$  

(17.5)

This trivial re-expression is that basis for system learning. If we rewrite it for more than two kinds of event (see fig. 17.1), using fixed classes $c_i$, for $i = 1 \ldots C$, and an unknown event $E$ that could be in any of them, we have

$$P(c_i|E) = \frac{P(E|c_i)P(c_i)}{P(E)} = \frac{P(E|c_i)P(c_i)}{\sum_{i=1}^{C} P(c_i, E)} = \frac{P(E|c_i)P(c_i)}{\sum_{i=1}^{C} P(E|c_i)P(c_i)}.$$  

(17.6)

This is Bayes’ formula. The usefulness of this much adored result lies in a special interpretation of the learning process. We assume that the ‘true’ classification of observed information is represented by an event of type $c_i$, and that this is a fixed classification. Our aim in learning is to determine the most probable classification of information, given new information.

The Bayesian philosophy distinguishes between $a$ priori probabilities (i.e. our initial estimate or guess, which is based on imperfect information), and $a$ posteriori probabilities (i.e. our revised estimates of likely classification after receiving new information). The formula is then interpreted as follows.

- The uncorrelated probabilities of seeing different events $P(c_i)$ and $P(E)$ are assumed to be known. We can guess these initially, or simply admit that our previous experience is always limited and that they are estimates of the truth, and might not be very accurate. They are based on prior knowledge, i.e. the experience we have before making any new observations.
The conditional probability $P(E|c_i)$ is interpreted as the likelihood that we are able to classify an event $E$, given that we know the classes. This is a likelihood function. As this probability increases, our guesses become closer and closer to the truth. This is treated as though it were a function of $E$, and is sometimes written $L(E|c_i)$ to mark this special interpretation.

The derived (a posteriori) probability $P(c_i|E)$ is the best estimate of the probability of seeing the classification $c_i$, given the evidence $E$ is $P(c_i|E)$. It can be thought of as $P(H_i|O)$, the probability of hypothesis being the correct explanation, given the observation $O$. This result can be used directly, and it can also be used, with a little subtlety, to replace our initial estimate of $P(c_i)$ to reiterate the procedure once more.

**Example 186.** Consider a single cause-effect relationship $C = 1$, and let $c_1$ be computer failure by disk error, and let $E$ be the probability of a computer being down, by any cause. The probability of a disk error, on any computer, is found over time to be $P(c_1) = 0.02$.

The probability that a computer will be down due to disk error is initially unknown, so the system administrator sucks a finger and pokes it into the air, declaring that $P(E|c_1) = L(E|c_1) = 0.6$, i.e. there is a sixty percent chance that disk error will be the cause of computer failure. The probability that disk failure is not the cause is thus $P(E|\neg c_1) = 0.4$, since probabilities sum to
one. We can now combine these uncertainties to find the probability that disk error will be the cause, given that a computer is observed to be down:

\[
P(c_1 | E) = P(H | O) = \frac{L(E | c_1) P(c_1)}{P(c_1) L(E | c_1) + P(\neg c_1) L(E | \neg c_1)}
\]

\[
= \frac{0.6 \times 0.02}{0.6 \times 0.02 + 0.4 \times 0.98}
\]

\[
= 0.03.
\]  

(17.7)

The probability that the true cause is a disk failure is really only as low as 0.02. The uncertainty flies in the face of the system administrators finger estimate, and reflects the fact that 98% of computers do not show the symptoms of disk failure and yet 40% of computers will be down anyway, due to other causes. Note that the result is larger than the independent estimate, but only slightly: 0.03 is now our belief of the probability of disk error, rather than 0.02 that was measured. This tells us that our initial finger-in-the-air estimate was badly wrong and it has been adjusted almost all the way down to the independent measurement. We should therefore replace this new value with the old \( P(E | c_1) = L(E | c_1) = 0.03 \) and use the formula again, if we obtain new data.

### 17.4 Belief versus Truth

It is a philosophical conundrum for science that, in spite of a search for absolute truth, one is forced to settle for making a value judgement about belief. This is an inevitable consequence of unsupervised learning: the world does not give up its secrets easily, and never according to a classification that is preordained. It is therefore up to a process of inference to determine what is a reasonable belief about the truth and falsity of our hypotheses. This realisation is both liberating and complicating for the administration of human-computer systems, since so much of human involvement is based on belief.

This should not be received as a signal to abandon method however. As always, we are in the business of reducing uncertainty by proper observation and analysis. Rather, it opens up an alternative viewpoint in the form of Bayesian statistical methods. There is only space to mention these briefly here. Readers are referred to [Bre70, Pea88] and [DHS01] for more details.

**Example 187.** Network Intrusion Detection Systems (NIDS) examine arriving packets of data on the network and examine them using pattern matching rules, for possible intrusion attempts by crackers. Suppose the IDS signals an alarm. What is the likelihood that an intrusion has taken place? Clearly, we cannot be certain about this; indeed, does the question even have any meaning. We can talk about our degree of belief in the matter.

Suppose that there is a 95% chance that an attempted intrusion will trigger an alarm:

\[
P(\text{alarm} | \text{intrusion}) = 0.95.
\]  

(17.8)
Based on the possibility of false alarms, there is a 6% chance that an alarm will be false:

\[ P(\text{alarm} | \neg \text{intrusion}) = 0.06. \]  \hspace{1cm} (17.9)

Figures from security watchdog organizations indicate that there is a 0.01% chance of being attacked on a given day, so \( P(\text{intrusion}) = 0.0001 \). What is our basis for belief that the alarm is a true signal of attack? Let \( A \) stand for alarm, and \( I \) for intrusion.

\[ P(I | A) = \frac{P(A | I)P(I)}{P(A | I)P(I) + P(A | \neg I)P(\neg I)} = \frac{0.95 \times 0.0001}{0.95 \times 0.0001 + 0.06 \times (1 - 0.0001)} = 0.00158. \]  \hspace{1cm} (17.10)

Thus, this additional information about the behaviour of the alarm in response to data, has increased our belief in the validity of the alarm from 0.0001 to 0.00158, some sixteen-fold. Although the likelihood of truth is still tiny, there is a significant improvement due to the constraints on uncertainty implied by the conditional probabilities.

### 17.5 Decisions Based on Expert Knowledge

Decision making is central to system administration. A system that claims to perform an expert function cannot afford to be as badly wrong as in the finger-sucking example 186 above, so it is crucial to make reasoned estimates of the truth of our hypotheses. Bayes formula tells us how close our estimate of knowledge is to independently measured values. The Bayesian method allows iteration of the formula, and it identifies this iterative revision of probabilities with learning, or the refinement of probable hypothesis fitting (data recognition). The likelihood function \( L(H | O) \) tells us the latest state of our knowledge about the reasonableness of a hypothesis \( H \), given the observed data \( O \). The programme is:

1. Formulate \( C \) hypotheses to classify data.
2. Formulate a discriminant criterion for deciding when a hypothesis is actually true. This is needed to train the likelihood function, otherwise we can never make decisions.
3. Train the likelihood function by working out probabilities from independent measurements, as far as possible. Even if these are imperfect, they allow us to make a start.
4. Work out the revised estimates probability for probable hypothesis, or update the likelihood function with new data.
Consider the following example of using Bayes theorem to make a decision based on training, or supervised learning, in which a human is able to pass on expert judgement to a computer program in order to manage a flow of data.

**Example 188.** One of the classic examples of Bayesian hypothesis discrimination in system management is the filtering of junk mail by mail servers. There are various approaches to codifying the hypotheses about what constitutes junk mail, some are rule based and others are probabilistic.

Suppose that we initially separate junk mail by hand and collect statistics about the percentage of junk mail arriving. Using the mail history, we train the likelihood function $P(\text{data}|\text{junk}) = L(\text{junk}|\text{data})$, giving

$$L(\text{junk}|\text{data}) = 0.95$$
$$L(\neg\text{junk}|\text{data}) = 0.1$$

(17.11)

This tells us that, when we feed junk into the likelihood function, and update the probabilities based on certain knowledge (discrimination of junk from non-junk) we obtain a probability estimate of seeing the discriminating features in the data. In reality, there will be many triggers that cause us to classify mail as junk mail (spam), but they all result in a final classification of either junk or NOT junk (in set notation “$\neg$ junk”). Note that some mail is not junk, even though it passes all the tests that we make for it; thus, the separation is not a clean one.

The data we use to train our likelihood function are the test results of expert probes over thousands of E-mails. The data that we use to discriminate each incoming mail then comes from each individual E-mail, after training has ceased. We thus compare the likelihood that each individual message is spam, based on what has been learned from all the others.

On examining a new message, we test the hypothesis that it is junk. We find that about sixty percent of the junk-tests on this mail are found to be positive: $P(c_1) = P(\text{junk}) = 0.6$, $P(\neg\text{junk}) = 0.4$, where the $\neg$ symbol means NOT or complement.

$$P(\text{junk}|\text{training}) = \frac{L(\text{junk}|\text{data})P(\text{junk})}{L(\text{junk}|\text{data})P(\text{junk}) + L(\neg\text{junk}|\text{data})P(\neg\text{junk})}$$
$$= \frac{0.95 \times 0.6}{0.95 \times 0.6 + 0.4 \times 0.1}$$
$$= 0.93.$$  

(17.12)

The probability that this is actually junk is slightly less than the training estimate, i.e. we have an almost maximally high degree of likelihood that the message is spam. In this example, our belief is amplified by the high proportion of E-mail messages that are correctly identified by the data sample tests. The main effect of the Bayesian formula is to go from a simple true/false picture of whether an E-mail is junk or not, to a more refined threshold based decision that can be dealt with by making a policy decision about the correct threshold for discarding a message.
The previous example is rather simplistic. In general, effects have many causes, each of which is a potential hypothesis for failure. Bayesian networks are a way of modelling the dependencies in more involved cause trees. Here we shall offer only a simple example.

**Example 189.** Consider a situation in which a worker is trying to establish a Virtual Private Network connection with their company from home, over an Internet Service Provider (ISP) line. The probability of a fault $F$ (not being able to establish a connection) depends on two factors: either the server is busy, or the network in between is too slow. Let us denote these two causes by $B$ and $S$ respectively. From monitoring software, the probabilities of these independent events can be estimated:

\[
P(B) = P(\text{busy}) = 0.1, \quad P(\neg B) = 0.9
\]
\[
P(S) = P(\text{slow}) = 0.4, \quad P(\neg S) = 0.6.
\] (17.13)

Moreover, we might believe the following probabilities about how these probabilities link up the independent events in a dependency network.

\[
P(F|BS) = 0.8 \quad P(F|B\neg S) = 0.6
\]
\[
P(F|\neg BS) = 0.5 \quad P(F|B\neg S) = 0.5
\]
\[
P(\neg F|BS) = 0.2 \quad P(\neg F|B\neg S) = 0.4
\]
\[
P(\neg F|\neg BS) = 0.5 \quad P(\neg F|B\neg S) = 0.5
\]

Note that, when the server is busy, we don’t really have any prior knowledge of the cause of the failure: it’s 50/50.

We begin by calculating the likelihood of a fault given these rough estimates of our beliefs. This is surprisingly complicated to take account of these biases fairly.

\[
P(F) = P(FBS) + P(FB\neg S) + P(FB\neg S) + P(FB\neg S)
\]
\[
= P(F|BS)P(BS) + P(F|B\neg S)P(\neg BS)
\]
\[
+ P(F|B\neg S)P(B\neg S) + P(F|B\neg S)P(\neg B\neg S)
\] (17.14)
Now, since \( B \) and \( S \) are independent, \( P(\text{BS}) = P(B)P(S) \).

\[
P(F) = P(F|\text{BS})P(B)P(S) + P(F|\neg\text{BS})P(\neg B)P(S) \\
    + P(F|B\neg S)P(B)P(\neg S) + P(F|\neg B\neg S)P(\neg B)P(\neg S) \\
    = (0.8 \times 0.1 \times 0.4) + (0.5 \times 0.9 \times 0.4) + (0.6 \times 0.1 \times 0.6) + (0.5 \times 0.9 \times 0.6) \\
    = 0.518 \quad (17.15)
\]

Thus, the best we can conclude is that the probability that there will be a fault is a little over even odds. The reason for this is our lack of knowledge about the network hypothesis.

Suppose now, we make more observations. This would be expected to reduce the uncertainty and we have the opportunity to learn from the added evidence. We use the Bayes formula for each branch of the tree in fig 17.2.

If we learn the \( B \) caused the time out, i.e. the server is busy, then we apply the Bayes formula for propagating that fact into the probability for \( B \) being the correct hypothesis.

\[
P(B|F) = \frac{P(F|B)P(B)}{P(F)} \\
    = \frac{[P(F|\text{BS})P(S) + P(F|B\neg S)P(\neg S)] P(B)}{P(F)} \\
    = \frac{(0.8 \times 0.4 + 0.6 \times 0.6)}{0.518} \\
    = 0.131. \quad (17.16)
\]

This is the probability that it was \( B \) that caused the fault \( F \). It should be compared to \( P(B) \) — it is slightly larger, i.e. the new evidence makes the likelihood of this hypothesis more likely (though still not very likely). If we learn, on the other hand, that the slow network \( S \) is the cause of the error, we use:

\[
P(S|F) = \frac{P(F|S)P(S)}{P(F)} \\
    = \frac{[P(F|\text{BS})P(B) + P(F|B\neg S)P(\neg S)] P(S)}{P(F)} \\
    = \frac{(0.8 \times 0.1 + 0.5 \times 0.9)}{0.518} \\
    = 0.409. \quad (17.17)
\]

This can be compared to \( P(S) \). Again it is slightly larger, adding more support to that hypothesis. To take account of this learned experience, we might consider replacing \( P(B), P(S) \) with the new estimates 0.131, 0.409 and using the Bayesian rule again. This process of iteration can sometimes result in the determination of optimal estimates, in which case one speaks of Bayesian learning.
Taking the new estimate and feeding it back into the formula to obtain a new one forms the basis of what is known as Bayesian learning. It is a gradual refinement of certainty about estimated values. To make this idea precise, we must stray into more technical discussions about parameter estimation that are beyond the scope of this book, so we leave this as an open problem for the reader to investigate further. The implications for measurement are that even an imperfect experiment is better than no experiment. We can make precise use of imperfect data, in a way that can be revised later, if more observations can be made.

17.6 Knowledge Out of Date and the Importance of Forgetting

The process of learning is not only advantageous; an excess of knowledge amounts to a prejudice. Some kinds of knowledge have a sell-by date, after which the knowledge no longer applies. One example of this is in anomaly detection, where the distant past history is of little interest to the recent past. If knowledge accumulates for longer than the period over which one expects policy to be constant, then it becomes anomalous itself according to the new policy.

There are two approaches to retaining finite window knowledge: a fixed width sliding window can be used to eliminate any data points that were accumulated before a certain fixed width interval, measured backwards in time from the present; alternatively, old knowledge can gradually be degraded by assigning newer knowledge a higher weight (see [Bur02b]).

**Example 190.** Anomaly detection is usually performed using one of two techniques: off-line time-series analysis, or real-time event threshold processing. Time-series data consume a lot of space and the subsequent calculation of local averages costs a considerable amount of CPU time as the window of measurement increases. In [Bur02b] it is shown how compression of the data can be achieved, and computation time can be spared by the use of iterative updating, by geometrical series convergence. The key to such a compression is to update a sample of data iteratively rather than using an off-line analysis based on a complete record.

The approximate periodicity observed in computer resources allows one to parameterize time in topological slices of period $P$, using the relation

$$t = nP + \tau.$$  \hspace{1cm} (17.18)

This means that time becomes cylindrical, parameterized by two interleaved coordinates $(\tau, n)$, both of which are discrete in practice. This parameterization of time means that measured values are multi-valued on over the period $0 \geq \tau < P$, and thus one can average the values at each point $\tau$, leading to a mean and standard deviation of points. Both the mean and standard deviations are thus functions of $\tau$, and the latter plays the role of a scale for fluctuations at $\tau$, which can be used to grade their significance.
The cylindrical parameterization also enables one to invoke a compression algorithm on the data, so that one never needs to record more data points than exist within a single period. It thus becomes a far less resource intensive proposition to monitor system normalcy.

An iteration of the update procedure may be defined by the combination of a new data point \( q \) with the old estimate of the average \( \bar{q} \).

\[
\bar{q} \rightarrow \bar{q}' = (q|\bar{q})
\]  

(17.19)

where

\[
(q|\bar{q}) = \frac{w q_1 + w \bar{q}}{w + w}.
\]  

(17.20)

This is somewhat analogous to a Bayesian probability flow. The repeated iteration of this expression leads to a geometric progression in the parameter \( \lambda = \frac{w}{w + w} \):

\[
(q_1|(q_2|\ldots(q_r|\ldots|q_n))) = \frac{w}{w + w} q_1 + \frac{w w}{(w + w)^2} q_2 + \ldots + \frac{w w^{r-1}}{(w + w)^r} q_r + \ldots + \frac{w^n}{(w + w)^n} q_n.
\]  

(17.21)

Thus on each iteration, the importance of previous contributions is degraded by \( \lambda \). If we require a fixed window of size \( N \) iterations, then \( \lambda \) can be chosen in such a way that, after \( N \) iterations, the initial estimate \( q_N \) is so demoted as to be insignificant, at the level of accuracy required. For instance, an order of magnitude drop within \( N \) steps means that \( \lambda \sim 10^{-N} \). Using the definition of the pseudo-fixed-window average \( \langle \ldots \rangle_N \), we may now define the average standard deviation, or error, by

\[
\langle \sigma(\tau) \rangle \equiv \sqrt{\langle (\delta q(\tau))^2 \rangle_N}
\]

\[
\delta q(\tau) = q(t) - \langle q(\tau) \rangle_N
\]  

(17.22)

This has similar properties to the degrading average itself, though the square root makes the accuracy more sensitive to change.

In order to satisfy the requirements of a decaying window average, with determined sensitivity \( \alpha \sim 1/N \), we require,

1. \( \frac{w}{w + w} \sim \alpha \), or \( w \sim w/N \).

2. \( \left( \frac{w}{w + w} \right)^N \ll \frac{1}{N} \), or \( wN \ll w \).

Consider the ansatz \( w = 1 - r \), \( \bar{w} = r \), and the accuracy \( \alpha \). We wish to solve

\[
r^N = \alpha
\]  

(17.23)
for $N$. With $r = 0.6$, $\alpha = 0.01$, we have $N = 5.5$. Thus, if we consider the weekly update over 5 weeks (a month), then the importance of month old data will have fallen to one hundredth. This is a little too quick, since a month of fairly constant data is required to find a stable average. Taking $r = 0.7$, $\alpha = 0.01$, gives $N = 13$. Based on experience with offline analysis, this is a reasonable arbitrary value to choose.

17.7 Convergence of the Learning Process

The process of learning should converge with time; i.e. we would like the amount of information on which a system depends to be non-increasing. If the information required to perform a function diverges, we have no hope of managing the process. Then, we must be contented with watching the system in either despair or wonder as it continues to operate. If, however, the information base of the system is constant or even diminishing, then there is a possibility to learn sufficient information to manage system expertise.

Feeding the a posteriori estimates back into the input of Bayes formula provides a method (with some reservations) of iteratively determining the ‘true’ cause of the data, as one of the available hypotheses. It is when the prior probabilities (in their $n$th iteration) tend towards a very specific answer (low uncertainty or low entropy) that one says the true value has been learned. In section 9.11, we noted that the principle of maximum informational entropy is a way of modelling the effect of maximum uncertainty, under given conditions. The maximum entropy principle is a model for forces that tend to increase uncertainty, such as random fluctuations and errors. Bayesian learning is the logical opposite of this; thus we can think of learning and entropy as being competing forces of uncertainty and certainty in a system.

Finally, note that this brings us back, once again, to the idea of convergence and policy (see sections 5.8 and 10.4). If a well posed system, learning will tend towards a unique result and stay there in the limit of many observations. This is a stability criterion for the algorithmic flow of cause and effect in the system. Creating such systems predictably is still an open problem in many areas, but note the approaches of convergent configuration management in [Bur04, CS03].
Chapter 17. Knowledge, Learning and Training

17.8 From Pattern Recognition to Semantic Reasoning

Applications and Further Study 17.

- Discussion of knowledge and expertise in a quantitative fashion.
- A strategy for collecting data as a basis for expert knowledge.
- Automation of data acquisition (machine learning).
Chapter 18

Policy transgressions, promises not kept, and fault modelling

“And now remains
That we find out the cause of this effect
Or rather say, the cause of this defect,
For this effect defective comes by cause.”
– Shakespeare, (Hamlet II. ii.100-4).

Non-deterministic systems are usually affected by the arrival of events or random occurrences, some of which are acceptable to system policy and some of which are not. Events which are not acceptable may be called faults. The occurrence of system faults is an extensive and involved topic that is the subject of whole texts (see, for instance [NRC81] and [SS03]). A fuller discussion in terms of promise theory is given in volume 2. This chapter could not cover the breadth of this subject in any detail, rather we attempt to distill one aspect of its essence in relatively simple terms and extract some conclusions from a management perspective, for general synthesis in this work. Readers may consult [SS03] for a survey of technological methods of fault localization in systems.

18.1 Faults and failures

System faults fall into three main categories:
• Random faults: unpredictable occurrences or freaks of nature.

• Emergent faults: faults which occur due to properties of the system which it was not designed for. These usually come about once a system is in contact with an environment.

• Systemic faults: faults which are caused by logical errors of design, or insufficient specification.

How we classify and define faults, seek their causes and remedies, is a difficult topic that we cannot cover here (however, note the remarks in section 9.13, especially example 96). We pursue the subject further in volume 2.

The IEEE classification of computer software anomalies ([IEE]) includes the following issues: operating system crash, program hang-up, program crash, input problem, output problem, failed required performance, perceived total failure, system error message, service degraded, wrong output, no output. This classification touches on a variety of themes all of which might plague the interaction between users and an operating system. Some of these issues encroach on the area of performance tuning, e.g. service degraded. Performance tuning is certainly related to the issue of availability of network services and thus this is a part of system administration. However performance tuning vis-à-vis only peripheral importance compared to the matter of possible complete failure.

Many of the problems associated with system administration can be attributed to input problems (incorrect or inappropriate configuration) and failed performance through loss of resources. Unlike many software situations these are not problems which can be eliminated by re-evaluating individual software components. In system administration the problems are partly social and partly due to the cooperative nature of the many interaction software components. The unpredictability of operating systems is dominated by these issues.

Another source of error is found at the human edge of the system:

• Management errors.

• Forgetfulness/carelessness.

• Misunderstanding/miscommunication.

• Confusion/stress/intoxication.

• Ignorance.

• Personal conflict.

• Slowness of response.

• Random or systematics procedural errors.
Inability to deal with complexity.

Inability to cooperate with others.

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Humans filter all communication through their own view of the world. We respond to things that make sense to us, and we tend to reject things that do not. This can lead to misunderstanding, or only partial understanding of a communicated message (see chapter 15). It can be modelled as the projection of a signal into a digital alphabet that describes our limited domain of understanding. We match input to the closest concept we already know.

Unlike machines, humans do not generally use reliable protocols for making themselves understood (except perhaps in military operations). A system administrator or user can easily misunderstand an instruction, or mis-diagnose a problem.

When a problem arises, it means that an undesirable change has occurred in the system. Debugging is a meta process - a process about the system, but not generally within the system itself. It involves gathering evidence and tracing cause-relationships. A fault might be

- A conflict with policy.
- A logical error.
- A random error.
- An emergent fault.

We can adopt different strategies for solving a problem:

- Mitigate the damage by relieving symptoms.
- Fix the cause of the problem at source.

In many complex systems, it is profitable to employ both of these.

**Example 191.** The immune system which protects higher animals from infections, uses short term counter measures to prevent the spread of infection (chemicals which retard cell replication processes, for instance), while at the same time it is synthesizing a counter-agent to the specific threat (antibodies and killer cells).
18.2 DETERMINISTIC SYSTEM APPROXIMATION

A simplistic, but effective, mode of analysis of systems is to treat them as deterministic boolean directed graphs. These are system representations that classify systems into those that either work or don’t work. It is essentially a dependency analysis of systems, composed of networks of components.

Consider a system of order $n$ components, labelled by a vector or coordinates $\vec{x} = (x_1, x_2, \ldots, x_n)$, joined together by links that transmit information. It is normal to have one of more entry points to this network, and one more more exit points. Information flows along the links from input to output.

A deterministic analysis assumes a two state model, in which:

$$x_i = \begin{cases} 
1 & \text{– if component works} \\
0 & \text{– if component is broken} 
\end{cases}$$

(18.1)

This appears, at first, to be rather simplistic; however, its chief value is in showing that the concept of ‘not working’ can be broken down into a more detailed view in which the source of failure is determined to be a single component. An alternative interpretation of this model is to view the connections between components to mean ‘depends on’ rather than ‘results in’.

Let us define a system configuration in terms of the so-called structure function $\phi(\vec{x})$ that summarizes the dependencies of the system on its components. (When we extend this analysis to include stochastic (random) failure events, this will become a macrostate function.)

**Definition 74 (Structure function).** The system’s state of repair is described uniquely by $\phi(\vec{x})$:

$$\phi = \begin{cases} 
1 & \text{– if the system works} \\
0 & \text{– if the system is broken} 
\end{cases}$$

(18.2)

**Example 192.** Let $x_i, (i = 1, 2, 3, 4)$ describe the CPU, disks and memory, and kernel of a computer database system. The structure function for this system is:

$$\phi(\vec{x}) = x_1 \cdot x_2 \cdot x_3 \cdot x_4.$$  

(18.3)

According to this deterministic model, if a single component is not working, then the entire system is not working.

The example above shows the simple view taken by this analysis. Clearly, there are various degrees by which the memory or disks of a computer system might not work, but we are not able to describe that yet. Nonetheless, there is something to be learned from this simple approach, before we extended it to cope with partial or probabilistic failures.
A serial dependency structure (fig. 18.1) works if and only if each of the components works. The combination is by the boolean AND operation:

\[
\phi(\vec{x})_{\text{serial}} = x_1 \text{ AND } x_2 \text{ AND } \ldots x_n = \prod_{i=1}^{n} x_i = \min_i x_i.
\] (18.4)

A parallel dependency structure (fig. 18.2) works if at least one of its components works. The combination is by the boolean OR operation:

\[
\phi(\vec{x})_{\text{parallel}} = x_1 \text{ OR } x_2 \text{ OR } \ldots x_n = 1 - \prod_{i=1}^{n} (1 - x_i) = \bigvee_{i=1}^{n} x_i.
\] (18.5)

The ‘voting’ gate or \( k \) of \( n \) requires \( k \) out of the \( n \) components to work. If \( k = 1 \), this is a parallel connection; if \( k = n \) it is a serial connection. Clearly this interpolates between these two cases.

So far we have looked at the system from the viewpoint of a random failure of a whole component. The dual description of the system describes the viewpoint of an attacker or saboteur.

**Definition 75** (Dual structure). Given a structure function \( \phi(\vec{x}) \), we can define the dual function \( \phi^D \), or the dual vector \( \vec{x}^D \):

\[
\phi^D(\vec{x}^D) = 1 - \phi(\vec{x}),
\] (18.6)

and

\[
\vec{x}^D = (1 - x_1, 1 - x_2, \ldots, x_n) = \vec{1} - \vec{x}.
\] (18.7)
From the dual viewpoint, we see the vulnerabilities of the system more explicitly. If a system works when only a single component functions (parallel system), then it doesn’t work if \((n - 1)\) additional components are destroyed. If a system works only when all components function, then we have only to destroy 0 additional components to destroy the system.

**NORMALIZATION CRITERIA**

The concept of irrelevant components is a way of identifying and eliminating redundant parameterization in the structure function. It is a way of pruning the graph of irrelevant nodes. We are interested in the relevant components only.

**Definition 76** (Relevant components). The \(i\)th component of a system is relevant to its structure if \(\phi(\vec{x})\) is a non-constant function of \(x_i\), i.e.

\[
\frac{\partial \phi}{\partial x_i} \neq 0,
\]

i.e. the function depends non-trivially on \(x_i\).

Another criterion for only discussing rationally constructed system is to consider only those systems where the repair of a component never makes the system worse\(^1\). Such systems are said to be coherent or monotonic, in reliability analysis.

**Definition 77** (Coherent and monotonic systems). A system of components is said to be coherent if and only if \(\phi(\vec{x})\) is a non-decreasing function of \(x_i\), and all the components are relevant, i.e. iff

\[
\frac{\partial \phi}{\partial x_i} > 0.
\]

It is additionally monotonic if we have \(\phi(\vec{0}) = 0\) and \(\phi(\vec{1}) = 1\), which is equivalent to requiring at least one relevant component.

The requirement of coherence might seem superficially obvious, but if there are mutually exclusive events in a system, parameterized by separate coordinates \(x_i\), the positivity of the structure dependence is not guaranteed.

**Example 193.** Consider a system for providing fault free access to a network server, using a fail-over server. Let \(x_1\) be non-zero if server 1 is active and \(x_2\) be non-zero if server 2 is active. Since the events are mutually exclusive, the structure function is a convex mixture of these:

\[
\phi(\vec{x}) = x_1(1 - x_2) + x_2(1 - x_1).
\]

\(^1\)Note the similarity of this concept to that of convergence in section 5.8
Clearly
\[
\frac{\partial \phi}{\partial x_i} < 0, \quad i = 1, 2. \tag{18.11}
\]

The same example could be applied to different shifts of human workers, in providing a 'round-the-clock service. The fact that the system is non-coherent means that the failure of one server does not leave the system in a non-vulnerable state.

**Redundancy folk theorem**

A folk theorem about redundancy that follows from this simple deterministic model concerns where to arrange for redundancy in a system. Roughly speaking it says that a parallel coupling of components (i.e. a low level parallelism) is never worse than a high level parallelism. In formal terms this follows from two inequalities. Using the notation,
\[
x_i \prod y_j \equiv 1 - (1 - x_i)(1 - y_j), \tag{18.12}
\]
and
\[
\bar{x} \prod \bar{y} = (x_1 \prod y_1, x_2 \prod y_2, \ldots, x_n \prod x_n), \tag{18.13}
\]
we have the inequalities for parallelization
\[
\phi(\bar{x} \prod \bar{y}) \geq \phi(\bar{x}) \prod \phi(\bar{y}). \tag{18.14}
\]
This tells us that the working condition of a system with redundant components is never worse than a redundant combination systems of non-redundant components. For serialization, the opposite is true:
\[
\phi(\bar{x} \cdot \bar{y}) \leq \phi(\bar{x}) \cdot \phi(\bar{y}), \tag{18.15}
\]

i.e. the working condition of a system with strong serial dependencies is never better than a series of dependent systems. The message in both cases is the the lower the level at which we can introduce redundancy, the better.

**Example 194.** Keeping a server reliable using RAID disks arrays, multiple CPUs and error correcting memory, is never worse than keeping two independent systems with single disks, single CPUs and so on.

**Pathways and cut sets**

Two concepts about flow graphs that illuminate their vulnerabilities are pathways and cut sets. We can partition the components in a system into two sets: those that work and those that don’t. If \(\phi(\bar{x})\)
be a monotonic system, then a vector $\vec{x}: \phi(\vec{x}) = 1$, i.e. there is a pathway through the system that works. A minimal path vector is one in which all the components work along the path. A vector is a cut vector is $\phi(\vec{x}) = 0$, i.e. if the vector leads to a broken system. The minimal cut set can then be defined in relation to this, as a set of components such that, if all components in the set are broken, the system is broken.

**Example 195.** Consider the network in fig. 18.3, taking an input on the left of the graph to an output on the right hand side. The minimal path sets are seen by referring to the figure.

$$P_1 = \{1, 4\}, \ P_2 = \{2, 5\}, \ P_3 = \{1, 3, 5\}, \ P_4 = \{2, 3, 4\}. \tag{18.16}$$

The minimal cut sets are

$$C_1 = \{1, 2\}, \ C_2 = \{4, 5\}, \ C_3 = \{1, 3, 5\}, \ C_4 = \{2, 3, 4\}. \tag{18.17}$$

### 18.3 Stochastic System Models

The deterministic analysis above is a useful point of reference that can sometimes be applied directly to human-computer systems. The weakness of the deterministic view of systems is that one cannot ask questions like the following: only 0.1 percent of my disk has an error, what is the likelihood that this will prevent the whole system from working at any given time? Or the probability of reaching the customer help-desk service on the telephone within 30 minutes is 0.3; what is the likelihood that the customer’s enterprise will lose business?

A more flexible analysis of the system treats component states as probabilistic variables. Such a stochastic analysis can also be used in a predictive way, to develop architectural strategies (see chapter 13). It is important for continuum approximations of systems, as only probabilities or averages can vary smoothly and characterize the changing external conditions of a system.
Using the notation of the previous section, we define the probability of a component’s working (i.e. the reliability of the component) as the expectation value of the component’s state:

\[ p_i = P(x_i = 1) = \langle x_i \rangle. \]  

(18.18)

The reliability of the whole system is then given by

\[ \rho \equiv P(\phi(\vec{x}) = 1) = \langle \phi(\vec{x}) \rangle. \]  

(18.19)

\( \rho \) is called the reliability function; it is a generalization of the structure function from boolean estimates to real-valued probabilities.

**Relationship between \( \phi \) and \( \rho \)**

We can see the relationship between these by taking the deterministic limit of the stochastic model, i.e. the limit in which probabilities are either zero or one. Then it follows that they are functionally the same:

\[ \rho(\vec{p}) = \phi(\vec{p}). \]  

(18.20)

This follows since, if \( p_i = \{0, 1\} \), \( x_i = p_i \)

\[ \rho(\vec{p}) = \langle \phi(\vec{x}) \rangle = \phi(\vec{x}) = \phi(\vec{p}). \]  

(18.21)

This tells us how to compute the probabilities of system reliability, given component reliabilities.

**Example 196.** For a serially coupled system:

\[ \rho(\vec{p}) = \langle x_1 \text{ AND } x_2 \text{ AND } \ldots \text{ AND } x_n \rangle \]

\[ = \langle \prod_{i=1}^{n} x_i \rangle \]

\[ = \prod_{i=1}^{n} p_i, \]  

(18.22)

i.e. the combined probability is the product of the component reliabilities.

**Example 197.** For a system with a parallel coupling of components:

\[ \rho(\vec{p}) = \langle \prod x_i \rangle \]

\[ = \langle 1 - \prod_{i=1}^{n} (1 - x_i) \rangle \]

\[ = 1 - \prod_{i=1}^{n} (1 - \langle x_i \rangle) \]

\[ = \prod_{i=1}^{n} p_i. \]  

(18.23)
These rules allow us to generalize the folk theorems about system redundancy in a straightforward manner:

\[
\rho \left( \prod \vec{p} \cdot \vec{p}' \right) \geq \rho \left( \prod \vec{p} \right) \cdot \rho \left( \prod \vec{p}' \right)
\]

\[
\rho \left( \vec{p} \cdot \vec{p}' \right) \leq \rho (\vec{p}) \cdot \rho (\vec{p}').
\]  

(18.24)

In other words: the probability that a single parallelized system is working is greater than or equal to the probability that parallel components will both be working. Conversely, the probability that a system that depends on serialization is working, is always less than the probability that the components are working. Note that there is no impediment to making probabilities into functions of time, in order to track changing conditions.

**Birnbaum Measure of Structural Importance**

One measure of component importance that is related to the minimal pathways and cut sets of the deterministic analysis is the Birnbaum measure, defined as follows.

**Definition 78** (Birnbaum importance). *The partial rate of change of the system reliability \( \rho \) with respect to a given component reliability \( p_i \) indicates its structural dependence on the component.*

\[
I_B^{(i)} = \frac{\partial}{\partial p_i} \rho (\vec{p}).
\]  

(18.25)

This measure is easily calculated from a knowledge of the structure function of the system, and it describes the probability that the \( i \)th component lies in a critical path vector of the system, i.e. that a failure of component \( i \) would lead to a failure of the system. To see this, we note that, for a monotonic system,

\[
\rho (\vec{p}) = p_i \rho (1_i, p_{-i}) + (1 - p_i) \rho (0_i, p_{-i}),
\]  

(18.26)

i.e. The reliability of the system is equal to the probability that \( i \) is working multiplied by the reliability of the system, given that \( i \) is working, plus (OR) the probability that component \( i \) is not working and that the rest of the components’ states are unknown. Using this expansion that gives special prominence to \( i \), we can now examine the Birnbaum measure for component \( i \):

\[
I_B^{(i)} = \frac{\partial}{\partial p_i} \rho (\vec{p})
= \rho (1_i, p_{-i}) - \rho (0_i, p_{-i}).
\]  

(18.27)

This is an expression of the conditional probability that the system is working, given that \( i \) is working, minus the conditional probability that the system is working given that \( i \) is not working; i.e. it is the change in reliability as a result of \( i \) being repaired.
We can rewrite this as follows:

\[
I^{(i)}_B = \Delta_i \rho (\vec{p}) = \langle \phi (1, x_{-i}) \rangle - \langle \phi (0, x_{-i}) \rangle = P ((\phi (1, x_{-i}) - \phi (0, x_{-i}) = 1) = P ((1_i, x_{-i}) is a critical pathway).
\] (18.28)

Put another way, if the change in structure \( \Delta_i \phi = 1 \), then it must mean that the system breaks when \( i \) is destroyed. Thus, the Birnbaum measure tells us the probable importance of the \( i \)th component to overall reliability, assuming that we know the structural form of the system.

**Example 198.** For a system of components in series:

\[
I^{(i)}_B = \frac{\partial}{\partial p_i} \left( \prod_j p_j \right) = \prod_{i \neq j} (1 - p_j).
\] (18.29)

Suppose we order the reliabilities of the components (this results in no loss of generality), so that

\[
p_1 \leq p_2 \leq p_3 \ldots \leq p_n.
\] (18.30)

This implies that

\[
I^{(1)}_B \geq I^{(2)}_B \geq I^{(3)}_B \ldots \geq I^{(n)}_B.
\] (18.31)

Or, the component with the lowest reliability has the greatest importance to the reliability of the system, i.e. “a chain is only as strong as its weakest link”.

**Example 199.** For a system of components in parallel:

\[
I^{(i)}_B = \frac{\partial}{\partial p_i} \left( \prod_j p_j \right) = \prod_{i \neq j} (1 - p_j).
\] (18.32)

Ordering probabilities again,

\[
p_1 \leq p_2 \leq p_3 \ldots \leq p_n.
\] (18.33)

we have that

\[
I^{(1)}_B \leq I^{(2)}_B \leq I^{(3)}_B \ldots \leq I^{(n)}_B.
\] (18.34)

Or, the component with the highest reliability has the greatest structural importance to the overall reliability since, if it has failed, it is likely that all the others have failed too.
Correlations and Dependencies

The interdependence of components can be important in a system in a number of ways. For instance, if one component fails, others might fail too. Or, related components might experience heavy loads or stresses together. In subsystems that are used to balance load between incoming information, the failure of one component might lead to an extra load on the others.

The correlation of component reliabilities is an indication of such interdependence. One measures this using the statistical covariance or un-normalized correlation function of the variables. If variables are associated, then

\[
\text{cov}(x, x') = \langle (x - \langle x \rangle)(x' - \langle x' \rangle) \rangle = \langle xx' \rangle - \langle x \rangle \langle x' \rangle \geq 0.
\] (18.35)

Readers are referred to texts on reliability theory, e.g. [Nat98, HR94], for details on this. We note in passing that ignoring correlations can lead to erroneous conclusions about system reliabilities. The assumption of independence of components \(x_i\) in a serial structure leads to an underestimation of the reliability \(\rho\), in general, whereas the assumption of independence of components in a parallel structure leads to an overestimation of \(\rho\), in general.

18.4 Approximate Information Flow Reliability

One of the aims of building a sturdy infra-structure is to cope with the results of failure. Failure can encompass hardware and software. It includes downtime due to physical error (loss of power, communications, etc.) and also downtime due to software crashes. The net result of any failure is loss of service.

Our main defences against actual failure are parallelism (redundancy) and maintenance. When one component fails, another can be ready to take over. Often it is possible to prevent failure altogether with pro-active maintenance (see the next chapter for more on this issue). For instance, it is possible to vacuum clean hosts, to prevent electrical short-circuits. It is also possible to perform garbage collection which can prevent software error. System monitors (e.g. cfengine) can ensure that crashed processes get restarted, thus minimizing loss. Reliability is clearly a multifaceted topic. We shall return to discuss reliability more quantitatively in section 18.4.

Component failure can be avoided by parallelism, or redundancy. One way to think about this, is to think of a computer system as providing a service which is characterized by a flow of information. If we consider figure 18.2, it is clear that a flow of service can continue, when servers work in parallel, even if one or more of them fails. In figure 18.1 it is clear that systems which are dependent on other series are coupled in series and a failure prevents the flow of service. Of course,
servers do not really work in parallel. The normal citation is to employ a fail-over capability. This means that we provide a backup service. If the main service fails, we replace it with a backup server. The backup server is not normally used however. Only in a few cases can one find examples of load-sharing by switching between (de-multiplexing) services.

Reliability cannot be measured until we define what we mean by it. One common definition uses the average (mean) time before failure as a measure of system reliability. This is quite simply the average amount of time we expect to elapse between serious failures of the system. Another way of expressing this is to use the average uptime, or the amount of time for which the system is responsive (waiting no more than a fixed length of time for a response). Another complementary figure is then, the average downtime, which is the average amount of time the system is unavailable for work (a kind of informational entropy). We can define the reliability as the probability that the system is available:

\[ \rho = \frac{\text{Mean uptime}}{\text{Total elapsed time}} \]  (18.36)

Some like to define this in terms of the Mean Time Before Failure (MTBF) and the Mean Time To Repair (MTTR), i.e.

\[ \rho = \frac{\text{MTBF}}{\text{MTBF} + \text{MTTR}}. \]  (18.37)

This is clearly a number between 0 and 1. Many network device vendors quote these values with the number of 9’s it yields, e.g. 0.99999.

FLOW OF SERVICES

The effect of parallelism, or redundancy on reliability can be treated as a facsimile of the Ohm’s law problem, by noting that service provision is just like a flow of work (see also section B.27 for examples of this).

Rate of service (delivery) = rate of change in information / failure fraction

This is directly analogous to Ohm’s law for the flow of current through a resistance:

\[ I = \frac{V}{R} \]  (18.38)

The analogy is captured in this table:

<table>
<thead>
<tr>
<th>Potential difference V</th>
<th>Change in information</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current I</td>
<td>Rate of service (flow of information)</td>
</tr>
<tr>
<td>Resistance R</td>
<td>Rate of failure or delay</td>
</tr>
</tbody>
</table>
This relation is simplistic. For one thing it does not take into account variable latencies (although these could be defined as failure to respond). It should be clear that this simplistic equation is full of unwarranted assumptions, and yet its simplicity justifies its use for simple hand-waving. If we consider figure 18.2, it is clear that a flow of service can continue, when servers work in parallel, even if one or more of them fails. In figure 18.1 it is clear that systems which are dependent on other series are coupled in series and a failure prevents the flow of service. Because of the linear relationship, we can use the usual Ohm’s law expressions for combining failure rates:

$$R_{\text{series}} = R_1 + R_2 + R_3 + \ldots$$  \hspace{1cm} (18.39)

and

$$\frac{1}{R_{\text{Parallel}}} = \frac{1}{R_1} + \frac{1}{R_2} + \frac{1}{R_3} \ldots$$  \hspace{1cm} (18.40)

These simple expressions can be used to hand-wave about the reliability of combinations of hosts. For instance, let us define the rate of failure to be a probability of failure, with a value between 0 and 1. Suppose we find that the rate of failure of a particular kind of server is 0.1. If we couple two in parallel (a double redundancy) then we obtain an effective failure rate of

$$\frac{1}{R} = \frac{1}{0.1} + \frac{1}{0.1}$$  \hspace{1cm} (18.41)

i.e. $R = 0.05$, the failure rate is halved. This estimate is clearly naive. It assumes, for instance, that both servers work all the time in parallel. This is seldom the case. If we run parallel servers, normally a default server will be tried first, and, if there is no response, only then will the second backup server be contacted. Thus, in a fail-over, model, this is not really applicable. Still, we use this picture for what it is worth, as a crude hand-waving tool.

The Mean Time Before Failure (MTBF) is used by electrical engineers, who find that its values for the failures of many similar components (say light bulbs) has an exponential distribution. In other words, over large numbers of similar component failures, it is found that the probability of failure has the exponential form

$$P(t) = \exp(-t/\tau)$$  \hspace{1cm} (18.42)

or that the probability of a component lasting time $t$ is the exponential, where $\tau$ is the mean time before failure and $t$ is the failure time of a given component. There are many reasons why a computer system would not be expected to have this simple form. One is dependency, which causes events to be correlated rather than independent.

Thus the problem with these measures of system reliability is that they are difficult to measure and assigning any real meaning to them is fraught with subtlety. Unless the system fails regularly, the number of points over which it is possible to average is rather small.
18.5 Fault correction by monitoring and instruction

Let us now use the flow approach of described above to analyze the likely success of a number of common network topologies. Many systems rely on centralized management, but we know that centralization, while a cheap strategy, is fragile since it results in many points of failure. The efficiency of fault correction models has been estimated under different communication infra-structures to analyze the scalability of the solutions (see [BC03]). A simple estimate of the scalability of fault correction can be found by using the system model in chapter 15 for time evolution and error correction.

The simplest estimate is made by assuming that the reliability of each component in a system and each channel is independent of all others, so that the probabilities of resource availability are all independent random variables. This suffices to discuss many aspects of reliability and scaling. If a system component or dependency fails or becomes outdated, a ‘repair’ or update requires a communication with the component from some source of ‘correctness’ or policy.

Let a set of components or resources in a system be defined by a column vector of probabilities

\[ \vec{C} = \begin{pmatrix} p_1 \\ p_2 \\ \vdots \\ p_N \end{pmatrix} \] (18.43)

where \( p_i (i = 1 \ldots N) \) is the probability that component \( i \) is available. If the probabilities are 1, the hosts are said to be reliable, otherwise they are partially reliable.

The channels of information and flow that link the components are represented in the adjacency matrix of the network. This matrix need not be symmetrical in practice, but we shall not address that issue here.

We define a simple measure of the availability of a service, using the connectivity of the graph \( \chi \) (see section 6.2). \( \chi \) has a maximum value of 1, when every node is connected to every other, and a minimum value of zero when all nodes are disconnected.

For a fixed topology and time-independent node availabilities, \( \chi \) is a constant characterizing the network. In general \( \chi \) is time-dependent, as the system evolves; one then obtains a static figure for the network by taking the long-time average:

\[ \langle \chi \rangle = \lim_{n \to \infty} \frac{1}{n} \sum_{i=1}^{n} \chi(t_i). \] (18.44)

The purpose of this measure is that it enables us to gauge and compare different network configurations on equal terms. It is also a measure for comparison by which we can map the problem of unreliable components in a fixed network onto a corresponding problem of reliable components in an ad hoc network.
A DUALITY: AD HOC NETWORKS AND UNRELIABLE COMPONENTS

Ad hoc networks are networks whose adjacency matrices are subject to a strong, apparently random time variation. If we look at the average adjacency matrices, over time, then we can represent the probability of connectivity in the network as an adjacency matrix of probabilities.

Example 200. In an ad hoc communications network, with a fixed number of components, the links are not independent variables. They are constrained both by the physical geography in which the components move (only nearby components are candidates for links), and by interference effects among the set of components near to a given component. Any given component thus may or may not establish a working link with a near component, depending on interference from other near components.

For our purposes here, these dependencies are not important; the important property of the ad-hoc net is the intermittency of the links, due to the components’ mobility.

Definition 79 (Ad hoc adjacency matrix). An ad hoc network is represented by a symmetric matrix of probabilities for adjacency. Thus the time average of the adjacency matrix (for, e.g., four components) may be written as

\[
\langle A \rangle = \begin{pmatrix}
0 & p_{12} & p_{13} & p_{14} \\
p_{21} & 0 & p_{23} & p_{24} \\
p_{31} & p_{32} & 0 & p_{34} \\
p_{41} & p_{42} & p_{43} & 0
\end{pmatrix}
\]  

An ad hoc network is therefore a partially reliable network.

To motivate our discussion further, we note that:

**Theorem 8.** A fixed network of partially-reliable components, \( C_i \), is equivalent to an ad hoc network of reliable components, on average.

**Proof.** This is easily seen from the definition of the connectivity, using a matrix component form:

\[
N(N - 1)\langle \chi \rangle = \bar{C}(p^\prime)^T \langle A(1) \rangle \bar{C}(p) \\
= \bar{C}(1)^T \langle A(pp^\prime) \rangle \bar{C}(1) \\
= \sum_{ij} C_i(p_i) \langle A_{ij}(1) \rangle C_j(p_j) \\
= \sum_{ij} C_i(1) \langle A_{ij}(p_ip_j) \rangle C_j(1). 
\]  

(18.46)
CHAPTER 18. POLICY TRANSGRESSIONS, PROMISES NOT KEPT, AND FAULT MODELLING

The proof demonstrates the fact that one can move the probabilities (uncertainties) for availability from the host vectors to the connectivity matrix and vice versa; for example

\[
\begin{pmatrix}
  p_1 \\
  p_2 \\
  p_3
\end{pmatrix}^T
\begin{pmatrix}
  0 & 1 & 1 \\
  1 & 0 & 1 \\
  1 & 1 & 0
\end{pmatrix}
\begin{pmatrix}
  p_1 \\
  p_2 \\
  p_3
\end{pmatrix} =
\begin{pmatrix}
  1 \\
  1 \\
  1
\end{pmatrix}
\begin{pmatrix}
  0 & p_1p_2 & p_1p_3 \\
  p_2p_1 & 0 & p_2p_3 \\
  p_3p_1 & p_3p_2 & 0
\end{pmatrix}
\begin{pmatrix}
  1 \\
  1 \\
  1
\end{pmatrix}.
\] (18.47)

Thus an array of system components with reliability probabilities \( p_i \), is equivalent to an array of completely reliable components in an unreliable network, where the probability of communication between them is the product of probabilities (assumed independent) from the reliability vector.

**POLICY CURRENT IN A GRAPH**

As networks grow, some system structure topologies do not scale well. We are interested in examining the scaling properties of different configuration management schemes, especially in the context of network models that look to the future of configuration management.

Using even the most simplistic analysis, we can consider a number of cases, in order of decreasing centralization, to find the worst case scaling behaviours. Our discussion follows [BC03].

We assume a simple linear relationship between the probability of successful maintenance and the rates of communication with the policy- and enforcement-sources. This need not be an accurate description of reality in order to lead to the correct scaling laws (see section 12.8). Let us suppose that a change of configuration \( \Delta Q \) is proportional to an average rate of information flow \( I \), over a time \( \Delta t \); that is

\[
\Delta Q = I \Delta t.
\] (18.48)

This equation says that \( I \) represents the time-averaged flow over the interval of time for which it acts. As we are interested in the limiting behaviour for long times, this is sufficient for the job.

Now we apply this simple picture to configuration management for dynamic networks. We take the point of view of a ‘typical’ or ‘average’ host. It generates error in its configuration at the (average) rate \( I_{err} \), and receives corrections at the rate \( I_{repair} \). Hence the rate of increase of error for the average node is:

\[
I_{fail} = (I_{err} - I_{repair}) \theta(I_{err} - I_{repair}).
\] (18.49)

This ordering also corresponds, roughly, to decreasing predictability. However this interpretation may be misleading, since centralized control schemes are also prone to noise, and local or even catastrophic system-wide failures.

The various cases that we consider are presented in Table 18.1 below.
The Heaviside step-function is only non zero when its argument exceeds zero:

$$\theta(x) = \begin{cases} 1 & x > 0 \\ 0 & x \leq 0 \end{cases}$$

and we use it to incorporate the fact that, if the maintenance rate exceeds the error rate, then (on average, over long times) nothing remains outstanding and there is no net rise in configuration error. Thus this averaged quantity is never negative.

If random errors and changes to configuration occur at a rate $I_{err}$ and the configuration agent is unavailable to correct them, then $I_{fail} = I_{err}$. If this holds during a time $\Delta t$, the configuration falls behind by an amount:

$$\text{Bytes missing} = \frac{\text{bytes/sec}}{I_{err}} \times \frac{\text{seconds available}}{\Delta t}.$$  

In the following we will use $p$ to denote the average (over time, and over all nodes) probability that configuration management information flow (repair current) is not available to a node. This unavailability may come from either link or node unreliability. We can lump all the unreliability into the links (see above) and so write

$$p = (1 - \langle A_{ij} \rangle),$$

where $\langle A_{ij} \rangle$ denotes both time and node-pair average. Each node then can only receive repair current during the fraction $(1 - p)$ of the total elapsed time.

The repair current is generated by two possible sources in our models: i) a remote source, and ii) a local source. In each case, the policy can be transmitted and/or enforced at a maximum rate given by the channel capacity of the source. We shall denote the channel capacities by $C_R$ and $C_L$ for remote and local sources for clarity, but we assume that $C_R \sim C_L$, since source and target machines are often comparable, if not identical. If the communication by network acts as a throttle on these rates, then one can further assume that $C_R < C_L$. In any case, the weakest link determines the effective channel capacity. Note that in the case of a confluence of traffic, as in the star models below, the channel capacity will have to be shared by the incoming branches.

We now have a criterion for eventual failure of a configuration strategy. If

$$I_{fail} = \frac{\Delta Q}{\Delta t} > 0,$$

the average configuration error will grow monotonically for all time, and the system will eventually fail in continuous operation. Our strategy is then to look at the scaling behaviour of $I_{fail}$ as the number of nodes $N$ grows large.
Table 18.1: Comparison of models from the viewpoint of the different dimensions: policy dissemination, enforcement, freedom of choice, whether hosts can exchange chosen policy ideas with peers and how political control flows. A ‘push’ model implies a forcible control policy, whereas ‘pull’ signifies the possibility to choose. Model 3 lies between these two, in having the possibility but not the inclination to choose.

<table>
<thead>
<tr>
<th>Model</th>
<th>Application</th>
<th>Enforcement</th>
<th>Policy</th>
<th>Policy Exchange</th>
<th>Control Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Star</td>
<td>Transmitted</td>
<td>No</td>
<td>No</td>
<td>Radial push</td>
</tr>
<tr>
<td>2</td>
<td>Star</td>
<td>Transmitted</td>
<td>No</td>
<td>No</td>
<td>Radial push</td>
</tr>
<tr>
<td>3</td>
<td>Mesh</td>
<td>Local</td>
<td>No</td>
<td>No</td>
<td>Radial pull</td>
</tr>
<tr>
<td>4</td>
<td>Mesh</td>
<td>Local</td>
<td>Yes</td>
<td>No</td>
<td>Radial pull</td>
</tr>
<tr>
<td>5</td>
<td>Mesh</td>
<td>Local</td>
<td>Yes</td>
<td>Yes</td>
<td>Hierarchical pull</td>
</tr>
<tr>
<td>6</td>
<td>Mesh</td>
<td>Local</td>
<td>Yes</td>
<td>Yes</td>
<td>P2P pull</td>
</tr>
</tbody>
</table>

18.6 POLICY MAINTENANCE ARCHITECTURES

Model 1: Star model

The traditional (idealized) model of host configuration is based on the idea of remote management (e.g. using SNMP). Here one has a central manager who decides and implements policy from a single location, and all networks and hosts are considered to be completely reliable. The manager must monitor the whole network, using bi-directional communication. This leads to an $N : 1$ ratio of clients to manager (see fig 18.4). This first model is an idealized case in which there is no

Figure 18.4: Model 1: the star network. A central manager maintains bi-directional communication with all clients. The links are perfectly reliable, and all enforcement responsibility lies with the central controller.
unreliability in any component of the system. It serves as a point of reference.

The topology on the left hand side of fig 18.4 is equivalent to that on the right hand side. We can assume a flow conservation of messages on average, since any dropped packets can be absorbed into the probabilities for success that we attribute to the adjacency matrix. Thus the currents must obey Kirchoff’s law:

$$I_{\text{controller}} = I_1 + I_2 + \ldots I_N.$$  \hspace{1cm} (18.53)

The controller current cannot exceed its maximum capacity, which we denote by $C_S$. We assume that the controller puts out a ‘repair current’ at its full capacity (since the Heaviside function corrects for lower demand), and that all nodes are average nodes. This gives that

$$I_{\text{repair}} = \frac{C_S}{N}. \hspace{1cm} (18.54)$$

The total current is limited only by the bottleneck of queued messages at the controller, thus the throughput per node is only $1/N$ of the total capacity. We can now write down the failure rate in a straightforward manner:

$$I_{\text{fail}} = \left( I_{\text{err}} - \frac{C_S}{N} \right) \theta \left( I_{\text{err}} - \frac{C_S}{N} \right). \hspace{1cm} (18.55)$$

As $N \to \infty$, $I_{\text{fail}} \to I_{\text{err}}$—that is, the controller manages only a vanishing repair current per node. The system fails however at a finite $N = N_{\text{thresh}} = C_S/I_{\text{err}}$. This highlights the clear disadvantage of centralized control, namely the bottleneck in communication with the controller.

**MODEL 2: STAR MODEL IN INTERMITTENTLY CONNECTED ENVIRONMENT**

The previous model was an idealization, and was mainly of interest for its simplicity. Realistic centralized management must take into account the unreliability of the environment.

In an environment with partially reliable links, a remote communication model bears the risk of not reaching every host. If hosts hear policy, they must accept and comply, if not, they fall behind in the schedule of configuration. Monitoring in distributed systems has been discussed in [ALB99].

The capacity of the central manager $C_S$ is now shared between the average number of hosts $\langle N \rangle$ that is available, thus

$$I_{\text{repair}} = \frac{C_S}{N\langle A_{ij} \rangle} = \frac{C}{\langle N \rangle}. \hspace{1cm} (18.56)$$

This repair current can reach the host, and serve to decrease its policy error $\Delta Q$, during the fraction of time $(1 - p)$ that the typical host is reachable. Hence we look at the net deficit $\Delta Q$ accrued
over one “cycle” of time $\Delta t$, with no repair current for $p\Delta t$, and a maximal current $C_S/\langle N \rangle$ for a time $(1-p)\Delta t$. This deficit is then

$$\Delta Q(\Delta t) = I_{err}p\Delta t + \left(I_{err} - \frac{C_S}{\langle N \rangle}\right)(1-p)\Delta t$$

(here it is implicit that a negative $\Delta Q$ will be set to zero). Thus, the average failure rate is

$$I_{fail} = I_{err}p + \left(I_{err} - \frac{C_S}{\langle N \rangle}\right)(1-p) = I_{err} - \frac{C_S}{N}$$

(Again there is an implicit $\theta$ function to keep the long-time average failure current positive.) This result is the same as for Model 1, the completely reliable star. This is because we assumed the controller was clever enough to find (with negligible overhead) those hosts that are available at any given time, and so to only attempt to communicate with them.

This model then fails (perhaps surprisingly), on average, at the same threshold value for $N$ as does Model 1. If the hunt for available nodes places a non-negligible burden on the controller capacity, then it fails at a lower threshold.

**Model 3: Mesh Topology with Centralized Policy and Local Enforcement**

The serialization of tasks in the previous models forces configuration ‘requests’ to queue up on the central controller. Rather than enforcing policy by issuing every instruction from the central source, it makes sense to download a summary of the policy to each host and empower the host itself to enforce it.
There is still a centrally determined policy for every host, but now each host carries the responsibility of configuring itself. There are thus two issues: i) the update of the policy and ii) the enforcement of the policy. A pull model for updating policy is advantageous here, because every host then has the option to obtain updates at a time convenient to itself, avoiding confluence contentions; moreover, if it fails to obtain the update, it can retry until it succeeds. We ask policy to contain a self-referential rule for updating itself.

The distinction made here between communication and enforcement is important, because it implies distinct types of failure, and two distinct failure metrics: i) distance of the locally understood policy from the latest version, and ii) distance of host configuration from the ideal policy configuration. In other words: i) communication failure, and ii) enforcement failure.

![Mesh topology diagram](image)

Figure 18.6: Model 3. Mesh topology. Nodes can learn the centrally-mandated policy from other nodes as well as from the controller. Since the mesh topology does not assure direct connection to the controller, each node is responsible for its own policy enforcement.

The host no longer has to share any bandwidth with its peers, unless it is updating its copy of the policy, and perhaps not even then, since policy is enforced locally and updates can be scheduled to avoid contention.

Let $I_{update}$ be the rate at which policy must be updated. This current is usually quite small compared to $I_{err}$. Based on the two failure mechanisms present here, we break up the failure current into two pieces:

$$ I_{fail} = I_{fail}(i) + I_{fail}(ii) . $$(18.59)
The former term is

\[ I_{\text{fail}}(i) = (I_{\text{err}} - C_L)\theta(I_{\text{err}} - C_L) \]  

(18.60)

this term is independent of \( N \) and may be made zero by design. \( I_{\text{fail}}(ii) \) is still determined by the ability of the controller to convey policy information to the hosts. However, the load on the controller is much smaller since \( I_{\text{update}} \ll I_{\text{err}} \). Also, the topology is a mesh topology. In this case the nodes can cooperate in diffusing policy updates, via flooding, i.e. by using each neighbour to pass on the policy to its neighbours, but never back in the direction it came from.

The worst case—in which the hosts compete for bandwidth, and do not use flooding over the network (graph)—is that, for large \( N \), \( I_{\text{fail}} \rightarrow I_{\text{update}} \). This is a great improvement over the two previous models, since \( I_{\text{update}} \ll I_{\text{err}} \). However note that this can be further improved upon by allowing flooding of updates: the authorized policy instruction can be available from any number of redundant sources, even though the copies originate from a central location. In this case, the model truly scales without limit, i.e. \( I_{\text{fail}} = 0 \).

There is one caveat to this result. If the meshed network of hosts is an *ad hoc* network of mobile nodes, employing wireless links, then connections are not feasible beyond a given physical range \( r \). In other words, there are no long-range links: no links whose range can grow with the size of the network. As a result of this, if the ad hoc network grows large (at fixed node density), the path length (in hops) between any node and the controller scales as a constant times \( \sqrt{N} \). This growth in path length limits the effective throughput capacity between node and controller, in a way analogous to the internode capacity. The latter scales as \( 1/\sqrt{N} \) (see [GK00, LNP90]). Hence, for sufficiently large \( N \), the controller and AHN will fail collectively to convey updates to the net. This failure will occur at a threshold value defined by

\[ I_{\text{fail}}(ii) = I_{\text{update}} - \frac{C_g}{c\sqrt{N_{\text{thresh}}}} = 0 \]  

(18.61)

where \( c \) is a constant. The maximal network size \( N_{\text{thresh}} \) is in this case proportional to \( \left( \frac{C_g}{r_{\text{update}}} \right)^2 \) still considerably larger than for Models 1 and 2.

**MODEL 4: MESH TOPOLOGY WITH PARTIAL HOST AUTONOMY AND LOCAL ENFORCEMENT**

As a variation on the previous model, we can begin to take seriously the idea of distance from a political centre. In this model, hosts can choose not to receive policy from a central authority, if it conflicts with local interests. Hosts can make their own policy, which could be in conflict or in concert with neighbours. Communication thus takes the role of conveying ’suggestions’ from the central authority, in the form of the latest version of the policy.

\(^3\)Note, flooding in the low-level sense of a datagram multicast is not necessarily required, but the effective dissemination of the policy around the network is an application layer flood.
For instance, the central authority might suggest a new version of widely-used software, but the local authority might delay the upgrade due to compatibility problems with local hardware. Local enforcement is now employed by each node to hold to its chosen policy \( P_i \). Thus communication and enforcement use distinct channels (as with Model 3); the difference is that each node has its own target policy \( P_i \) which it must enforce.

![Controller](image)

Figure 18.7: Model 4. As in Model 3, except the hosts can choose to disregard or replace aspects of policy at their option. Question marks indicate a freedom of hosts to choose.

Thus the communications and enforcement challenges faced by Model 4 are the same (in terms of scaling properties) as for Model 3: i.e. \( I_{\text{fail}} \) is the same as that in Model 3. Hence this model can in principle work to arbitrarily large \( N \).

Model 4 is the model used by cfengine ([Bur95, Bur04]). The largest current clusters sharing a common policy are known to be of order \( 10^4 \) hosts, but this could soon be of order \( 10^6 \), with the proliferation of mobile and embedded devices.

**MODEL 5: MESH, WITH PARTIAL AUTONOMY AND HIERARCHICAL COALITION**

An embellishment of Model 4 is to allow local groups of hosts to form policy coalitions that serve to their own advantage. Such groups of hosts might belong to one department of an organization, or to a project team, or even to a group of friends in a mobile network.

Once groups form, it is natural to allow sub-groups and hence a generalized hierarchy of policy refinement through specialized social groups.

If policies are public then the scaling argument of Model 3 still applies since any host could cache any policy; but now a complete policy must be assembled from several sources. Once can thus imagine using this model to distribute policy so as to avoid contention in bottlenecks, since load is automatically spread over multiple servers. In effect, by delegating local policy (and keeping
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Figure 18.8: Model 5. Communication over a mesh topology, with policy choice made hierarchically. Sub-controllers (dark nodes) edit policy as received from the central controller, and pass the result to members of the local group (as indicated by dashed boxes). Question marks indicate the freedom of the controllers to edit policy from above.

a minimal central policy) the central source is protected from maximal loading. Specifically, if there are $S$ sub-controllers (and a single-layer hierarchy), then the effective update capacity is multiplied by $S$. Hence the threshold $N_{\text{thresh}}$ is multiplied (with respect to that for Model 3) by the same factor.

MODEL 6: MESH, WITH PARTIAL AUTONOMY AND INTER-Peer POLICY EXCHANGE

The final step in increasing autonomy is the free exchange of information between arbitrary hosts (peer to peer). Hosts can now offer one another information, policy or source materials in accordance with an appropriate trust model. In doing so, impromptu coalitions and collaborations wax and wane, driven by both human interests and possibly machine learning. A peer-to-peer policy mechanism of this type invites trepidation amongst those versed in traditional control mechanisms, but it is really no more than a distributed genetic algorithm. With appropriate constraints it could equally be made to lead to sensible convergent behaviour, or to catastrophically unstable behaviour.

Before a distributed policy exchange nears a stable stationary point, policy updates could be much more numerous here than for the previous models. This could potentially dominate configuration management behaviour at early times.

Example 201. A collaborative network that has led to positive results is the Open Source Com-
The lesson of Open Source Software is that it leads to a rapid evolution. A similar rapid evolution of policy could also be the result from such exchanges. Probably policies would need to be weighted according to an appropriate fitness landscape. They could include things like shared security fixes, best practices, code revisions, new software, and so on.

Note that this model has no centre, except for a dynamically formed centre represented by centrality (see section 6.5). Hence it is, by design, scale-free: all significant interactions are local. Therefore, in principle, if the model can be made to work at small system size, then it will also work at any larger size.

In practice, this model is subject to potentially large transients, even when it is on its way to stable, convergent behaviour. These transients would likely grow with the size of the network. Here we have confined ourselves to long-time behaviour for large \( N \)—hence we assume that the system can get beyond such transients, and so find the stable regime.

Finally we note that we have only assessed the success of the given models according to their ability to provide an integrity preserving, or error correcting stream, as discussed in chapter 15, that communicates and enforces policy.

18.7 CRITIQUE OF A CURRENT APPROXIMATION

The scaling approximation used in the foregoing models 1-6 is based on the notion of a neutral information current, basically similar to Kirchoff’s laws in physics. The virtue of the model is its
simplicity, but it also glosses over many important features of what is going on in a system. We are justified in using this kind of model when there is weak coupling between the specific semantics of interactions between agents, and the agents themselves are homogeneous in their treatment of the information.

The models do not address semantic aspects of the architectures, such as the degree of centralization (centralization is known to assist in calibration and consistency of shared input and output). Conversely, it does not address the robustness of the peer models, where there are no privileged roles to act as single points of failure. A discussion of these points must wait until volume 2, where we can delve into more details.

18.8 Diagnostic Cause Trees

From the previous sections, we recognize that the causal relationships within a system can form complex networks. Unravelling such networks is difficult. In many cases we can simplify the causal structure by replacing part of the network with an effective tree that more clearly describes the causal relationships. The price for this simplification is that the events are non-deterministic; by hiding details, we lose complete information about the system, but achieve the illusion of a higher level understanding.

Charting cause-trees is a systematic method used in fault diagnosis. The idea is to begin by building lists of possible causes, then causes of those causes, and so on, until one has covered an appropriate level of detail. Once a cause tree has been constructed for a system, it becomes a road-map for fault finding for the future also. The use of cause trees is sometimes called Root Cause Analysis (RCA). A related method called Event Tree Analysis (ETA) maps out every single eventuality, as a true/false binary tree, where every possibility is documented, but only certain pathways actually occur. The latter is mainly a way of documenting the extent of a system; it has little analytical value.

Many of the techniques described in this chapter were pioneered over the last half century by authorities working with nuclear power, where the risk of accidents takes on a whole different level of importance. The keyword in causal analyses is dependencies. All of the immediate causes of an phenomenon or event are called dependencies, i.e. the event depends on them for its existence. The cause tree for the diagnostic example 202 is shown in fig. 18.10. The structure is not completely hierarchical, but it is approximately so.

Example 202 (Network services become unavailable.). A common scenario is the sudden disappearance of a network service, like, say, the WWW. If a network service fails to respond it can only be due to a few possibilities:

- The service has died on the server host.
• The line of communication has been broken.
• The latency of the connection is so long that the service has timed-out.

A natural first step is to try to send a network ping to server-host:

```
ping www.domain.country
```

to see whether it is alive. A ping signal will normally return with an answer within a couple of seconds, even for a machine halfway across the planet. If the request responds with

```
www.domain.country is alive
```

then we know immediately that there is an active line of communication between the our host and the server hosts and we can eliminate the second possibility. If the ping request does not return, then there are two further possibilities

- The line of communication is broken.
- The DNS lookup service is not responding.

The DNS service can hang a request for a long period of time if a DNS server is not responding. A simple way to check whether the DNS server is at fault or not is to bypass it, by typing the IP address of the WWW server directly:

```
ping 128.39.74.4
```

If this fails to respond then we know that the fault was not primarily due to the name service. It tends to suggest a broken line of communication. The traceroute command on Unix-like operating systems, or tracert on Windows can be used to follow a net connection through various routers to its destination. This often allows us to narrow down the point of failure to a particular group of cables in the network. If a network break has persisted for more than a few minutes, a ping or traceroute will normally respond with the message

```
ICMP error: No route to host
```

and this tells us immediately that there is a network connectivity problem.

But what if there is no DNS problem and ping tells us that the host is alive? Then the natural next step is to verify that the WWW service is actually running on the server host. on a Unix-like OS we can simply log onto the server host (assuming it is ours) and check the process table for the httpd daemon which mediates the WWW service.

```
ps aux | grep httpd  BSD
ps -ef | grep httpd  Sys V
```
On a Windows machine, we would have to go to the host physically and check its status. If the WWW service is not running, then we would like to know why it stopped working. Checking log files to see what the server was doing when it stopped working can provide clues or even an answer. Sometimes a server will die because of a bug in the program. It is a simple matter to start the service again. If it starts and seems to work normally afterwards, then the problem was almost certainly a bug in the program. If the service fails to start, then it will log an error message of some kind which will tell us more. One possibility is that someone has changed something in the WWW service’s configuration file and has left an error behind. The server can no longer make sense of its configuration and it gives up. The error can be rectified and the server can be restarted.

What if the server process has not died? What if we cannot even log onto the server host? The latter would be a clear indication that there was something more fundamentally wrong with the server host. Resisting the temptation to simply reboot it, we could then try to test other services on the server host to see if they respond. We already know that the ping service is responding, so the host is not completely dead. There are therefore several things which could be wrong:

- The host is unable to respond (e.g. it is overloaded)
- The host is unwilling to respond (e.g. a security check denying access to our host)

We can check that the host is overloaded by looking at the process table, to see what is running. If there is nothing to see there, the host might be undergoing a denial of service attack. A look at `netstat` will show how many external connections are directed towards host and their nature. This might show something that would confirm or deny the attack theory. An effective attack would be difficult to prevent, so this could be the end of the line for this particular investigation and the start of a new one, to determine the attacker. If there is no attack, we could check that the DNS name service is working on the server-host. This could cause the server to hang for long periods of time. Finally, there are lots of reasons why the kernel itself might prevent the server from working correctly: the TCP connection close time in the kernel might be too long, leading to blocked connections; the kernel itself might have gone amok; a full disk might be causing errors which have a knock-on effect (the log files from the server might have filled up the disk), in which case the disk problem will have to be solved first. Notice how the DNS and disk problems are problems of dependency: a problem in one service having a knock-on effect in another.

A cause tree for diagnosing a full disk is shown in fig. 18.11. This is a particularly simple example; it simply becomes a flat list. Causal analyses can be used at different levels. At the level of human management, it takes on a more heuristic role, e.g.

- Inadequate procedures.
- Inadequate training.
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Figure 18.10: Attempt at cause tree for a missing network service.

Figure 18.11: Attempt at cause tree for a full disk.

- Quality Control.
- Miscommunication.
- Poor management.
- Social/human engineering.
- Supervision error.
- Preventative maintenance lacking.

Information is collected about an incident or phenomenon and this is broken down into cause-effect relationships. Analysts must understand the systems they model thoroughly from the highest levels, down to the component level.

The construction of an event tree is just like the top-down analysis performed in programming. Breaking the event up into component causes is like breaking up a task into subroutines. The benefit is the same: a complex problem is reduced to a structured assembly of lesser problems.
18.9 Probabilistic Fault Trees

How can we go beyond mapping cause and effect to calculating the likely outcomes of the different pathways through a cause tree, to include some of the stochastic reliability analysis from the start of this chapter? This would give us an approximate way of performing reliability analysis based on a kind of spanning tree approximation to diagnosis. To accomplish this, we must acknowledge that not all of the possible pathways occur all of the time: some occur only infrequently, some are mutually exclusive, some are co-dependent and others are uncorrelated. To make serious headway in estimating likely cause, we thus need to add probabilities and combinatorics to the discussion. This is the value of fault tree analysis. The discussion here follows that of Apthorpe ([Apt01]), based on [NRC81].

18.9.1 Faults

For the purposes of modelling, fault tree analysis distinguishes between:

- **Failures**: abnormal occurrences.
- **Faults**: systemic breakdowns within the system.

An important subset of faults is formed by *component faults*.

Component faults fall into three categories:

- **Primary faults**: occur when a component is working within its design limits, e.g. a web server that is rated at 50 transactions per second fails when it reaches 30 transactions per second.
- **Secondary faults**: occur when a fault is operating outside its design specification. e.g. a web server that is rated at 50 transactions per second fails when it reaches 90 transactions per second.
- **Command faults**: are faults that occur when a system performs its specified function, but at the wrong time or place. e.g. a Web server that begins querying a database persistently when no request is being made by an external agent.

Faults occur in response to events. The events are also categorized, this time depending on their position within the tree structure:

- **Top**: This is the top of the tree – the end phenomenon that we are trying to explain. It is analogous to the ‘main’ function in a computer program.
- **Intermediary**: This is a dependency within the tree, but not a root cause of the phenomenon. It is analogous to a subroutine of the main program, it has deeper dependencies that are subroutines of itself.
• Primary: This is an event that is either a root cause, or — as deep an explanation as we can manage to determine. In a computer program analogy, it is like a basic library function, i.e. the lowest level of control available. Events that we cannot say much about are called undeveloped events because although we cannot dig any deeper, we know that there is more going on than we can say. Events that have no further explanation are called basic events. These are the primitive atoms of causality: the very root causes.

Events are drawn according to the symbols in fig. 18.12.

![Figure 18.12: A basic symbols for fault trees.](image)

18.9.2 CONDITIONS AND SET LOGIC

When several smaller causes lead to an intermediate event or phenomenon, there arises a question about how many of the sub-events were needed to trigger the higher level event – all of them? Any of them? a certain number? Events thus combine in ways that can be represented by simple combinatoric set notation – with ‘AND’ and ‘OR’ or other conditions. These are best known to computer scientists in the form of logic gates\(^4\). Figure 18.13 shows the standard symbols for the gates types. Although there are many gate types, for a richness of expression, in practice AND and OR suffice for most cases.

The properties of the gates, in combining the probabilities are noted below. Note that it makes a difference whether or not events are independent, in the probabilistic sense: i.e. the occurrence of one event does not alter the probability of occurrence of another.

\(^4\)One might be forgiven for believing that Boolean logic arrived with digital computers, but this is not the case. Mechanical logic gates may be created, e.g. with hydraulics.
• In OR gates, probabilities combine so as to get larger.

\[ P(A \ OR \ B) = P(A) + P(B) - P(A \ AND \ B). \]  \hspace{1cm} (18.62)

In general,

\[
P(A_1 \ OR \ A_2 \ OR \ \ldots \ A_n) = \sum_{i=1}^{n} P(A_i) - \sum_{i=1}^{n-1} n \sum_{j=i+1}^{n} P(A_i)P(A_j) + \ldots + (-1)^{n+1} P(A_1)P(A_2) \ldots P(A_n). \]  \hspace{1cm} (18.63)

• In AND gates, probabilities combine so as to get smaller.

\[ P(A \ AND \ B) = P(A)P(B|A), \]  \hspace{1cm} (18.64)

or in general:

\[
P(A_1 \ AND \ A_2 \ AND \ \ldots \ A_n) = \prod_{i=1}^{n} P(A_i). \]  \hspace{1cm} (18.65)

If \( A \) and \( B \) are independent, then

\[ P(A)P(B|A) = P(A)P(B), \]  \hspace{1cm} (18.66)
which is smaller than $P(A)$ or $P(B)$; but if the events are not independent, the result can be much greater than this.

- XOR gates have no predictable effect on magnitudes.

$$P(A \text{ XOR } B) = P(A) + P(B) - 2P(A \text{ AND } B) \quad (18.67)$$

Thus if we see many OR pathways, we should be scared. If we see many AND pathways, we should be pleased — the latter means that things are tied down quite tightly with redundancy or protections.

### 18.9.3 Construction

![Fault Tree Diagram](Image)

Figure 18.14: A simple fault tree for an unavailable service.

As a simple example, consider how to work out the probability of failure for a system attack, where an attacker tries the obvious pathways of failure: guessing the root password, or exploiting some known loopholes in services which have not been patched.
We split the tree into two main branches: first try the root password of the system, ‘OR’ try to attack any services which might contain bugs.

- The two main branches are "independent" in the probabilistic sense, because guessing the root password does not change the sample space for attacking a service and vice versa (it’s not like picking a card from a deck).

- On the service arm, we split (for convenience) this probability into two parts and say that hosts are vulnerable if they have a service which could be exploited AND the hosts have not been patched or configured to make them invulnerable.

- Note that these two arms of the AND gate are time-dependent. After a service vulnerability becomes known, the administrator has to try to patch/reconfigure the system. Attackers therefore have a window of opportunity. This adds a time dimension to the fault analysis which we might or might not wish to address.

Since all the events are independent, we have:

\[
P(\text{break in}) = P(A \ OR \ (\ NOT \ A \ AND \ (B \ AND \ C)))
\]

\[
= P(A) + (1 - P(A)) \times P(B) \times P(C)
\]

Suppose we have, from experience, that

- Chance of router problem \(P(A) = 5/1000 = 0.005\).
- Chance of server problem \(P(B) = 50/1000 = 0.05\).
- Chance that server is misconfigured \(P(C) = 10\% = 0.1\).

\[
P(T) = 0.005 + 0.995 \times 0.05 \times 0.1
\]

\[
= 0.005 + 0.0049
\]

\[
= 0.01
\]

\[
= 1\% \quad (18.70)
\]

Notice how, even though the chance of guessing the root password is small, it becomes an equally likely avenue of attack, due to the chance that the host might have been upgraded. Thus we see that the chance of break in is a competition between an attacker and a defender.

A cutset is a set of basic events that are essential for a top level fault to occur. A minimal cutset is a cutset in which the removal of a single event no longer guarantees the occurrence of the top level the event. One of the aims of fault tree analysis is to identify these cut sets. They represent the critical dependencies of the system.
Applications and Further Study 18.

- Analysis of structure and its effects on failure modes.
- Determining the likely place and time window of fault occurrence.
- Quantifying reliability.
- Avoidance of flawed structures.
- Choosing reliable strategies in policy decisions.
CHAPTER 19

DECISION AND STRATEGY

Decisions are made at many levels in network systems, from the top level decisions about policy to the low level choices made during diagnostics. Rational decision making is often assumed central to the optimal performance of human-computer systems. However, humans do not always behave rationally, and even computers do not always behave predictably, so one must take into account the possibility that systems will not always be deterministic.

This chapter is about how one can evaluate the rational limits of strategy in a system, and how one sets choosing the best strategies to maximize the result delivered by the system.

19.1 RATIONAL ACTORS

The dream of rational decision making has a long history in philosophy, going far back in time[Mir89]. The idea that there exists a rational function, which might be minimized to determine the best possible solution to any problem is a part of the heritage that comes from the laws of physics, where energy conservation principles reveal such methods to us in the so-called action principle. Beyond this, the social sciences like economics and sociology have often attempted to describe human behaviour as if it were based on rational minimization of effort. Research in psychology has hopefully disproved this idea once and for all[Kah11, Ari12, Sti90], though it persists especially in the field of economics[NM44].

On short timescales, we know that humans are driven by emotion, not rational computation. If there is an approximation in which human behaviour may be considered rational, it would have to be averaged over time and space at a large ensemble size. Such behaviours might be able to describe crowds, species, or civilizations over evolutionary timescales, but not the minutae of systemic decision making.
Why then should we look at such methods to study systems? One reason is to understand whether the machine aspects of systems, which do not respond emotionally, could be modelled as rational actors. We surely cannot deny that they behave rationally, in the sense of being free of emotion, but we cannot really be sure that they behave optimally, or that the degrees of freedom available to them are free of human bias.

19.2 Causal Analysis

All human and computer systems satisfy basic natural ‘laws’ that tell us how the world works. We cannot escape such laws; they bind us to basic truths about systems, even when the physics of systems seems utterly buried from view. Causality is the term used to express a basic truth about the world: that for every effect there is a cause that precedes it.

Sometimes authors confuse the necessity of this basic law with the ability to identify the precise root of a cause. It is important to realize that effects need not have one simple cause, but that even complex systems that are practically unpredictable obey the law of causality. Causality says simply that physics is a directed graph, at a low level. However, we know from weather charts that when we combine a lot of arrows pointing in different directions, the result can be far from easy to predict. This is why we are often more interested in systems that are predictable than in necessarily being able to trace the exact sequence of changes that led to the state of the system. Still, we are bound by the basic constraint of causality in every decision we make.

When a causally directed system ends up in a state that is not going anywhere, we say that it has reached an equilibrium, as discussed in chapter 10. Such a state is stable to small perturbations in the conditions under which it formed. This concept returns in this chapter in connection with decision making. We would like to ask: what is the likelihood of being able to base decisions on causal, rational information that can tell us a stable kind of ‘truth’ that will not be undone by the smallest change in the environment? The weather forecast suggests to us that this is not going to be easy, but can we at least minimize the uncertainty in decision making by using all of the available information about a human-computer system?

19.3 Decision Making

The simplest decisions are made by associating an action with the occurrence of a state in a chain of events\(^1\). States are identified with actions by combining predicates about the world using logical operations such as **AND**, **OR** and **NOT**. These state-based classifiers are set constructions (see chapter 5).

\(^1\)This is like a switch-case or if-then-else construction in programming
Sets effectively make decisions because they classify the world into regions: inside the set, outside the set, inside this set and that set, and so on. Thus, if we give every set a name, then we have labelled all of the objects that lie within the sets also, and this is what we use to sift through members and identify their properties in terms of a state that is given by the label of the set.

**Example 203.** The system administration tool cfengine ([Bur93]) makes decisions by classifying systems according to their set membership. When the cfengine agent starts executing on any computer, it tries to identify the sets or classifiers to which it belongs:

Defined Classes = ( Saturday Hr12 Min10 Min10_15 Day7 June Yr2003 solaris nexus 32_bit sunos_5_9 sun4u sunos_sun4u_5_9 sparc myname_domain ipv4_128_39_89_10 )

This list of sets identifies the type of operating system, the name of the computer, its Internet address and so on. Notice that even the time is classified into sets that describe the days of the week, the hours of the day and intervals of minutes etc. Any property can be classified in this way, using sets. That is the essence of logic and reasoning.

Some sets do not overlap, or are mutually exclusive:

\[ \text{linux AND windows} = \text{linux} \cap \text{windows} = \emptyset \]  \hspace{0.5cm} (19.1)

Others do overlap and provide nuances of description to decide whether or not to take one course of action or another:

\[
\begin{align*}
\text{linux AND Saturday} & \quad :: \quad \text{action 1} \\
\text{linux AND Sunday} & \quad :: \quad \text{action 2} \\
(\text{linux OR windows}) \text{ AND Monday} & \quad :: \quad \text{action 3}
\end{align*}
\]

Another way in which decisions are made is by statistical confidence: we have processes such as *voting*. If sufficient *support* is given to a certain proposition, we can view it as being statistically ‘true’, or at least *significant*.

Finally we have *games*. The point of game theory is to find out the limits of rational choices with varying degrees of information about competing inputs. If one player has a high expectation of payoff from a particular decision, the game can provide a reality check on those expectations. The results of the game set the baseline for expectation. There will then be noise on top of this, since not all agents in the game were likely to behave rationally.

### 19.4 Game Theory

The Theory of Games was first significantly developed by J. Von Neumann and O. Morgenstern ([NM44]) and later developed by Nash (see [Nas96]) and many others. It is a method for pitting a
set of pre-emptive and defensive strategies against one another, and finding their point of balance in order to see how they fare against one another. By doing this one tries to maximize gain and minimize loss in a competitive setting.

Games are played in many contexts: in fact, a wide variety of interactive processes can be formulated as some kind of game. The simple pendulum, mentioned in chapter 4 is a game that is played between gravity and motion for winning ‘energy’. At a supermarket, customers and merchants play a game with each other in which prices are used to lure customers to a particular seller but also to maximize profits. A balance must be then found between setting the prices as high as possible without losing customers and using a cheaper strategy to sell more items but with a lesser profit per item.

Game theory is applicable in all cases where it is difficult to evaluate the gains generated by following particular policies. This occurs whenever the number of choices is large or the effects are subtle. Contests which are caused by conflicts of interest between system policy and user wishes, unfold in this framework as environmental interactions which tend to oppose convergence and stability (see [NM44, Dre61]). Game theory introduces ‘players’, with goals and aims, into a scheme of rules and then analyzes how much each player can win, according to those restrictions. Each pair of strategies in a game affords the players a characteristic value, often referred to as the ‘payoff’. Game theory has been applied to warfare, to economics (commercial warfare) and many other situations.

Game theory is a vast subject, with many technical challenges. Here, we shall restrict our examples to games between two players, since this is adequate for many situations and presents a sufficient range of issues to exhaust the time allowed in an introductory text.

**WHO ARE THE PLAYERS?**

The players in a game are any actors or influences that affect the transfer of value to any of the other players; i.e. they are entities who exchange some form of system currency. Each player has their own viewpoint of what is best (most valuable) for them but that viewpoint is constrained to work within and share the same system as all of the other players. In some cases, players have opposing interests in which case we speak of non-cooperative games. In other cases players share some common interests and can collaborate leading to partially cooperative games.

Players are labelled by Roman indices $i = 1 \ldots n$ and abstract players can be made to represent many opposing viewpoints about a system:

- System users versus system policy (or system administrator),
- System policy versus entropy - chance degradation of the system,
- A rogue user versus the other $n - 1$ users.
In many games, it is not necessary to interpret a player as a person or rational entity, as one does in classical game theory. As we have seen in chapter 16, the random forces of disorder, measured as entropy, are a sufficient counter-player in many situations. The principle of maximizing entropy is sufficient to make even random chance into a 'rational' player in the game theoretical sense: it is an influence that cares nothing for the other players and which is statistically biased against them. In that sense, it can be viewed as seeking to maximize its own gains. We can therefore think of system administrators as playing a two-person game against 'gremlins' in the system, and this will be a profitable way of formulating the strategic problem.

However, we should be cautious with this viewpoint. There is a slight difference between playing a game against a rational user and playing a game against chance. It is not necessarily true that the most likely outcome of chance is an optimal strategy in a game. When we look for the strategies played by the forces of chance, the most reliable guide is to measure them with experimental data to find out what they actually do, rather than necessarily trusting the formalism of the game that would like to assume the worst. In either case, it is instructive to assume the worst and then compare how efficient chance is at maximizing its effect. If actual data are procured later, they can be substituted and the table elements with a known (sub-optimal) strategy can be summed over to reduce the problem to one of optimization with respect to one less variable.

**Definition 80 (Worst case scenario).** We define this to be the mixed-strategy minimax solution of a zero-sum game. Even if our counter-player is 'nature' or the forces of chance, the equilibrium solution tells us that chance is playing its most destructive hand.

**Reefinement of reasonable belief**

In chapter 17, the issue of learning or the refinement of belief was raised for the attainment of expert knowledge. Decisions must clearly be made based on this kind of expert knowledge. This applies to games or to any other form of decision. If one does not have a reasonable observational basis for making decisions, a strategy of confining the limits of possibility can be used. For example, one begins with the worst case scenario, and then refines it as more data become available. The worst case scenario is bound to be pessimistic, but it is the only rational starting point in the absence of more data.

**Principle 9 (Policy confinement).** In the absence of perfect information about a problem, one adopts a strategy of finding the bounds of reason, and refining these as new information is acquired.

There is thus a synergy between decision theory and Bayesian learning.
Payoff or ‘utility’

What is it that players win or lose in a system administration game? In classical game theory, the payoff has often been money, as game theory was employed as a means for studying economic competition. In section 4.9, we looked at the ways of measuring gain in a human-computer system in terms of system resources or even social capital (status and privilege). There is no simple answer, nor recipe for what the payoff is in a game within the system. If we can formulate a game in which players compete for something, then that is a sufficient justification for doing so. One can imagine payoff being formed from a combination of several importance factors: e.g., memory share, CPU resources, disk space, money, privilege, time for human recreation, and so on.

Payoff is represented by a function $\Pi_i$ for each player $i$. For two players, this function is a matrix with a finite number of pure strategies $s_i$. Games fall into two distinct types: constant sum games and non-constant sum games. In a constant sum game, each element of the payoff sums to a constant value over all players:

$$\sum_{i=1}^{n} \Pi_i = \text{const} \times 1,$$  \hspace{1cm} (19.2)

where $1$ is a matrix or table filled with ones.

What is a strategy?

A pure strategy is a single course of action taken by a player throughout a game. We can think of it as a mode of behaviour. There are two interpretations of strategies: in the extensive form of a game, a strategy represents a single set of moves by a single player from start to finish; in the strategic or ‘normal’ form of the game, the strategy represents an average mode of play, without specifying the details of individual moves.

Different courses of action lead to different returns or payoffs, and the point of the game is to compare the results of using different strategies on the final payoff. The method of solution of a game is to vary each player’s strategy, or mixture of strategies so as to optimize the amount they can win.

Suppose that the set of all pure strategies $s_i$ for player $i$ is denotes by $S_i$, so that $s_i \in S_i$. The set of all players’ strategies is denoted by the outer product:

$$S = S_1 \times S_2 \times \ldots \times S_n.$$  \hspace{1cm} (19.3)

Sometimes it is not advisable for players to play a single strategy, but to mix several different approaches to playing a game. For instance, we might discover that it pays more to play one strategy half the time and a different strategy the remainder of the time. This is expressed by defining mixtures of pure strategies.

\[^2\text{The payoff is also called the player’s utility in many texts.}\]
A mixed strategy is a probability distribution over pure strategies, and is denoted \( \sigma_i \) for player \( i \). In other words, if player \( i \) players strategy \( \alpha \) with probability \( P(\alpha) \) then,

\[
\sigma_i(\alpha) = P_i(\alpha). \tag{19.4}
\]

Clearly the sum of probabilities for all alternative strategies is one, for every player, so:

\[
\sum_\alpha \sigma_i(\alpha) = 1, \forall i. \tag{19.5}
\]

Mixed strategy probabilities can be interpreted in various ways:

- The average play over time within a single execution of a game.
- The likelihood of choosing a particular pure strategy on repeated invocations of similar games.
- The average strategies of multiple players of a game, over multiple trials.

Mixed strategies are important because they make the theory of games into a tool for statistical inference. A certain randomness of strategy can often compensate for uncertainty by randomly hitting a randomly moving target.

**The value for a player**

The value of what is earned or ‘won’ by a player in a game is given by the scalar product of the payoff function \( \Pi_i \) with the strategy profiles of the users.

The value of any player is weighted by the choices made by all the players. Thus no player can win an arbitrary amount, without other players being able to downgrade their potential payoff by counter-play.

**Example 204. Consider a two person game with payoff matrix**

\[
\Pi_1 = \begin{pmatrix} 4 & 5 & 6 \\ 2 & 8 & 3 \\ 3 & 9 & 2 \end{pmatrix} \tag{19.6}
\]

for player 1 and payoff matrix

\[
\Pi_2 = \begin{pmatrix} 3 & 1 & 2 \\ 1 & 4 & 6 \\ 0 & 6 & 8 \end{pmatrix} \tag{19.7}
\]
for player 2. These two matrices are often combined as follows:

\[
\Pi_{(1,2)} = \begin{pmatrix}
(4, 3) & (5, 1) & (6, 2) \\
(2, 1) & (8, 4) & (3, 6) \\
(3, 0) & (9, 6) & (2, 8)
\end{pmatrix}.
\]  

(19.8)

Now let \( \sigma_1^T \) be the transpose of a general mixed strategy vector for player 1, i.e.

\[
\sigma_1^T = (P(s_1), P(s_2), P(s_3)) = \left(\frac{1}{3}, \frac{1}{3}, \frac{1}{3}\right),
\]  

(19.9)

and let \( \sigma_2 \) be the mixed strategy vector for player 2, i.e.

\[
\sigma_2 = \begin{pmatrix}
P(s_1) \\
P(s_2) \\
P(s_3)
\end{pmatrix} = \begin{pmatrix}
0 \\
\frac{1}{2} \\
\frac{1}{2}
\end{pmatrix}.
\]  

(19.10)

The value of the payoff to player 1

\[
v_1 = \sigma_1^T \Pi_1 \sigma_2 = \left(\frac{1}{3}, \frac{1}{3}, \frac{1}{3}\right) \begin{pmatrix}
4 & 5 & 6 \\
2 & 8 & 3 \\
3 & 9 & 2
\end{pmatrix} \begin{pmatrix}
0 \\
\frac{1}{2} \\
\frac{1}{2}
\end{pmatrix} = \frac{11}{2}.
\]  

(19.11)

The value of the game for player 2 is

\[
v_2 = \sigma_1^T \Pi_2 \sigma_2 = \left(\frac{1}{3}, \frac{1}{3}, \frac{1}{3}\right) \begin{pmatrix}
3 & 1 & 2 \\
1 & 4 & 6 \\
0 & 8 & 2
\end{pmatrix} \begin{pmatrix}
0 \\
\frac{1}{2} \\
\frac{1}{2}
\end{pmatrix} = \frac{27}{6}.
\]  

(19.12)

19.5 The Strategic Form of a Game

The strategic or normal form of a game consists of a number of players, strategies and rewards or payoffs that result from the use of the strategies.

1. A set of players \( i = 1, 2, \ldots n \).

2. Sets of pure strategies \( S_i \) for each player \( i \). The vector \( \bar{s} = (s_1, s_2, \ldots, s_n) \), where \( s_i \in S_i \) is called a strategy profile for the game, i.e. a choice of strategies for each player. Note that each \( s_i \) is also a vector whose number of elements is the number of pure (independent) strategies available to player \( i \).

3. A function \( \Pi_i(s) \) for player \( i \) that describes the player’s payoff when a certain combination of strategies \( s \) is played by all the players.
Example 205. A simplistic formulation of a game to weigh the advantages and disadvantages of upgrading software by various methods. The payoff can be thought of as the level of ‘convenience’ to the players. Thus the row player, who is the system administrator, considers the advantage to the system, while the column player, who represents the users of the system, considers the advantage to themselves.

<table>
<thead>
<tr>
<th></th>
<th>Security hole</th>
<th>Bug in function</th>
<th>Missing function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upgrade version now</td>
<td>(10,5)</td>
<td>(10,0)</td>
<td>(5,-5)</td>
</tr>
<tr>
<td>Test then upgrade</td>
<td>(5,5)</td>
<td>(3,9)</td>
<td>(0,8)</td>
</tr>
<tr>
<td>Keep parallel versions</td>
<td>(-10,5)</td>
<td>(-1,10)</td>
<td>(0,10)</td>
</tr>
</tbody>
</table>

The system administrator believes that the maximum advantage (10) to the system arises from upgrading software immediately when faced with a security hole, while the hostile user is dealt a maximum blow by a quick upgrade so this is also of advantage to the users of the system, who do not have the same level of advantage (5) from the strategy since they are perhaps protected in other ways, with redundancy and backup. If the users are missing some important functionality that only exists in a newer version of the program they have a high level of payoff by getting the upgrade quickly (8), however new versions often incorporate new bugs, so parallel versions give the maximum benefit to users (10), but neither of these cases are of any great interest to the system administrator who does not use the software.

If there is a bug in the software, the administrator benefits from upgrade by not having to deal with irate users (3), while users clearly benefit from upgrade (9). Again parallel versions suit users (10) but might disadvantage administrators (-1) since multiple versions often present technical and administrative challenges to administrators.

We can continue in this way, posing values for the payoffs. The payoff values here certainly depend on other factors than we have considered in the primitive example, e.g. how reliable the new versions tend to be from the software producer. A better model of this game would take these explicitly into account.

The real challenge in formulating strategic form games is how to model the payoffs using actual numbers. Inspired guesswork is the most primitive level of approximation. One might imagine that this would not lead to any useful result, but often surprises result when the game is solved. Games automatically take into account all of the competing forces in the game, which often leads to results that are not easily seen or guessed from the individual estimations of payoffs.

Measurement of the system over time is another way to develop payoff models. A semi-empirical model can easily be used to gauge the relative advantages of different strategies. This also allows one to introduce dimensions such as time variation into the games. Note however that a game has no concept of causal time in its strategic form, with one strategy leading to another. Time becomes at best an average parameter changing conditions for the whole game.
19.6 THE EXTENSIVE FORM OF A GAME

How do we find the strategic payoffs from a game that has many complicated moves and courses of action? In a board game, such as chess for instance, we normally think of a complex interaction of the players’ moves and counter-moves. In system administration, games between users and the system might extend over considerable numbers of moves and counter moves. The extensive form of a game is thus based on a complete tree of every detailed possible move or sub-decision between the players (fig. 19.1). It is a way of mapping out the behaviour of the players causally,

![Game Tree Diagram]

Figure 19.1: The extensive form of a game is a complete history of the state space of the players. By tracing the choices made from the start of the game, a player can examine the payoff accrued and predict the best course of action for the remaining moves.

though not necessarily deterministically. The decision trees in game theory are related to the fault trees in chapter 18: we can examine the tree for the players’ moves in turn (one moves, then the other counter-moves etc), or we can separate the two decision trees of the players, if we do not necessarily know the order in which the moves are made. The two cases are referred to as games with perfect and imperfect information, respectively.

The extensive form consists of a number of players, a game tree and a set of payoffs. A game tree is a graph \((x, \Gamma)\) consisting of nodes and edges or arcs (see section 6.2). Each edge represents a move in the game, and each node is a player whose turn it is to make a decision. The graph for the game must be a tree, i.e. every node must have exactly one path back to a root node, i.e. must have a unique parent. This is somewhat like a state diagram, except that the same ‘state’ in a system can appear several times in the game tree, if it can be reached by a number of different means. The game tree codifies the history of the transitions as well as the actual states of the players.
A player in a game might have complete knowledge of the game tree when it is his or her turn to move, but he might also not be able to remember how he arrived at a given node – thus, if several nodes have the same state, there might be several alternatives to play that seem appropriate. In fact, only one choice is the true choice, but the player might not be able to determine this and could believe that the best course of action forward is from one of the other nodes that look the same. Game play can therefore also describe situations where the players have perfect recall of the entire game history, and situations where that information is lost.

Example 206. In an organization, records are usually kept only for a certain length of times. In a computer system, logs of system events are rotated and deleted after a certain length of time. In each case, information must eventually be forgotten due to limited resources. This might affect the ability of system administrators and managers to determine the best course of action forward in time.

The extensive form of a game can finally be reduced to a strategic form, by summing the payoffs of the individual pathways in the game. Two extensive forms of a game are said to be equivalent if they reduce to the same strategic form.

In this book, we shall not look at extensive games in any detail; such a topic could probably be a book in its own right. The strategic form of game theory will be the most immediately useful form for ordinary decision making, however the extension to using causal trees as an interpretation of the extensive form of the game allows more detailed analyses of system dependencies by game theoretical means.

19.7 SOLVING ZERO SUM GAMES

Zero sum games are games that satisfy conservation of pay-off constraints, i.e. we are neither allowed to create nor destroy the currency of the game. This is a familiar idea in the physical world, where energy is conserved for closed systems. Certain simplifications arise for this type of game, as a result of this constraint. The basic approach to solution of zero-sum games begins with the minimax theorem, due to J. Von Neumann.

Consider a two person zero sum game, with payoff matrices \((\Pi_1, \Pi_2)\) and pure strategy sets \((S_1, S_2)\). The pay-off matrices satisfy

\[
\Pi_2(s_1, s_2) = -\Pi_1(s_1, s_2), \forall s_1 \in S_1, s_2 \in S_2.
\]  

(19.13)

The minimax theorem tells us that all games have a solution that is expressible in terms of mixed strategies, and games that have an immediate saddle-point have a solution in terms of pure strategies. The minimax theorem says that it is always possible to find a pair of mixed strategies \((\sigma_1, \sigma_2)\) such that there is a unique equilibrium between the players that gives value \(v_1\) to player 1 and \(-v_2\) to
CHAPTER 19. DECISION AND STRATEGY

player 2:

\[ v_1 = \max_{\sigma_1} \min_{\sigma_2} \sigma_1^T \Pi_1 \sigma_2 = \min_{\sigma_2} \max_{\sigma_1} \sigma_1^T \Pi_1 \sigma_2. \]  \tag{19.14}

Moreover, the limiting case of the theorem occurs when the payoff matrix alone has a saddle-point, i.e.

\[ \max_{\sigma_1} \min_{\sigma_2} \Pi_1 = \min_{\sigma_2} \max_{\sigma_1} \Pi_1. \]  \tag{19.15}

For the two player game, this condition is very easy to check, by looking along the rows and the columns of the payoff matrix for either of the players.

**Example 207.** The following game has a pure strategy saddle-point. Let the payoff or utility matrix for player A be given by:

\[ \Pi_A = \begin{pmatrix} 1 & -3 & -2 \\ 2 & 5 & 4 \\ 2 & 3 & 2 \end{pmatrix}. \]  \tag{19.16}

This game is zero-sum, so \( \Pi_B = -\Pi_A \). We look for a saddle-point:

\[ \max \min \Pi_A = \max \begin{pmatrix} -3 \\ 2 \\ 2 \end{pmatrix} = 2. \]  \tag{19.17}

\[ \min \max \Pi_A = \min(2, 5, 4) = 2. \]  \tag{19.18}

Since these two values are equal, we have found the value of the game \( v = 2 \), and see two optimal strategy saddle-points, with row-column coordinates: \( (r^*, c^*) = (2, 1) \) and \( (3, 1) \).

**19.8 DOMINATED STRATEGIES**

We wish to discuss varying the strategies of a single player \( i \), while holding the opponents strategies fixed. Let \( s_i \) be an element of \( S_i \), the strategy space of player \( i \), and let \( s_{-i} \) denote a strategy selection for every player other than \( i \) (i.e. using the set notation from section 5.1, this is the strategy for the set \( -i \) or ‘not’ \( i \)). We can now write a complete strategy selection for all players, but singling out \( i \) as \( (s_i, s_{-i}) \). Better still, we can draw attention to the fact that we are looking at a trial strategy for \( i \) by writing \( s_i \rightarrow t_i \), so that a complete strategy profile is given by:

\[ \vec{s} = (t_i, s_{-i}). \]  \tag{19.19}
Similarly, for mixed strategies, we can write $(\sigma_i, \sigma_{-i})$ or

$$\bar{\sigma} = (\tau_i, \sigma_{-i}).$$  \hspace{1cm} (19.20)

We say that a pure strategy $t_i$ is strictly dominated for player $i$ if

$$\Pi_i(\sigma_i, s_{-i}) > \Pi_i(t_i, s_{-i}), \forall s_{-i},$$  \hspace{1cm} (19.21)

i.e. a player is always better off using some other mixture of strategies than choosing $t_i$, regardless of what the other players do. If the strict inequality above is replaced by a weak inequality $\geq$, the strategy is said to be weakly dominated. Notice that means that the definition in eqn. (19.21) also applies for any opponent mixed strategies, since $\sigma_{-i} = \sum_i p_i s_{-i}$, but all $s_i$ are covered in this relation, and all $p_i \leq 1$, hence $\sigma_{-i} \leq s_{-i}$. Similarly, given any pure strategy that is dominated, a mixed strategy that gives non-zero weight to this strategy is also dominated.

### 19.9 Nash Equilibria

The Nash equilibrium is probably the most important solution concept in game theory. For two-person zero-sum games, it corresponds to the minimax saddle-point for mixed strategies; however, it also generalizes this concept as it is not limited to zero sum games. Nash proved that all games have at least one equilibrium in terms of mixed strategies.

The concept of a Nash equilibrium is related to the idea of fixed points and equilibria encountered in chapter 10. It is most easily formulated for the strategic form of the game. Suppose a game has $n$ players with pure strategy sets $S_i$, and payoff functions $\Pi_i : S \rightarrow R^1$ for $i_1, \ldots, n$. The set or space of all random strategy profiles is defined by

$$\Sigma = \sigma(s_1) \times \sigma(s_2) \times \cdots \times \sigma(s_n) = \times_{i \in n} \sigma(s_i).$$  \hspace{1cm} (19.22)

A Nash equilibrium is a mixed strategy profile $\sigma^*$ for each and every player, such that each player’s strategy is an optimal response to all of the other’s strategies, i.e.

$$\Pi_i(\tau^*_i, \sigma^*_{-i}) \geq \Pi_i(\sigma_i, \sigma^*_{-i}), \forall \sigma_i \in \Sigma_i.$$  \hspace{1cm} (19.23)

The Nash equilibrium is related to the Kakutani fixed point theorem, by forming a correspondence between every mixed strategy and its optimal response. Let us define the optimal response mapping as the function $R_i$ that maps a certain combination of opponents strategies $\sigma_{-i}$ to an optimal strategy $\sigma_i$ for player $i$:

$$\sigma_i = R_i(\sigma_{-i}) = \arg \max_{\tau \in \Sigma} \Pi_i(\tau, \sigma_{-i}),$$  \hspace{1cm} (19.24)
i.e. it selects the value of the argument that maximizes the payoff and returns it as its value. Although this function does not need to know the value of \( \sigma_i \), since it actually selects it, it does no harm to make \( R_i \) functionally dependent on it in a trivial way, i.e. we can equally well write this for all the players’ \( \sigma_i \):

\[
\sigma_i = R_i(\sigma).
\] (19.25)

Finally, we can form the product correspondence of all of these functions for all the players:

\[
R(\sigma) = R_1(\sigma) \times R_2(\sigma) \times \ldots \times R_n(\sigma).
\] (19.26)

The Nash equilibrium is then defined as strategy profile \( \sigma^* \) which is the fixed point of this correspondence:

\[
\sigma^* = R(\sigma^*).
\] (19.27)

Nash used this construction to prove that such a fixed point must exist in a finite game, for mixed strategies.

**Example 208** (Competition or cooperation for service?). Consider, for simplicity, just two customers or users \( A \) and \( B \) who wish to share a service resource. We shall assume that the service ‘market’ is a free-for-all; i.e. no one player has any a priori advantage over the other, and that both parties behave rationally.

The users could try to cooperate and obtain a ‘fair’ share of the resource, or they could let selfish interest guide them into a competitive battle for largest share. The cooperation or collaboration might, in turn, be voluntary or it might be enforced by a service provider.

These two strategies of competition and collaboration are manifestly reflected in technologies for networking, for instance:

- **Voluntary sharing:** Ethernet is an example of voluntary sharing, in which any user can grab as much of a share as is available. There is a maximum service rate that can be shared, but it is not necessarily true that what one user loses is automatically gained by the other. It is not a zero-sum game.

- **Forced sharing:** Virtual circuits (like MPLS, ATM or Frame Relay networking technologies) are examples of forced sharing, over parallel circuits. There are thus fixed quotas that enforce users’ cooperation. These quotas could be allocated unevenly to prioritize certain users, but for now we shall assume that each user receives an equal share of the resource pot.

We analyze this situation, in a very simple way, using a classic game theoretical approach. The customers can ‘win’ a certain amount of the total service rate \( R \) (e.g. bytes per second, in the case
Table 19.1: A’s payoff matrix in two customer sharing.

<table>
<thead>
<tr>
<th></th>
<th>B Cooperate</th>
<th>B Compete</th>
</tr>
</thead>
<tbody>
<tr>
<td>A Cooperate</td>
<td>( \frac{R}{2} )</td>
<td>( \frac{R}{2} - \delta R )</td>
</tr>
<tr>
<td>A Compete</td>
<td>( \frac{R}{2} + \delta R )</td>
<td>( R_c )</td>
</tr>
</tbody>
</table>

Table 19.2: B’s payoff matrix in two customer sharing.

<table>
<thead>
<tr>
<th></th>
<th>B Cooperate</th>
<th>B Compete</th>
</tr>
</thead>
<tbody>
<tr>
<td>A Cooperate</td>
<td>( \frac{R}{2} )</td>
<td>( \frac{R}{2} + \delta R )</td>
</tr>
<tr>
<td>A Compete</td>
<td>( \frac{R}{2} - \delta R )</td>
<td>( R_c )</td>
</tr>
</tbody>
</table>

Table 19.3: A, B combined payoff matrix in two customer sharing. This is the usual way of writing the payoff matrices. We see that, when both customers collaborate (either willingly or by forced quota), they obtain equal shares. If one of them competes greedily, it can obtain an extra \( \delta R \) that is then subtracted from the other’s share. However, if both users compete, the result is generally worse \( (R_c) \) than an equal share.

<table>
<thead>
<tr>
<th></th>
<th>B Cooperate</th>
<th>B Compete</th>
</tr>
</thead>
<tbody>
<tr>
<td>A Cooperate</td>
<td>( \frac{R}{2} ), ( \frac{R}{2} )</td>
<td>( \frac{R}{2} - \delta R, \frac{R}{2} + \delta R )</td>
</tr>
<tr>
<td>A Compete</td>
<td>( \frac{R}{2} - \delta R, \frac{R}{2} + \delta R )</td>
<td>( R_c, R_c )</td>
</tr>
</tbody>
</table>

Thus, we assume that each of the users assumes an equal share \( R/2 \) when they cooperate with one another. The relative sizes of the payoff are important. We have:

\[
\delta R \leq \frac{R}{2} \tag{19.28}
\]

\[
\left( \frac{R}{2} - \delta R \right) \leq \frac{R}{2} \leq \left( \frac{R}{2} + \delta R \right) \tag{19.29}
\]

In other words, by competing, a selfish user might be able to gain an additional amount of the service capacity \( \delta R \) to the other’s detriment. The sum of both users’ shares cannot exceed \( R \). If both users choose to compete, the resulting competition might lead to an amount of waste that goes to neither of the users. This is the case in Ethernet, for instance, where collisions reduce the efficiency of transmission for all parties equally. We model this by assuming that both users then obtain a share of \( R_c < R/2 \).

This leaves us with two separate cases to analyse:

1. \( R_c > R/2 - \delta R \): If the result from competitive ‘attacks’ against one another is greater than the result that can be obtained by passively accepting the other customer’s aggressiveness, then we are inclined to retaliate. This becomes an instance of the Prisoner’s Dilemma game. It has a solution in terms of Nash equilibria by dominant strategies.

2. \( R_c < R/2 - \delta R \): If the payoff for mutual competition is less than than penalty for
CHAPTER 19. DECISION AND STRATEGY

Figure 19.2: With $R_c > R/2 - \delta R$, the game becomes a classic game theoretical problem of ‘Prisoner’s Dilemma’. The dominant Nash equilibrium is where both players decide to compete with one another. If the customers are altruistic and decide to collaborate (or are forced to collaborate) with one another, they can win the maximum amount. However, if they know nothing about each other’s intentions then they realize, rationally, that they can increase their own share by $\delta R$ by choosing a competitive strategy. However, if both choose to be competitive, they cannot achieve exactly this much: the balance point for mutual competition is $R_c$. This value is determined by the technology used by the service provider. If either one of the players decided to cooperate with the other, they would lose.

Figure 19.3: With $R_c < R/2 - \delta R$ the game becomes another classic game of Dove-Hawk. If both players are ‘hawkish’ and attack greedily, they both lose out. The stable equilibria are that one player is greedy and the other is submissive.

We can ask if there is a mixed strategy of partial cooperation that would succeed at counter-
ing the poor result from mutual competition, but which yield slightly more. To show that this is not the case, let us pick B’s strategy and then allow A to choose cooperation with probability p:

(a) B cooperates: compare then the payoffs for A and B and ask, is there a value of p such that

\[ p \frac{R}{2} + (1 - p) \left( \frac{R}{2} + \delta R \right) > p \frac{R}{2} + (1 - p) \left( \frac{R}{2} - \delta R \right)? \] (19.30)

Either we must have \( \delta R = 0 \) or \( p = 0 \), so the answer is no: there is no way to improve on this strategy as long as there is something to gain from competition.

(b) B competes: compare then the payoffs for A and B and ask, is there a value of p such that

\[ p \left( \frac{R}{2} - \delta R \right) + (1 - p)R_c > p \left( \frac{R}{2} + \delta R \right) + (1 - p)R_c? \] (19.31)

Again, consistency forces us to take \( p = 0 \) or \( \delta R = 0 \).

These simple games capture the essence of the issues involved in sharing. They reflect both human strategies for competition and technological ones. We see that there is no clear answer as to whether Ethernet (Hawkish) or fixed quota virtual circuit (Dove-like) behaviour is preferable, it depends on the level of traffic.

Example 209. The model in the previous example addresses many situations. We can use it for human scheduling too. The basic result from game theory, although simplistic, tells us that random event competition works well as long as the number of jobs is small, i.e. as long as there is plenty of time and no one is taxed to the limit. However, as a human becomes very busy, it becomes better to schedule fixed quota time-slices for jobs, otherwise all jobs tend to suffer and nothing gets done. The overhead of swapping tasks at random can lead to great inefficiency if time is short.

19.10 A SECURITY GAME

Electronic banking, and other on-line services that require users to identify themselves in secure way, use a variety of mechanisms to achieve this authentication. Many banks issue ‘calculator’-like smartcards that generate one-time passwords based on a personal code and the time of day. Others use a Transmission Layer Security (TLS) certificate mechanism to download a secret key from the bank, using a login and Personal Identification Code that they receive from the bank by postal mail. At the lowest end level, some sites are password protected, or could use some kind of biometric

---

3TLS was formerly known as Secure Socket Layer (SSL).
secret. The attacks that can be levelled against these measures include actually stealing a user’s credentials, hijacking the TCP session, spoofing the web site and placing a Trojan horse in the user’s browser.

Let us formulate a game based on the a simple system versus adversary model. At the most simplistic level, security can be regarded as a zero-sum game. What is secured properly, is lost to the potential attacker and what is not secured is gained. However, if we add in the cost of buying or developing and then maintaining the security system, the ‘gains’ of the defender are somewhat mitigated by the cost of ownership. This would make the game non zero-sum. We can therefore examine the game in stages. We begin by defining the payoffs in the ‘game’.

We estimate the payoffs on a scale of 0-100%. We shall consider the payoff to the defender of the system, so that the payoff is essentially the same as the estimated level of security. Suppose that the security levels are given as in table below for defence and attack \((D, A)\).

<table>
<thead>
<tr>
<th>Security</th>
<th>Steal credentials</th>
<th>Hijack session</th>
<th>Spoof site</th>
<th>Trojan in browser</th>
</tr>
</thead>
<tbody>
<tr>
<td>Smartcard</td>
<td>(90,10)</td>
<td>(90,10)</td>
<td>(40,60)</td>
<td>(30,70)</td>
</tr>
<tr>
<td>Certificate</td>
<td>(70,30)</td>
<td>(80,20)</td>
<td>(70,30)</td>
<td>(10,90)</td>
</tr>
<tr>
<td>Password</td>
<td>(50,50)</td>
<td>(70,30)</td>
<td>(10,90)</td>
<td>(10,90)</td>
</tr>
</tbody>
</table>

These values are posed only for illustration, not actually measurements or estimates, but we can justify them approximately as follows. Smart cards are difficult to steal, since they can be carried with a person and kept safe; thus they offer a high level of security to theft. Certificates are a little easier to steal, since they reside on the user’s computer and could therefore be eavesdropped or extracted somehow. Passwords are easier still to steal or tease out of users by social engineering or eavesdropping.

If an attacker can hijack a session smartcard systems require code confirmations of all important operations so without the smartcard such session hijacking would not be a useful strategy. Certificates are slightly less secure, since they can sometimes be extracted from transactions; again, passwords offer the weakest security.

An attacker who spoofs a web site can trick a user with a smartcard to enter a code that could be used by the attacker instead for a short interval of time, so the security level to spoofing is quite low. A certificate a somewhat better here, however, since it is used to encipher information for the specific site, and thus would provide only nonsense to a spoofed web site. Passwords would be trivially collected at a spoof site. They offer essentially no security.

Finally, if an attacker can sneak Trojan code into the user’s browser or computer, then none of the security mechanisms are really secure, but the smartcard is a little more secure than either certificates or passwords, since it is external to a user’s computer and can only be abused for a limited time window when a user enters a code from the smartcard.
19.10.1 Zero sum approximation

In order to apply the minimax theorem, we renormalize the matrix elements so that the sum of attack and defence is not 100 but zero, thus we subtract 50 from all the values, giving

\[
\Pi_D = -\Pi_A \begin{pmatrix}
50 & 50 & -10 & -20 \\
20 & 30 & 40 & -40 \\
0 & 20 & -40 & -40
\end{pmatrix}.
\]

(19.32)

We begin by looking for a saddle-point in the game.

\[
\max \min \Pi_D = \max \left( \begin{pmatrix}
-20 \\
-40 \\
-40
\end{pmatrix} \right) = -20.
\]

(19.33)

\[
\min \max \Pi_D = \min(50, 50, 40, -20) = -20.
\]

(19.34)

The matrix yields a single saddle-point and the value of the game as \(-20\), with the optimal strategies being smartcards versus Trojan horses. This is an surprising result that is not clearly obvious from the original payoff table. First of all, the value of the is a negative number, which means that the result is in the attacker’s favour. This might not have been expected by looking at the original estimates of security, but the conclusion is the stable equilibrium of the two opposing sides, and therefore reveals the ‘rational’ conclusion in the data.

We might however consider this to be a unfair conclusion, since the likelihood that an attacker will be able to insert a Trojan horse into a user’s computer is quite low and might be detected by anti-virus software. We could therefore delete the Trojan strategy and recompute the equilibrium. Thus, the game is still zero-sum, but we now have

\[
\Pi_D' = -\Pi_A' \begin{pmatrix}
50 & 50 & -10 \\
20 & 30 & 40 \\
0 & 20 & -40
\end{pmatrix}.
\]

(19.35)

We now find that there is no saddle point equilibrium in the matrix, thus there is no optimal pure strategy contest here. This means that the solution must be in terms of a mixture of strategies and is therefore rather harder to solve. We begin by trying to eliminate any obviously weak strategies from the mixture, since these cannot lead to any optimal behaviour. Examining the matrix in eqn. (19.35), we see that rows \(i = 1, i = 2\) both strictly dominate row \(i = 3\), hence \(i = 3\) is a weak strategy for the defending row player “α”; we can delete it, giving:

\[
\Pi_D'' = -\Pi_A'' \begin{pmatrix}
50 & 50 & -10 \\
20 & 30 & 40
\end{pmatrix}.
\]

(19.36)
There is still no saddle point, and there are no more cases of strict dominance. We must now find 
a way of solving the correct linear combinations. There are many ways one might proceed, but this 
early quite simple. The value of the payoff for the defending row player is
\[ v = \bar{\alpha}^T \Pi^w \bar{\beta}, \]  
where \( \bar{\alpha} \) is the vector of strategies for the defence that now lies between a convex mixture of the 
smartcard strategy and the certificate strategy, and \( \bar{\beta} \) is a convex mixture of the attack strategies, all 
of which are active at present:
\[
\bar{\alpha}^T = (\alpha, 1 - \alpha), \quad \bar{\beta} = \begin{pmatrix} \beta_1 \\ \beta_2 \\ \beta_3 \end{pmatrix},
\]
where \( \beta_1 + \beta_2 + \beta_3 = 1 \) and \( \alpha > 0, \beta_i > 0 \). We can try to maximize the payoff for the 
defending player, by examining the rates of change and looking for any stationary values. This is 
not a solution method, but it does provide an indication of how to choose the values of the free 
parameters \( (\alpha, \beta_1, \beta_2, \beta_3) \). The partial derivatives are:
\[
\frac{\partial v}{\partial \beta_1} = 30\alpha + 20 \\
\frac{\partial v}{\partial \beta_2} = 20\alpha + 30 \\
\frac{\partial v}{\partial \beta_3} = -50\alpha + 40 \\
\frac{\partial v}{\partial \alpha} = 30\beta_1 + 20\beta_2 - 50\beta_3 = 0.
\]  
We set the last derivative to zero, since we are looking for a mixture of \( \bar{\beta} \) that makes the payoff 
stationary for the players. Thus, whatever the attacker would like to play, this condition will limit 
the success of the attack, by virtue of the defending player’s optimal responses. We notice first that 
the derivative with respect to \( \beta_3 \) has a possibility of going negative. This means that the \( \beta \)-attack 
player will try to play this strategy to make the defence players payoff \( v \) less. Using the fact that 
the \( \beta_i \) sum to one, we have
\[
\frac{\partial v}{\partial \alpha} = 30\beta_1 + 20\beta_2 - 50(1 - \beta_1 - \beta_2) = 0 \\
= 80\beta_1 + 70\beta_2 - 50 = 0.
\]

Now let us examine some cases of this:

1. Suppose the attack player decides to avoid session hijack and play \( \beta_2 \), since it looks like a 
marginally weak strategy (it is a weakly dominant column of the defence player’s payoff –
and this is a zero sum game), then we have:

$$\beta_1 = \frac{5}{8}, \beta_3 = \frac{3}{8}. \quad (19.41)$$

If we choose this mixture of strategies for $\vec{\beta}$ in eqn. (19.37), then the value of the game is

$$v = \frac{1}{8}(\alpha, 1 - \alpha) \begin{pmatrix} 220 \\ 220 \end{pmatrix}. \quad (19.42)$$

i.e. the payoff to the defence player is $v = 27.5$ regardless of what mixture of strategies $\alpha$ is chosen. Thus both smartcards and certificates seem to be equally valid defences to this attack. This conclusion is not particularly intuitive.

2. What if we choose $\beta_1 = 0$? Then we have

$$\frac{\partial v}{\partial \alpha} = 70\beta_2 - 50 = 0. \quad (19.43)$$

i.e.

$$\beta_1 = \frac{5}{7}, \beta_3 = \frac{2}{7}. \quad (19.44)$$

and the value of the game for the defence player is

$$v = \frac{1}{7}(\alpha, 1 - \alpha) \begin{pmatrix} 230 \\ 230 \end{pmatrix}. \quad (19.45)$$

Again, surprisingly, the choice of $\alpha$ is irrelevant to the outcome. However, crucially, the value of the game, $v = 230/7$, is now greater for the defence player. This is to the defence player’s linking, but alas he is not calling the shots and choosing $\vec{\beta}$. The attacker would not choose this strategy, since he is trying to minimize the defence player’s payoff. Remarkably, in both these cases, the defence player is a ‘sitting duck’, with no way of improving his defence by choosing one strategy or another, in any given mixture.

We have not proved that case 1 above is the optimal strategy, beyond doubt, but with a little work it is not difficult to see that it is indeed the solution to the game. Indeed, we can see from eqn. (19.40), the value of $80\beta_1 + 70\beta_2$ can never be greater than when $\beta_2 = 0$, thus the value of the denominator in the fraction cannot be made larger and the value of the game cannot be made smaller by choice of $\beta_i$.

What is interesting about this example is that the conclusion is not at all obvious from the original security level evaluations. This analytical procedure selects the limits of the tug-of-war contest in an impartial way.
19.10.2 Non-zero Sum Approximation

The zero sum approximation does not allow us to take into account other sources of loss and gain. What is lost to the defender is not necessarily gained by the attacker. The cost of implementing a technological solution should be factored into the calculations in considering "cost of ownership". It is not gratis to implement a certificate system, for example. We can add in costs of this kind by modifying the payoff:

\[ \text{Payoff} = \text{Security level} - \text{cost of strategy}. \]

What should the exact formula be for this cost? This depends on our estimation of the relative importance of these. We need to relate the currencies to one another, using the same scale (see section 11.3).

Let us define for clarity \( \alpha \) to be the defending player and \( \beta \) to be the attacking player. There is no rational way to relate security level to cost of implementation, so we must define this relationship as a matter of policy. For the defender:

\[ \Pi_1 = \Pi_1^{(0)} - k_1 C_\alpha, \quad (19.46) \]

where \( \Pi_1^{(0)} \) is the basic constant sum estimation of payoff, \( k_1 \) is a policy constant, and \( C_\alpha \) is the cost of investing in the security technology. Similarly, for the attacker:

\[ \Pi_2 = \Pi_2^{(0)} - k_2 C_\beta, \quad (19.47) \]

where \( \Pi_2^{(0)} \) is the attackers basic payoff, \( k_2 \) is the attackers own estimate of how gain relates to invested time \( C_\beta \) in carrying out the attack.

The addition of the cost of strategy term is a perturbation to the basic payoff. One can use the solution for the Nash equilibrium of the resulting game to test how much of a perturbation must be added to the constant-sum conclusion, before the conclusion about optimal strategies is altered.

**Add cost to defender** \( \sigma_1 = \alpha \)

We now try to combine the information about the different strategies to modify the payoff, as above. What is the cost of implementing the security technologies?

Smart cards cost money and need to be replaced sometimes, so there is an expense; however we can make customers pay for these, so there is no effect on the payoffs for the bank. A certificate system, on the other hand, is costly since it must be set up and maintained, depending on the local web services. Also, cryptographic certificates do not work consistently\(^4\) in all browsers, so there is much programming, debugging and maintenance to keep the system working. Passwords are available to everyone with no investment, so these are also unaffected.

\(^4\)This scenario is based around an Internet Bank known to the author.
We shall thus suppose, as a combination of judgement and policy, that $k_1 C = -20$ for the certificate strategy, and zero for the others. Accordingly -20 is subtracted from the second row of the payoff table, for the defender:

<table>
<thead>
<tr>
<th>Security</th>
<th>Steal credentials</th>
<th>Hijack session</th>
<th>Spoof site</th>
<th>Trojan in browser</th>
</tr>
</thead>
<tbody>
<tr>
<td>Smartcard</td>
<td>(90,10)</td>
<td>(90,10)</td>
<td>(40,60)</td>
<td>(30,70)</td>
</tr>
<tr>
<td>Certificate</td>
<td>(50,30)</td>
<td>(60,20)</td>
<td>(50,30)</td>
<td>(-10,90)</td>
</tr>
<tr>
<td>Password</td>
<td>(50,50)</td>
<td>(70,30)</td>
<td>(10,90)</td>
<td>(10,90)</td>
</tr>
</tbody>
</table>

Now that the payoffs are not constant sum, different solution methods are required than those used for the zero sum case. The solution of non-constant sum games is beyond the scope of this book, however the open source Gambit software package [Gam] is a useful tool for solving for Nash equilibria. Feeding these data into the computer software, we obtain an answer that is, in fact, the same as for the zero-sum case: the pure strategy equilibrium is $(\alpha_1, \beta_4)$, i.e. use of smartcards for the defender and Trojan horse for the attacker.

**ADD COST TO ATTACKER $\sigma_2 = \beta$**

Now we consider the cost of engagement from the attacker’s viewpoint. The bank is not the only one in the game who needs to invest to use its available strategies. The average attacker is poorly inclined to invest a huge effort in preparing an attack of the system, thus he judges that the cost $-k_2 C \beta$ of developing the Trojan horse strategy to be $-40$. This is a relatively high price, but then the attacker is somewhat lazy and judges the effort to be more than his time is worth. The value is thus subtracted from the final column. Similarly, he judges the cost of the hijack strategy to be $-10$, on the payoff scale. This is less expensive to him than the Trojan horse, because there are tools already available on the Internet to help him. The resulting table is now like this:

<table>
<thead>
<tr>
<th>Security</th>
<th>Steal credentials</th>
<th>Hijack session</th>
<th>Spoof site</th>
<th>Trojan in browser</th>
</tr>
</thead>
<tbody>
<tr>
<td>Smartcard</td>
<td>(90,10)</td>
<td>(90,0)</td>
<td>(40,60)</td>
<td>(30,30)</td>
</tr>
<tr>
<td>Certificate</td>
<td>(50,30)</td>
<td>(60,10)</td>
<td>(50,30)</td>
<td>(-10,50)</td>
</tr>
<tr>
<td>Password</td>
<td>(50,50)</td>
<td>(70,20)</td>
<td>(10,90)</td>
<td>(10,50)</td>
</tr>
</tbody>
</table>

After these alterations, there is no pure strategy equilibrium. Instead, there is now a mixed strategy equilibrium, $(\frac{1}{2} \alpha_1 + \frac{1}{2} \alpha_2, \frac{1}{5} \beta_3 + \frac{4}{5} \beta_4)$, with the defender mixing 50-50 between smartcards and certificates, i.e. and the attacker mixing $\frac{1}{5} \beta_3$ (site spoofing) and $\frac{4}{5} \beta_4$ (Trojan horse). The analysis mixes these as probabilities, but suppose we decide that a one in five chance of using a Trojan makes it worth disregarding this attack strategy altogether, then the game is solved by a pure strategy equilibrium of certificates versus site-spoofing.
CONCLUSIONS

This example, although somewhat contrived, tells us the relative stability of the conclusions drawn from placing value on different aspects of the system solution. It is not necessary to have precise information about the different payoffs in order to make a reasonably informed decision about the optimal strategies. Why? In this case the reason is that the payoffs are quite stable to small perturbations. If we use our best guesses in order to find a suitable stable equilibrium, we can then test each assumption by saying: what if I perturb the value by a small amount? Is the equilibrium robust under this change? If it is, we have a good idea what conclusion the model predicts. On the other hand, if a small change leads to a quite different solution, then the onus is on the system analyst to find an accurate payoff model, or to find additional strategies that can result in a more stable conclusion.

19.11 THE GARBAGE COLLECTION GAME

The difficult aspect of game theoretical modelling is turning the high level concepts and aims listed above, into precise numerical values. This is particularly true when the values that govern a game change over time.

To illustrate a possible solution to this problem we consider an example of some importance, namely the clearing of garbage files from user disks (see fig. 19.4). The need for user garbage collection (called tidying) has been argued by several authors (see [Zwi89, Bur95, BR97]), but users do not like having even the most useless of files deleted from their home areas.

We shall model this game as a zero sum game. The currency of this game must first be agreed upon. What value will be transferred from one player to the other in play? There are three relevant measurements to take into account: (i) the amount of resources consumed by the attacker (or freed by the defender); sociological rewards: (ii) ‘goodwill’ or (iii) ‘privilege’ which are conferred as a result of sticking to the policy rules. These latter rewards can most easily be combined into an effective variable ‘satisfaction’. A ‘satisfaction’ measure is needed in order to set limits on individuals’ rewards for cheating, or balance the situation in which the system administrator prevents users from using any resources at all. This is clearly not a defensible use of the system, thus the system defences should be penalized for restricting users too much. The characteristic matrix now has two contributions,

\[ \pi = \pi_r(\text{resources}) + \pi_s(\text{satisfaction}). \]  

(19.48)

It is convenient to define

\[ \pi_r \equiv \pi(\text{resources}) = \frac{1}{2} \left( \frac{\text{Resources won}}{\text{Total resources}} \right). \]  

(19.49)
Figure 19.4: Payoff matrix and a fault tree showing how the fault tree feeds into the game as probabilities, and vice versa. The values in the matrix are probabilistic expressions expressing the likelihood of achieving each strategic goal, weighted by a currency scale for its relative importance. See [Bur00c] for details of this game.

Satisfaction \( \pi_s \) is assigned arbitrarily on a scale from plus to minus one half, such that,

\[
-\frac{1}{2} \leq \pi_s \leq +\frac{1}{2} \\
-\frac{1}{2} \leq \pi_r \leq +\frac{1}{2} \\
-1 \leq \pi \leq +1.
\]  

(19.50)

The pay-off is related to the movements made through the lattice \( \vec{d} \). The different strategies can now be regarded as duels, or games of timing.

<table>
<thead>
<tr>
<th>Users/System</th>
<th>Ask to tidy</th>
<th>Tidy by date</th>
<th>Tidy above Threshold</th>
<th>Quotas</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tidy when asked</td>
<td>( \pi(1,1) )</td>
<td>( \pi(1,2) )</td>
<td>( \pi(1,3) )</td>
<td>( \pi(1,4) )</td>
</tr>
<tr>
<td>Never tidy</td>
<td>( \pi(2,1) )</td>
<td>( \pi(2,2) )</td>
<td>( \pi(2,3) )</td>
<td>( \pi(2,4) )</td>
</tr>
<tr>
<td>Conceal files</td>
<td>( \pi(3,1) )</td>
<td>( \pi(3,2) )</td>
<td>( \pi(3,3) )</td>
<td>( \pi(3,4) )</td>
</tr>
<tr>
<td>Change timestamps</td>
<td>( \pi(4,1) )</td>
<td>( \pi(4,2) )</td>
<td>( \pi(4,3) )</td>
<td>( \pi(4,4) )</td>
</tr>
</tbody>
</table>

These elements of the characteristic matrix must now be filled, using a model and a policy. A
general expression for the rate at which users produce files is approximated by:

\[ r_u = \frac{n_b r_b + n_g r_g}{n_b + n_g}, \tag{19.51} \]

where \( r_b \) is the rate at which bad users (i.e. problem users) produce files, and \( r_g \) is the rate for good users. The total number of users \( n_u = n_b + n_g \). From experience, the ratio \( n_b/n_g \) is about one percent. The rate can be expressed as a scaled number between zero and one, for convenience, so that \( r_b = 1 - r_g \).

The payoff in terms of the consumption of resources by users, to the users themselves, can then be modelled as a gradually accumulation of files, in daily waves, which are a maximum around midday:

\[ \pi_u = \frac{1}{2} \int_0^T dt \frac{r_u (\sin(2\pi t/24) + 1)}{R_{tot}}, \tag{19.52} \]

where the factor of 24 is the human daily rhythm, measured in hours, and \( R_{tot} \) is the total amount of resources to be consumed. Note that, by considering only good user or bad users, one has a corresponding expression for \( \pi_g \) and \( \pi_b \), with \( r_u \) replaced by \( r_g \) or \( r_b \) respectively. An automatic garbage collection system (cfengine) results in a negative pay-off to users, i.e. a pay-off to the system administrator. This may be written

\[ \pi_a = -\frac{1}{2} \int_0^T dt \frac{r_a (\sin(2\pi t/T_p) + 1)}{R_{tot}}, \tag{19.53} \]

where \( T_p \) is the period of execution for the automatic system. This is typically hourly or more often, so the frequency of the automatic cycle is some twenty times greater than that of the human cycle. The rate of resource-freeing \( r_a \) is also greater than \( r_u \), since file deletion takes little time compared to file creation, and also an automated system will be faster than a human. The quota payoff yields a fixed allocation of resources, which are assumed to be distributed equally amongst users and thus each quota slice assumed to be unavailable to other users. The users are nonchalant, so \( \pi_s = 0 \) here, but the quota yields

\[ \pi_q = +\frac{1}{2} \left( \frac{1}{n_b + n_g} \right). \tag{19.54} \]

The matrix elements are expressed in terms of these.

\( \pi(1, 1) \): Here \( \pi_s = -\frac{1}{2} \) since the system administrator is as satisfied as possible by the users’ behaviour. \( \pi_r \) is the rate of file creation by good users \( \pi_g \), i.e. only legal files are produced. Comparing the strategies, it is clear that \( \pi(1, 1) = \pi(1, 2) = \pi(1, 3) \).

\( \pi(1, 4) \): Here \( \pi_s = 0 \) reflecting the users’ dissatisfaction with the quotas, but the system administrator is penalized for restricting the freedom of the users. With fixed quotas, users cannot generate
large temporary files. \( \pi \) is the fixed quota payoff, a fair slice of the resources. Clearly 
\( \pi(4, 1) = \pi(4, 2) = \pi(4, 3) = \pi(4, 4) \). The game has a fixed value if this strategy is 
adopted by system administrators. However, it does not mean that this is the best strategy, 
according to the rules of the game, since the system administrator loses points for restrictive 
practices, which are not in the best interest of the organization. This is yet to be determined.

\[ \pi(2, 1): \] Here \( \pi_s = \frac{1}{2} \) since the system administrator is maximally dissatisfied with users’ refusal to 
tidy their files. The pay-off for users is also maximal in taking control of resources, since the 
system administrator does nothing to prevent this, thus \( \pi_r = \pi_u \). Examining the strategies, 
one find that \( \pi(2, 1) = \pi(3, 1) = \pi(3, 2) = \pi(3, 3) = \pi(4, 1) = \pi(4, 2) \).

\[ \pi(2, 2): \] Here \( \pi_s = \frac{1}{2} \) since the system administrator is maximally dissatisfied with users’ refusal to 
tidy their files. The pay-off for users is now mitigated by the action of the automatic system 
which works in competition, thus \( \pi_r = \pi_u - \pi_a \). The automatic system is invalidated by 
user bluffing (file concealment).

\[ \pi(2, 3): \] Here \( \pi_s = \frac{1}{2} \) since the system administrator is maximally dissatisfied with users’ refusal 
to tidy their files. The pay-off for users is mitigated by the automatic system, but this does 
not activate until some threshold time is reached, i.e. until \( t > t_0 \). Since changing the date 
cannot conceal files from the automatic system, when they are tidied above threshold, we 
have \( \pi(2, 3) = \pi(4, 3) \).

Thus, in summary, the characteristic matrix is given by:

\[
\pi(u, s) = \begin{pmatrix}
\frac{1}{2} + \pi_a(t) & \frac{1}{2} + \pi_u(t) + \pi_a(t) & \frac{1}{2} + \pi_u(t) + \pi_a(t) \theta(t_0 - t) & \pi_q \\
\frac{1}{2} + \pi_u(t) & \frac{1}{2} + \pi_u(t) + \pi_a(t) & \frac{1}{2} + \pi_u(t) + \pi_a(t) \theta(t_0 - t) & \pi_q \\
\frac{1}{2} + \pi_u(t) & \frac{1}{2} + \pi_u(t) & \frac{1}{2} + \pi_u(t) + \pi_a(t) \theta(t_0 - t) & \pi_q \\
\frac{1}{2} + \pi_u(t) & \frac{1}{2} + \pi_u(t) & \frac{1}{2} + \pi_u(t) & \pi_q
\end{pmatrix},
\]

(19.55)

where the step function is defined by,

\[
\theta(t_0 - t) = \begin{cases} 
1 & (t \geq t_0) \\
0 & (t < t_0) 
\end{cases},
\]

(19.56)

and represents the time-delay in starting the automatic tidying system in the case of tidy-above-
threshold. This was explained in more detail in [Bur03].

It is possible to say several things about the relative sizes of these contributions. The automatic 
system works at least as fast as any human so, by design, in this simple model we have

\[
\frac{1}{2} \geq |\pi_a| \geq |\pi_u| \geq |\pi_g| \geq 0,
\]

(19.57)
Figure 19.5: The absolute values of pay-off contributions as a function of time (in hours). For daily tidying $T_p = 24$. User numbers are set in the ratio $(n_g, n_b) = (99, 1)$, based on rough ratios from the author’s College environment, i.e. one percent of users are considered mischievous. The filling rates are in the same ratio: $r_b/R_{tot} = 0.99$, $r_g/R_{tot} = 0.01$, $r_a/R_{tot} = 0.1$. The flat dot-slashed line is $|\pi_q|$, the quota pay-off. The lower wavy line is the cumulative pay-off resulting from good users, while the upper line represents the pay-off from bad users. The upper line doubles as the magnitude of the pay-off $|\pi_a| \geq |\pi_u|$, if we apply the restriction that an automatic system can never win back more than users have already taken. Without this restriction, $|\pi_a|$ would be steeper.
for all times. For short times $\pi_q > \pi_u$, but users can quickly fill their quota and overtake this. In a zero-sum game, the automatic system can never tidy garbage faster than users can create it, so the first inequality is always saturated. From the nature of the cumulative pay-offs, we can also say that

$$\left(\frac{1}{2} + \pi_u\right) \geq \left(\frac{1}{2} + \pi_u + \pi_a (t_0 - t)\right) \geq \left(\frac{1}{2} + \pi_u + \pi_a\right),$$

(19.58)

and

$$\left|\frac{1}{2} + \pi_u\right| \geq \left|\pi_g - \frac{1}{2}\right|.$$

(19.59)

Applying these results to a modest strategy of automatic tidying, of garbage, referring to figure 19.5, one sees that the automatic system can always match users’ moves. As drawn, the daily ripples of the automatic system are in phase with the users’ activity. This is not realistic, since tidying would normally be done at night when user activity is low, however such details need not concern us in this illustrative example.

The policy created in setting up the rules of play for the game, penalizes the system administrator for employing strict quotas which restrict their activities. Even so, users do not gain much from this, because quotas are constant for all time. A quota is a severe handicap to users in the game, except for very short times before users reach their quota limits. Quotas could be considered cheating by the system administrator, since they determine the final outcome even before play commences. There is no longer an adaptive allocation of resources. Users cannot create temporary files which exceed these hard and fast quotas. An immunity-type model which allows fluctuations is a more resource efficient strategy in this respect, since it allows users to span all the available resources for short periods of time, without consuming them for ever.

According to the minimax theorem, if we have

$$\max \min \pi_{rc} = \min \max \pi_{rc},$$

(19.60)

it implies the existence of a pair of single, pure strategies $(r^*, c^*)$ that are optimal for both players, regardless of what the other does. If the equality is not satisfied, then the minimax theorem tells us that there exist optimal mixtures of strategies, where each player selects at random from a number of pure strategies with a certain probability weight.

The situation for our time-dependent example matrix is different for small $t$ and for large $t$. The distinction depends on whether users have had time to exceed fixed quotas or not; thus ‘small $t$’ refers to times when users are not impeded by the imposition of quotas. For small $t$, one has:

$$\max \min \pi_{rc} = \max \left(\begin{array}{c}
\pi_g - \frac{1}{2} \\
\frac{1}{2} + \pi_u + \pi_a \\
\frac{1}{2} + \pi_u
\end{array}\right).$$

$$= \frac{1}{2} + \pi_u.$$

(19.61)
The ordering of sizes in the above minimum vector is:

\[
\frac{1}{2} + \pi_u \geq \frac{1}{2} + \pi_u + \pi_a \theta(t_0 - t) \geq \pi_u + \pi_a \theta(t_0 - t) \geq \pi_g - \frac{1}{2}.
\]  

(19.62)

For the opponent’s endeavours one has

\[
\min \rightarrow \max \downarrow \pi_{rc} = \min \rightarrow \max \downarrow \frac{1}{2} + \frac{1}{2} + \frac{1}{2} + \frac{1}{2} + \pi_u, \pi_q
\]

\[
= \frac{1}{2} + \pi_u.
\]  

(19.63)

This indicates that the equality in eqn. (19.60) is satisfied and there exists at least one pair of pure strategies which is optimal for both players. In this case, the pair is for users to conceal files, regardless of how the system administrator tidies files (the system administrator’s strategies all contribute the same weight in eqn (19.63). Thus for small times, the users are always winning the game if one assumes that they are allowed to bluff by concealment. If the possibility of concealment or bluffing is removed (perhaps through an improved technology), then the next best strategy is for users to bluff by changing the date, assuming that the tidying looks at the date. In that case, the best system administrator strategy is to tidy indiscriminately at threshold.

For large times (when system resources are becoming or have become scarce), then the situation looks different. In this case one finds that

\[
\max \downarrow \min \rightarrow \pi_{rc} = \min \rightarrow \max \downarrow \pi_{rc} = \pi_q.
\]  

(19.64)

In other words, the quota solution determines the outcome of the game for any user strategy. As already commented, this might be considered cheating or poor use of resources, at the very least. If one eliminates quotas from the game, then the results for small times hold also at large times.

19.12 A SOCIAL ENGINEERING GAME

The extensive form of a game is the form in which all possible moves are documented. The extensive form is more often associated with \( N \)-person game theory (where \( N > 2 \)) than with simple 2-person games, since the extensive game tree of a two person game is often trivial, unless the moves are repeatable. A general introduction to \( N \)-person game theory and the extensive form is well beyond the scope of this book. However, we can examine some of the ideas through examples, as they contains some important insights.

So far, in decision making, we have ignored the causality of strategies and actions taken by the agents within a human-computer system. The extensive form brings us back to this issue and rounds off the topic of decision making by bringing together the concepts of information, causality and utility into a unified framework. Let us consider a simple example of causal decision making, with three participants.
The following example is a special case of a general decision game. A generic 3-person game tree for binary decision-making is shown in fig. 19.6. The yes-no decision can be interpreted in a number of ways to apply this to different scenarios.

Consider the following scenario: a company or other enterprise offers a training programme to its employees so as to instruct them in policy. One of the aims of this training is to prevent attacks of the company by social engineers. Whether or not employees are trained, some of them will choose to obey company policy while others will not. This decision can be based, or not, on the information from the training process. A potential attacker of the company can observe the employees and have knowledge of the decisions made by management and by the individual employees (he might be an insider). How shall the employer, employees and attacker make their decisions?

In order to make rational decisions, there has to be a payoff to each player. We shall consider first a game of perfect information, in which each player can see all of the moves and decisions throughout the game, in the order in which they occur. Note that the primary obstacle to understanding games in extensive form is in finding a suitable notation for describing the possible strategy combinations. On the one hand, a more complex notation can provide greater clarity, on the other hand it can also overwhelm.

\[
\begin{array}{c|c|c}
\text{Player} & \text{Yes move} & \text{No move} \\
\hline
\text{P1 = Employer} & \text{Train personnel} & \text{Don’t train personnel} \\
\text{P2 = Employees} & \text{Obey policy} & \text{Don’t obey policy} \\
\text{P3 = Attacker} & \text{Attack enterprise} & \text{Don’t attack enterprise} \\
\end{array}
\]

An alternative interpretation might be to imagine that the attacker is simply a bad case of ‘gremlins’, i.e. chance, and to consider the worst case scenario for the enterprise’s training policy given that chance error plays its ‘worst’ hand.

A first step in solving this game is to change it into strategic form. One might suppose that the game tree as shown in fig. 19.6 catalogues all possible strategies, but this is not the case. The tree of moves is not the same as the tree of decisions about play of an entire game. A game has two trees: a tree of moves (the game tree) and a tree of strategies that is not normally drawn. The strategy tree determines a complete set of contingencies for every player in every situation, and is often drawn as a table or payoff matrix; however, the tree form preserves some of the causal structure in the game.

Any player’s pure strategy must specify a course of action in each and every contingency that can occur throughout the game, given the information that is available to each player when he or she commences play. After all, it is not known at the outset of the game what will actually transpire between the players. A pure strategy must therefore correspond to a complete play of the game, by the player concerned, with all reasoning implicit. This means that, even if the response of a player is just one move, a strategy must specify alternative moves concomitant with the actual state.
CHAPTER 19. DECISION AND STRATEGY

Figure 19.6: The extensive form of a 3-person game with binary decision-making.

The states of the binary decision game are described by the information strings using the binary symbols Y and N (for yes and no), or equivalently operations $\hat{O}_1$ and $\hat{O}_2$. Each player $\alpha = 1, 2, 3$ can send only one symbol or perform one operation, thus all game plays consist of possible sequences of these two symbols: YYY, YNY, YNN, YYN, NYY, NNY, NYN, and NNN. However, this is not the same as the number of decisions, given that each player makes a move based on past information. Each decision has one of two forms: either the player decides on a move (operation), given that the game has arrived in a particular state,

$$S_\alpha(\hat{O}_1 \text{else } \hat{O}_2 \mid \text{state } = Q), \tag{19.65}$$

or the player decides to make a move, ignoring the information about past history:

$$S_\alpha(\hat{O}_1), \tag{19.66}$$
$$S_\alpha(\hat{O}_2). \tag{19.67}$$

Thus, as the state space grows exponentially, so the number of decisions grows exponentially. For player 1 (the employer) there is only one state — the starting state, so the conditional moves make no sense, or one can say that they are not independent of the unconditional moves. Using $Y, N$ notation for simplicity, these are:

$$S_1(Y), S_1(N) \tag{19.68}$$
Player 2 inherits two possible states from player 1 and thus can choose between

\[ S_2(Y, N \mid Q = Y), \]
\[ S_2(N, Y \mid Q = Y), \]
\[ S_2(Y, N \mid Q = N), \]
\[ S_2(N, Y \mid Q = N), \]
\[ S_2(Y, Y \mid Q = Y), \]
\[ S_2(N, N \mid Q = N). \] (19.69)

The first four strategies can be summarized as two: i.e. do the same as player 1 or do the opposite of player 1 (Employer). The latter two strategies are: do Y or N regardless of what player 1 does, e.g. do Y else do Y implies ‘do anyway’.

Player 3 inherits four possible states from player 2 and thus can choose all of the four choices at each node.

\[ S_3(Y, N \mid Q = YY), \]
\[ S_3(N, Y \mid Q = YY), \]
\[ S_3(Y, Y \mid Q = YY), \]
\[ S_3(N, N \mid Q = YY) \]
\[ S_3(Y, N \mid Q = YN), \]
\[ S_3(N, Y \mid Q = YN), \]
\[ S_3(Y, Y \mid Q = YN), \]
\[ S_3(N, N \mid Q = YN) \]
\[ S_3(Y, N \mid Q = NY), \]
\[ S_3(N, Y \mid Q = NY), \]
\[ S_3(Y, Y \mid Q = NY), \]
\[ S_3(N, N \mid Q = NY) \]
\[ S_3(Y, N \mid Q = NN), \]
\[ S_3(N, Y \mid Q = NN), \]
\[ S_3(Y, Y \mid Q = NN), \]
\[ S_3(N, N \mid Q = NN). \] (19.70)

Notice that there must be an ‘else’ alternative at each branch: i.e. each strategy is of the form, “if the state of the system is Q do X else do Y”. If, for whatever reason, a player formally chooses a
strategy based on an irrelevant state, we must be able to evaluate the game nevertheless⁵. What is the point of this? An intelligent player would hardly choose a strategy corresponding to a given state unless the system were in that state — but what if the player is somehow prevented from knowing the state, or suffers a lapse of judgement? This is where one strays into the realms of imperfect information: we shall not go down that path, but mention only the possibility in passing. Completeness requires us to catalogue all possible pathways in a game.

If we imagine that each path through the tree is a sequence of operations $\hat{O}_3\hat{O}_2\hat{O}_1$, then paths through the tree are represented, with dependencies explicit as

$$(3|21)(2|1)(1),$$

where each parenthesis is an operator, and we borrow the notation of conditional probability to make dependencies explicit. We must now transfer the payoffs from the tree of moves to the table of payoffs (now a three dimensional table). Let us suppose hypothetically that an employer can earn 100 credits from a business in total. It costs 10 credits to train the staff in the company policy and procedures. Moreover, staff wages amount to a constant 10 credits, regardless of profits. If staff are found to be not complying with policy, they are docked half their earnings, and thus they receive only 5 credits. An attacker is likely to corrupt an employee into cooperating with an attack if the employee does not follow policy, and might earn 20 credits from this, but must pay 10 to the corrupt employee who was the ‘insider’ to compensate for losses in wages. Clearly this is rather simplistic, but serves to illustrate a point.

<table>
<thead>
<tr>
<th>Path</th>
<th>Employer</th>
<th>Employee</th>
<th>Attacker</th>
<th>Payoff vector</th>
</tr>
</thead>
<tbody>
<tr>
<td>NNN</td>
<td>100-5</td>
<td>5</td>
<td>0</td>
<td>(95,5,0)</td>
</tr>
<tr>
<td>NNY</td>
<td>100-5-20</td>
<td>5+10</td>
<td>20-10</td>
<td>(75,15,10)</td>
</tr>
<tr>
<td>NYN</td>
<td>100-10</td>
<td>10</td>
<td>0</td>
<td>(90,10,0)</td>
</tr>
<tr>
<td>NYY</td>
<td>100-10-20</td>
<td>10</td>
<td>20</td>
<td>(70,10,20)</td>
</tr>
<tr>
<td>YNN</td>
<td>90-5</td>
<td>5</td>
<td>0</td>
<td>(85,5,0)</td>
</tr>
<tr>
<td>YNY</td>
<td>90-5-20</td>
<td>5+10</td>
<td>20-10</td>
<td>(65,15,10)</td>
</tr>
<tr>
<td>YYN</td>
<td>90-10</td>
<td>10</td>
<td>0</td>
<td>(80,10,0)</td>
</tr>
<tr>
<td>YYY</td>
<td>90-10-20</td>
<td>10</td>
<td>20</td>
<td>(60,10,20)</td>
</tr>
</tbody>
</table>

These are the payoffs to the various users. It is not clear from the table exactly what strategy is best for any of the players pursuing selfish interest. Thus, a method that can tell us the best rational choice in terms of the payoff currency is of great interest, especially if it tells us the effect

⁵Note that the simple form “If (Q)... else...” is possible here due to the binary nature of the decisions. The tree grows very complex if there are more than two choices. In a game with three or more decisions at each node, there must be sufficient state information in the decision tree to distinguish a unique strategy path, in every decision, otherwise choices must be made ad hoc, at random.
of changing the relative payoffs between users. Could an employer maximize likely profits by paying employees a little more, faced with possible attack?

The game presented is an almost constant sum game, except that the constant is different in the two choices of player 1 (the employer). If the employer trains the employees then total profit is only 90 rather than 100, since this costs him the outlay of training\(^6\).

The payoff matrix is three dimensional, so we must be split into two slices for \( S_1 = Y \):

\[
\begin{pmatrix}
S_3 / S_2 & (Y, Y | Y Y) & (Y, N | Y Y) & (N, Y | Y Y) & (N, N | Y Y) \\
(Y, Y | Y Y) & Y Y Y & Y Y Y & Y N Y & Y N Y \\
(Y, N | Y Y) & Y Y Y & Y Y Y & Y N N & Y N N \\
(N, Y | Y Y) & Y Y N & Y Y N & Y N Y & Y N Y \\
(N, N | Y Y) & Y Y N & Y Y N & Y N N & Y N N \\
(Y, Y | Y N) & Y Y Y & Y Y Y & Y N Y & Y N Y \\
(Y, N | Y N) & Y Y Y & Y Y Y & Y N N & Y N N \\
(N, Y | Y N) & Y Y N & Y Y N & Y N N & Y N N \\
(N, N | Y N) & Y Y N & Y Y N & Y N N & Y N N \\
(Y, Y | N N) & Y Y Y & Y Y Y & Y N Y & Y N Y \\
(Y, N | N N) & Y Y N & Y Y N & Y N N & Y N N \\
(N, Y | N N) & Y Y Y & Y Y Y & Y N Y & Y N Y \\
(N, N | N N) & Y Y N & Y Y N & Y N N & Y N N
\end{pmatrix}
\]

\[\Pi_{S_1 = Y, S_2, S_3} = \] (19.72)

and \( S_1 = N \):

---

\(^6\)This could be handled by introducing a fourth player to whom we pay this value, but that would only serve to complicate matters here, since the fourth player plays no strategic role in the security situation being analyzed here.
The outcomes of this game can be calculated (see [Gam]), with the result that there are many possible equilibrium strategies. The payoffs to the players are all pure strategy equilibria and are all quite similar, since there the pure strategies of the extensive game are not independent, in spite of the great number of distinguishable combinations. The one that is best for the employer assigns payoffs \((75, 15, 10)\) to the players. This comes from no staff training, no policy conformance and attack by an attacker. This is not a bad outcome perhaps, but the employer is losing ten units of profit to the attacker and has disloyal staff. This is not desirable.

Perhaps of greater concern, the employer is powerless to decide the outcome of the game in the equilibria. All of the solutions assume that no staff training is performed; thus it is the actions of the other players who determine the employer’s payoff. There is no point to the employer in training staff, because they can get more by corrupt means. This is not a desirable situation, so we consider how to modify the rules of the game to achieve a more desirable result, and restore some of the control the employer has over destiny. Suppose, for instance, the employer doubles the wages of the employees and maintains the policy of halving of wages for failure to comply with company training.
We recompute the Nash equilibria with these values. The best new equilibrium is in fact worse than before for the employer. Again, there are several equilibria; the outcomes for the players are all close to (60,20,20) with equilibrium strategies: no training, break policy and attack respectively for the players. Clearly bribing the staff does not necessarily help. What else might we do? What we have not accounted for is the possible effect of training on deflecting the attacker. Suppose instead of paying employees more, the employer seeks more effective training that halves the gain of the attacker. This needn’t cost any more — the employer simply finds a competent staff trainer. Now we have:

<table>
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</tr>
</tbody>
</table>

With these payoffs we have, again, many equilibria; however, amongst the equilibria is now the possibility of the equilibrium payoffs (70,10,10) to the players using moves Train staff, Staff obey policy, Attacker attack. Thus, while the employer does not increase total profit, he can reduce losses to attackers.

We can continue reevaluating how much resources to assign to each part of the system and indeed our own value systems (what is truly important to us), to see how far we must go before the balance of the game tips over to a cost-effective result. The main point to be derived from this toy model is that by changing value allocations in a game, one can explore the rational consequences, even when they are somewhat convoluted and counter-intuitive.
19.13 **Human Elements of Policy Decision**

Policy compliance by humans is a thorny issue, since humans do not always behave rationally. Game theory suggests that humans require a payoff in order to tip their judgement in favour of compliance, on the scales of rational judgement. While game theory always assumes that humans will pursue purely selfish interest, this is clearly not necessarily true (though more true than many of us would care to imagine), so the game once again gives us mainly an insight into the worst case scenario.

Humans are also multifaceted souls. The payoff to a human might be include a number of things. Sometimes thank are enough, other times monetary reward is craved. Choosing policy over humans is fraught with the irrationality of the possible response: humans require training or conditioning to accept policy. Whether one calls this training or brainwashing is a subject for an ethical debate, very interesting but somewhat outside the scope of this chapter. Choosing a policy that is not understood by humans can lead to actions that are ‘criminal’ according to policy. Peer review of policy prevents policies that are overtly contrary to group wishes, but a group’s wishes are not always rational or representative either. For humans to accept policy, they must often have the feeling of freedom to decide their own destiny, even if this is through a democratic process in which they actually have little influence. Responsibility within a system confers privilege; this can be used as a payoff to motivate humans; it could also be withdrawn as a punishment, though disgruntled humans tend to sabotage systems, or become criminals within its policies.

19.14 **CODA: Extensive versus Strategic Configuration Management**

Let us now put together many of the themes of this book to consider the entire development of a human-computer system in terms of a game for attaining an ideal state (see [Bur98b]). If it is a game of perfect information, it must be deterministic and the size of the information sets must grow exponentially with each move. There is thus, in the limiting case, an infinite number of pure strategies required to attain the desired configuration in every possible situation. We must ask, however, whether or not the state we acquire is desirable and by what criteria. The view taken in this book is that stability is a key component of this criterion.

The path dependent and path independent approaches can be described using the game theoretical language of this chapter. They also have their analogues in thermodynamics of equilibrium and non-equilibrium systems. Asking whether or not complete path is required is the same as asking whether or not configuration must be a game of perfect information in order to “win”. We have not properly discussed the matter of games of imperfect information here; this is a large and subtle topic which would require inappropriately long discussion for this book (see [Ras01]), so we shall
mention only the briefest sketch.

In a game of imperfect information, every decision in the game tree is made without knowing the whole history of the tree. All a player sees is the information in a coarse-grained form. The nodes of the game tree are grouped into sets called information sets that conceal the details of past history to the player. Information sets blur the distinction between individual nodes, making them indistinguishable to the player, by grouping together nodes in the game. Rasmusen calls these information sets ‘clouds’, since a player can tell that the game has reached an approximate location covered by a cloud, but cannot tell the exact node within the cloud.

Example 210. In the configuration management approach used by the software cfengine, only the current state of the system is known, not the path by which this was achieved. The distinct routes by which the state occurred are hidden, and thus cfengine operates a game of imperfect information.

Can a player win based on only imperfect information? This depends on whether there is a path to a node within the winning game equilibrium from the current location. The key to this is to disallow deep trees. If every final state is attainable from every intermediate state, then this condition is satisfied. If we use only commuting operators (see section 15.3) this will be true, since there is no impediment to implementing a string of operations that makes a move in the game. If we use path dependent operators, then the property of ‘can get there from here’ is spoiled unless we can steer the exact path throughout the game. If the something from the system environment should alter the state of the system, so that it is no longer in the equilibrium solution, it must be corrected by maintenance (see chapter 16), but now we are no longer guaranteed to be able to find a path back to where we were unless the operations are commutative. We can summarize this as follows.

**Principle 10** (Regulation with imperfect information). Only a system with finite state information (that functions with imperfect information) like a Markov process can be regulated indefinitely with a finite policy. Open systems must forget their history in order to satisfy this criterion.

We can also turn this argument around to make it sound less like a limitation and more like an advantage. A system that achieves equilibrium cannot depend on the route it took to reach equilibrium. A steady state, i.e. one that persists as the end-point of a process, (see section 9.10) is one in which the system has reached a stable dynamical equilibrium. That means that the system desires to change itself by amount zero in every time step once it has reached that state. Since zero does not depend on anything, this cannot depend on the path. Such a state is a property of a location, not a path. Consider the following analogy.

Example 211. Imagine a ball rolling into a valley: once it reaches the bottom of the valley, it stays there regardless of the path it took — assuming only that once such path exists. The bottom of the valley is the most stable state and we would normally choose this as policy. Now, suppose we decide to place the ball on a ledge and make this our policy. Now the ball might not be stable to
perturbations and if, by some perturbation, it falls out of that state, there is only a limited number of paths that will take it back there. If there is no single point of ‘bottomness’ in the valley (perhaps the whole base of the valley is at the same height), then these final states are an equivalence class and there is no need to distinguish them.

The example above tells us that maintaining a truly stable state requires no particular information about the path. However, if we ask to maintain an unstable state, precise information is required.

The reason why the extensive, path-dependent approach sometimes seems desirable in configuration management is that we seldom bother to construct the entire tree of actions or moves explicitly, or analyze the dependencies to determine their final importance. Moreover, if an extensive approach is to be coded as a policy then we must find the pre-determined strategies that describe the tree (as in eqns. 19.68, 19.69 and 19.70); these are even more numerous.

Traugott has argued that complete path information is vital to maintain the correct ‘congruent’ state of configuration of a host ([Tra02]). In Traugott’s proposal, one specifies a path always starting from a known base state so that the configuration follows the same path. The agrees with the view above, but is only distinct or special if the system is to be configured in a non-stable state, i.e. a state which does not have maximum stability. The viewpoint we must take here is that a sufficient description of policy must guarantee a best stable state and that any path to this state is good enough. Once the system arrives at the state, the path it took becomes irrelevant.

With the expert operator approach (see section 15.3), we provide a set of primitives that, while not providing the actual strategy set, is guaranteed to cover the possibility space for any policy (as orthogonal vectors span a vector space), thus provide a path to any final state. Choosing a policy then constrains the space of all possible policies along each orthogonal operations axis and reductions can be made rationally. It might seem like a lot of work to build up a configuration in this way, but once it is done, it is guaranteed to be stable to decision fluctuations.

19.15 VERDICT ON GAME THEORY

This chapter has gone to some lengths to attempt to apply simple game theoretic methods to plausible examples. It is worth now returning to the question posed at the start of the chapter: are such rational decision methods justified or beneficial.

The amount of preparation that goes into the formulation a problem game theoretically seems to weigh against it as a practical method. Although it might be possible to build software that automates the procedure for a limited domain in particular circumstances, it seems likely that other heuristic methods might well find more favour amongst system designers. If systems have the prerequisite stability to support reliable function, then they should not be strongly sensitive to the
precise optimization of system parameters. One would try to design for a certain resilience or insensitivity to such choices.

The formulation of games is facilitated by an understanding of the basic promises in a system[BB14]: this is the semantic connection between the interactions of agents and the formulation of a utility function to guide them. However, we would seem to be a long way from being able to use rational decision theory as one of the basic tools for systems engineers. The level of effort and mathematical methodology is simply too high, and we should focus our attention on fundamental stability.

Applications and Further Study 19.

- Creating decision models that relate dependencies to their payoff in a rational way.
- Understanding the mechanics of decision making.
- Investigating how changes in assumptions can lead to changes in the rational outcome of the decision procedure (hence provide a formal procedure for testing one’s assumptions).
- Evaluating the stability of policy conclusions to errors of assumption, by changing utility estimates until the rational outcome becomes significantly altered.
...to see the general in the particular and the eternal in the transitory
– Alfred North Whitehead

System administration or engineering has been presented in this book as a rational endeavour, whose aim is to provide checks and balances to the design, construction and running of human-computer systems. These checks and balances can be investigated and described by traditional scientific inquiry. Some specific tools and techniques have been presented here to guide and inspire their usage in future work.

In this book, we have assumed that no system can be isolated from its environment—that opening a system for input means opening it to unpredictability. The task of the theoretician is then to look for a formal language by which to describe the observable phenomena of interacting humans and computers, including all of the uncertainties and fluctuations. For this we need sets, graphs and functions of time and space (addresses), as well as statistical methods and the control of empiricism.

The approach taken here has been to allow the predictable and unpredictable meet, by mapping the ideal of maintenance onto the idea of error correction of strings of digital operations (as in the communications theory of Shannon). The challenge lies in defining complex, multi-dependent processes as discrete operators that work predictably in noisy environments; if one can do that, classical error correction methods automatically apply.

Key notions of stability, predictability, resource management, connectivity and flow then allow one to define the concept of a set of reasonable policies, or stable regions of behaviour for the system. Policies are alternative configurations that reflect system goals. By defining policy to be a point of stable operation for a system, one ensures that the system will not immediately come undone as soon as it is allowed to run. However, there might be several policies that can be
sustained, so there is a freedom that remains.

The remaining freedom can be partially eliminated by rational means; it requires a value judgement, and here the language of games and decision theory can be used. Once this has been allowed to play its role, we are left with a far smaller number of alternatives, that is sharpened by the decision constraints. This might still not select a unique policy; the remaining choices for policy are thus not rationally distinguishable. Human preference, or random choice, thus enters for the final selection of one or more of these policies; we refer to the remaining choice as a strategy.

Bringing the scientific tradition to a new field of research is not a trivial matter. The thickness of this brief, introductory volume is a testament to the effort required to make a precise, rational statements that can be challenged by experiment. Even so, this book is only a beginning: it suggests a platform or springboard in the form of a number of theorems and paradigms. For example,

- The maintenance theorem tells us what kinds of systems can have a probably stable policy.
- Convergence, closure and orthogonality tell us how systems can reach a stable fixed point of operation through a stochastic process, hence in a noisy environment.
- Shannon’s theorem shows us that error correction is possible in maintainable systems.
- Queueing and fault network theorems and centrality measures indicate where and problems and faults are likely to occur, as well as how resources should be deployed most efficiently.

Some readers will find the liberal mixture of technology and sociology in this book disconcerting. These traditionally incompatible domains are treated here as one, with common methods. Subjective concerns are made into rational choices by formulating the utility of the choices on a measurable scale. Ultimately this allows us to employ algebraic reasoning. Utility theory, as originated by Von Neumann and Morgenstern, extends far beyond the tiny introduction offered here and should provide research theses for decades to come.

There are plenty of reasons not to be judgemental about the inadequacies of treating the human part of systems with approximate characterizations. We have survived such inadequacies in other sciences with no great injury. Moreover, it is always good to remember that all scientific models are descriptive and have underlying uncertainties.

The analogies between system administration and economics are not accidental; nor are the clear parallels between system administration and physics. All these fields share the goal of describing systems in which accountable resources are distributed, flow and interact. Physics is a game in which different forces compete for their share of the energy available in a system; we can rightfully describe the contents of this book as a physics of human-computer systems\(^1\).\footnote{This viewpoint has often caused distress when I have expressed it in the past. “This is not physics, it’s computer science” say critics, but this artificial segregation misses the point completely. Science rarely advances by splintering off sub-cultures; it usually benefits when apparently dissimilar fields are fused.}
The main difference is that physics operates according to only one set of rules (one policy). In human-computer systems, each region can have its own laws: the question then becomes not only what happens within such regions, but also what happens where these regions meet. This is a fascinating challenge that should be of interest to computer scientists and physicists alike.

Security, as an explicit viewpoint, has been avoided in this book, with good reason. Once mentioned, it tends to dominate discussions, and deflect from core issues. However, security is, in fact, addressed implicitly in a number of the topics: accessibility, vulnerability of nodes, percolation and decision theory. For more specific discussions of security, readers are directed to [Bis02].

One of the key problems in security is not technical but sociological: the sabotage of systems by disgruntled users. The Romans are reputed to have claimed that civilization is never more than three meals away from anarchy. In other words, for humans, systematic cooperation depends on the subtle bribery of the people: give them what they want and they will play by the rules of the system. If the system should fail to give them what they want, it will degenerate into a free for all.

Finally, it is worth directing a critical eye on the substance of this book. What does it actually achieve? Does it advance our ability to describe human-computer systems, or is it simply an indulgence in applying familiar forms to an insubstantial field of research? I believe that it does represent an advance. In particular:

- It invalidates certain ad hoc discussions that recur in the literature on subjective issues and suggests a rational replacement.
- It provides a number of conclusions about what makes one system better than another, with measurable certainty.
- It documents a conceptual basis so that discussions can be made uniform.
- It identifies current weaknesses and new goals for technological development.

I have always been a theorist by inclination. When I practice or experiment, it is not for love of it, but in deference to its importance. Practice brings humility and experience, but only theory brings understanding. The trouble I have found is that few academics really believe in theory, when it comes down to it. They believe that it is acceptable in books, or in the classroom, but not in the infamous real world. I know colleagues who lecture of algorithms but will not apply the principles and knowledge to organize their surroundings. I know of economists who teach theory but resort to guesswork when it comes to application in "the real world". There are mathematicians who do not believe that any non-elementary mathematics should be applied to mundane problems. When it comes to "real life", too many academics are sceptical of the validity of the knowledge they teach!

As Faust exclaims in David Luke’s superior translation:

"And I fear...

Hard studies that have cost me dear."
And now I sit, poor silly man,
No wiser than when I began.
They call me Professor and Doctor, forsooth,
For misleading many an innocent youth..."

Some have said of the endeavour to bring some formality to engineering: "It is too early in the field to be talking of theory!" For others it is too late: 'If you’d told me that before it might have changed my ways.... Never was more effort expended in justifying a viewpoint than in the defence of the irrational.

In revising the text for this reincarnated edition, I have added mainly examples of modern relevance to a cultural survey of mainly historical analytical methods. The one exception has been to try to integrate a brief introduction to the concept of promises to the text, because the latter plays a major role in volume 2. Thus, here we leave this survey of methods, as a partially successful search for enlightenment and practicality. What came out of the writing of this volume was mainly a conviction that the role of intent was not properly or expediently captured within any of the methods reviewed here, and that something needed to be done to rectify that omission. Thus began a ten year effort to develop ‘Promise Theory’ and apply it to all forms of system—a effort which continues today. Indeed, the popular attention that Promise Theory has received has been a source of constant surprise. Part of the reason for that lies in its intuitive simplicity, even without attending to its formal rigour. I leave you, the reader, to discover that for yourself, in the subsequent volume.
APPENDICES
APPENDIX A

SOME BOOLEAN FORMULAE

The probability of that event $B$ follows event $A$ is

$$P(B|A)P(A),$$

i.e. the probability that $A$ happens, multiplied by the probability that $B$ happens, given that $A$ has already happened. There is an implicit causal sequence in these probabilities, because we are assuming that $B$ depends on $A$ somehow.

If the events are independent, i.e. if the probability of $A$ and $B$ occurring is independent of the order of measurement, then it makes sense to refer to the concept of “AND”, i.e. a symmetrical operator, with no memory of what came before. In this case, the probability of both events happening is merely coincidental, and is given by the overlap product of the probabilities:

$$P(A \text{ AND } B) = P(A \cap B) = P(A)P(B).$$

(A.2)

For independent events, the inclusive or exclusive ORs are the same:

$$P(A \text{ OR } B) = P(A \cup B) = P(A) + P(B) - P(A)P(B).$$

(A.3)

These expressions can be iterated, using the symmetry of their arguments for greater numbers of inputs. If the events are not independent, then the exclusive OR is given by

$$P(A \text{ XOR } B) = P(A \oplus B) = P(A) + P(B) - 2P(A)P(B).$$

(A.4)
**A.1 CONDITIONAL PROBABILITY**

Let the set $A$ consist of subsets $A = \{a_1, a_2, \ldots\}$, some of which might overlap. If the sets do not overlap, then:

$$P(a_1) = \frac{N(a_1)}{\sum_i N(a_i)} \quad (A.5)$$

In general, we can write the set that is complementary to $a_1$ as $\overline{a}_1$, i.e. all of the elements that are not in $a_1$. Then

$$P(a_1) = \frac{N(a_1)}{N(a_1) + N(\overline{a}_1)}, \quad (A.6)$$

The conditional probability of two overlapping events is

$$P(a_1|a_2) = \frac{N(a_1 \cap a_2)}{N(a_2)} = \frac{N(a_1 \cap a_2)/N}{N(a_2)/N} = \frac{P(a_1 \cap a_2)}{P(a_2)} \quad (A.7)$$

i.e., knowledge that the search space is within $a_2$ increases the likelihood of finding the result, so the conditional probability is greater.

Now, by symmetry

$$P(a_2|a_1) = \frac{N(a_1 \cap a_2)}{N(a_1)} = \frac{N(a_1 \cap a_2)/N}{N(a_1)/N} = \frac{P(a_1 \cap a_2)}{P(a_1)} \quad (A.8)$$

thus

$$P(a_1 \cap a_2) = P(a_2|a_1)P(a_1) = P(a_1|a_2)P(a_2), \quad (A.9)$$

thus one has Bayes formula

$$P(a_2|a_1) = \frac{P(a_1|a_2)P(a_2)}{P(a_1)} \quad (A.10)$$

This is really a definition of conditional probability.

**A.2 BOOLEAN ALGEBRA AND LOGIC**

Thanks to Von Neumann et. al. our present day idea of computers is that of binary digital devices which perform Boolean logical operations. Such a device can simulate any computational process.
in principle. What remains in order to create systems which compute the results of mathematical or logical problems is the ability to combine information streams into functions which are things we want to evaluate.

Modern computers are based on the use of binary data and Boolean algebra or logic. It is straightforward to show that a simple set of linearly independent operations on bits can be used to perform simple binary arithmetic, and thus more complex calculations in combination. The commonly referred to operations in Boolean algebra are the unary (1:1) operator

<table>
<thead>
<tr>
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<tr>
<td>In</td>
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<tr>
<td>1</td>
<td>0</td>
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<tr>
<td>0</td>
<td>1</td>
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</table>

and the binary (2:1) operators

<table>
<thead>
<tr>
<th>AND</th>
<th>(\cap)</th>
</tr>
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<tbody>
<tr>
<td>In1</td>
<td>In2</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
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<tr>
<td>0</td>
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<table>
<thead>
<tr>
<th>OR</th>
<th>(\cup)</th>
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<td>In1</td>
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<table>
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<tr>
<th>XOR</th>
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<tr>
<td>In1</td>
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In digital electronics, these are simulated using multi-transistor circuit blocks.

It is easy to show that any Boolean logic operation can be constructed from the two operations \(\cap\) (AND) and \(\neg\) (NOT). This may be seen from the following identities:

\[
\begin{align*}
P \cup Q &= \neg (\neg P \cap \neg Q) \\
P \rightarrow Q &= \neg P \cap Q \\
P \leftrightarrow Q &= (P \rightarrow Q) \cap (Q \rightarrow P) \\
P \oplus Q &= \neg (P \leftrightarrow Q). \tag{A.11}
\end{align*}
\]

or, in modern programming notation

\[
\begin{align*}
P | Q &= !(!P \& !Q) \\
P \rightarrow Q &= !P \& Q \\
P == Q &= (P \rightarrow Q) \& (Q \rightarrow P) \\
P ^{\wedge} Q &= ! (P == Q).
\end{align*}
\]
or, again, in more common notation

\[
P \text{ OR } Q = \text{ NOT ( NOT } P \text{ AND NOT } Q) \\
P \rightarrow Q = \text{ NOT } P \text{ AND } Q \\
P \text{ EQUA}LS \ Q = (P \rightarrow Q) \text{ AND } (Q \rightarrow P) \\
P \text{ XOR } Q = \text{ NOT (P EQUA}LS \ Q). \quad (A.12)
\]

The ‘implication’ symbol is defined by the truth table

\[
\begin{array}{ccc}
0 \rightarrow 0 &=& 1 \\
0 \rightarrow 1 &=& 1 \\
1 \rightarrow 0 &=& 0 \\
1 \rightarrow 1 &=& 1.
\end{array} \quad (A.13)
\]
Consider a stochastic dynamical variable $q(t)$, whose complete behaviour is unknown.

In the following sections we refer to a sample of data, measured by a sensor, or represented as an abstract function of time. The time span of the same is take to be from $t = 0$ to $t = T$. If the sample was measured at discrete regular time intervals, then $t$ is a subset of discrete values labelled $t = [i]$.

The average of a measured or represented function or sample, taken over all points, is denoted simply by the expectation value brackets; this is also denoted by $E(\cdot)$ in statistics literature:

$$\langle q(t) \rangle = E[q(t)].$$  \hfill (B.1)

These brackets have no subscript. Subscripts are used to denote the average taken over a limited subset of the points.

Similarly the variance over the entire sample is denoted by $\sigma^2$ with no subscript. Subscripts are used to denote the variance of a limited subset of the full sample, as defined below.

### B.1 Local Averaging Procedure

Let us define a local averaging procedure, or method of coarse-graining (see fig 3.10).

The local averaging procedure re-averages data, moving from a detailed view to a less detailed view, by grouping neighbouring data together. In practice one always deals with data which are sampled at discrete time intervals. We shall consider this case first, and then return to a continuous function approach, which is a useful approximation to the discrete case.
**APPENDIX B. STATISTICAL AND SCALING PROPERTIES OF TIME-SERIES DATA**

**DISCRETE TIME DATA**

Consider the function \( q(t) \) shown in fig. 3.10. Let the small ticks on the horizontal axis represent the true sampling of the data, and label these by \( i = 0, 1, 2, 3, \ldots, I \). These have unit spacing. Now let the large ticks, which are more coarsely spread out, be labelled by \( k = 1, 2, 3, \ldots, K \). These have spacing \( \Delta t = m \), where \( m \) is some fixed number of the smaller ticks. The relationship between the small and the larger ticks is thus:

\[
i = (k - 1)\Delta t = (k - 1)m. \tag{B.2}
\]

In other words, there are \( \Delta t = m \) small ticks for each large one. To perform a coarse-graining, we replace the function \( q(t) \) over the whole \( k \)th cell with an average value, for each non-overlapping interval \( \Delta t \). We define this average by

\[
\langle q(k) \rangle_m \equiv \frac{1}{\Delta t} \sum_{i=(k-1)\Delta t+1}^{k\Delta t} q(i). \tag{B.3}
\]

We have started with an abstract function \( q(t) \), sampled it at discrete intervals, giving \( q(i) \), and then coarse-grained the data into larger contiguous samples \( \langle q(k) \rangle_m \):

\[
q(t) \rightarrow q(i) \rightarrow \langle q(k) \rangle_m. \tag{B.4}
\]

The variance of data \( q(i) \) over the \( k \)th cell is thus

\[
\sigma^2(k) = \frac{1}{\Delta t} \sum_{i=(k-1)\Delta t+1}^{k\Delta t} (q(i) - \langle q(k) \rangle_m)^2 \tag{B.5}
\]

\[
= \langle q^2(k) \rangle_m - \langle q(k) \rangle_m^2. \tag{B.6}
\]

The mean of the entire set of samples (summed over either \( i \) or \( k \) variables) is the same:

\[
\langle q \rangle = \frac{1}{I} \sum_{i=0}^{I} q(i) = \frac{1}{K} \sum_{k=0}^{K} \langle q(k) \rangle_m, \tag{B.7}
\]

This follows from the linearity of the sums. The same is not true of the variances however.

**\( i \)-COORDINATES (SMALL TICKS)**

\[
\langle q \rangle = \frac{1}{I} \sum_{i=0}^{I} q(i) \tag{B.8}
\]

The variance:

\[
\sigma^2 = \frac{1}{I} \sum_{i=0}^{I} (q(i) - \langle q \rangle)^2
\]

\[
= \langle q^2 \rangle - \langle q \rangle^2. \tag{B.9}
\]

Recall this expression for comparison below.
**APPENDIX B. STATISTICAL AND SCALING PROPERTIES OF TIME-SERIES DATA** 387

**k-COORDINATES (LONG TICKS)**

The average is the same as for the small ticks:

\[
\langle q \rangle = \frac{1}{K} \sum_{k=0}^{K} \langle q(k) \rangle_m,
\]

\[
= \frac{1}{K} \sum_{k=0}^{K} \left( \frac{1}{\Delta t} \sum_{i=(k-1)\Delta t+1}^{k\Delta t} q(i) \right)
\]

\[
= \frac{1}{K\Delta t} \left( \frac{1}{K} \sum_{k=0}^{K} \sum_{i=(k-1)\Delta t+1}^{k\Delta t} q(i) \right)
\]

\[
= \frac{1}{I} \sum_{i=1}^{I} q(i)
\]

\[
= \langle q \rangle.
\]  \hspace{1cm} (B.10)

However, the variance is not the same:

\[
\sigma^2 = \frac{1}{K} \sum_{k=0}^{K} (\langle q(k) \rangle_m - \langle q \rangle)^2
\]

\[
\equiv \langle (\langle q(k) \rangle_m - \langle q \rangle)^2 \rangle_K
\]

\[
= \langle (\langle q(k) \rangle_m)^2 \rangle_K - 2\langle (\langle q(k) \rangle_m \langle q \rangle) \rangle_K + \langle q \rangle^2
\]

\[
= \langle (\langle q(k) \rangle_m)^2 \rangle_K - 2\langle (\langle q(k) \rangle_m \langle q \rangle) \rangle_K + \langle q \rangle^2
\]  \hspace{1cm} (B.11)

Now

\[
\langle \cdots \rangle = \langle \langle \cdots \rangle_m \rangle_K,
\]  \hspace{1cm} (B.12)

thus

\[
\sigma^2 = \langle (\langle q(k) \rangle_m)^2 \rangle_K - \langle q \rangle^2
\]

\[
\neq \sigma^2_1.
\]  \hspace{1cm} (B.13)

The two expressions thus differ by

\[
\sigma^2 - \sigma^2_K = \langle q^2 \rangle - \langle (\langle q(k) \rangle_m)^2 \rangle_K
\]

\[
= \langle (q^2)_m - (\langle q(k) \rangle_m)^2 \rangle_K
\]

\[
= \langle \sigma^2_m(k) \rangle_K,
\]  \hspace{1cm} (B.15)

which is the average variance of the coarse-grained cells.
CONTINUOUS TIME DATA

We can now perform the same procedure using continuous time. This idealization will allow us to make models using continuous functions and functional methods, such as functional integrals. Referring once again to the figure, we define a local averaging procedure by

\[
\langle q(\tilde{t}) \rangle_{\Delta t} = \frac{1}{\Delta t} \int_{\tilde{t} - \Delta t/2}^{\tilde{t} + \Delta t/2} q(t') \, dt'.
\] (B.16)

The coarse-grained variable \( \tilde{t} \) is now the more slowly varying one. It is convenient to define the parameterization

\[
\tilde{t} = (t - t'), \quad \tilde{t} = \frac{1}{2}(t + t'),
\] (B.17)

(B.18)
on any interval between points \( t \) and \( t' \). The latter is the mid-point of such a cell, and the former is the offset from that origin. The variance of the fundamental variable, over such a grain is

\[
\sigma^2(\tilde{t}) = \frac{1}{\Delta t} \int dt'(q(t') - \langle q(\tilde{t}) \rangle_{\Delta t})^2
\]

\[
= \langle q^2(\tilde{t}) \rangle_{\Delta t} - \langle q(\tilde{t}) \rangle_{\Delta t}^2.
\] (B.19)

\( t \)-COORDINATES (INFINITESIMAL TICKS)

Over a total sample, running from 0 to \( T \)

\[
\sigma^2 = \frac{1}{T} \int_0^T dt'(q(t') - \langle q(t') \rangle)^2
\]

\[
= \langle q^2 \rangle - \langle q \rangle^2.
\] (B.20)

\( \tilde{t} \)-COORDINATES (\( \Delta t \) TICKS)

Define the average over the \( \tau \) cells of width \( \Delta t \) by

\[
\langle \langle q(\tilde{t}) \rangle_{\Delta t} \rangle_{\tau} = \frac{1}{\tau} \int_0^\tau \langle q(\tilde{t}) \rangle_{\Delta t}.
\] (B.21)

Noting that \( \tau = T/\Delta t \) and \( d\tau = d\tilde{t}/\Delta t \), one confirms that

\[
\langle \langle q(\tilde{t}) \rangle_{\Delta t} \rangle_{\tau} = \langle q \rangle
\] (B.22)
i.e. that

\[
\langle \langle \cdots \rangle_{\Delta t} \rangle_{\tau} \equiv \langle \cdots \rangle.
\] (B.23)
Over a total sample, running from 0 to $T = \tau \Delta t$, the directly calculated variance is

$$\sigma^2 = \frac{1}{T} \int_0^T dt' (q(t') - \langle q(t') \rangle)^2 \equiv \langle q^2 \rangle - \langle q \rangle^2. \quad (B.24)$$

The variance of the coarse-grained variables differs, once again,

$$\sigma^2 = \frac{1}{\tau} \int_0^\tau (\langle q(\bar{t}) \rangle \Delta t - \langle q \rangle)^2 = \langle \langle q(\bar{t}) \rangle^2 \rangle \Delta t - \langle q(\bar{t}) \rangle^2. \quad (B.25)$$

The difference

$$\sigma^2 - \sigma^2_T = \langle q(\bar{t}) \rangle^2 \Delta t = \langle \sigma^2 (\bar{t}) \rangle_T. \quad (B.26)$$

which, again, is the average of the local variances.

### B.2 Scaling and Self-similarity

The scaling hypothesis, for a function $q(t)$, under a dilatation by an arbitrary constant $\alpha$, is expressed by:

$$q(\alpha t) = \Omega(\alpha) q(t). \quad (B.27)$$

In other words, the assumption is that stretching the parameterization of time $t \to \alpha t$, leads to a uniform stretching of the function $q(t)$, by a factorizable magnification $\Omega(\alpha)$. The function retains its same ‘shape’, or functional form; it is just magnified by a constant scale.

This property is clearly not true of an arbitrary function. For example, $q(t) = \sin(\omega t)$ does not satisfy the property. Our interest in such functions is connected with dynamical systems which exist and operate over a wide range of scales. Physical systems are always limited by some constraints, so this kind of scaling law is very unlikely to be true over more than a limited range of $\alpha$ values. Nevertheless, it is possible to discuss functions which, indeed, scale in this fashion, for all values of $\alpha$, as an idealization. Such functions are said to be scale invariant, dilatation invariant, or self-similar.

In addition to perfect self-similarity, or dilatation invariance of a function, physical systems sometimes exhibit other forms of self-similarity.

Dynamical invariance tells us that the equations which describe how the function $q(t)$ behaves, or is constrained, are invariant under the change of scale. This is a weaker condition, which means that the behaviour of a complete system is invariant, but that $q(t)$ itself need not be.

$$S[\Omega^{-1}(\alpha) q(\alpha t)] \to S[q(t)]. \quad (B.28)$$
Statistical invariance tells us that the average properties of a stochastic variable, or a physical system are invariant; i.e. the function need only satisfy the scaling law on average.

### B.3 Scaling of Continuous Functions

From eqn (B.27), the symmetry between $q(t)$ and $\Omega(s)$, tells us that

\[ q(x) \sim \Omega(x), \]  

(B.29)

i.e. that they must possess similar scaling properties. In fact, $q(t)$ and $\Omega(s)$ must be homogeneous functions, in order to satisfy this relationship:

\[ q(t) = t^H, \]
\[ \Omega(s) = s^H, \]  

(B.30)

for some power $H$. In other words, one has

\[ s^{-H}q(st) = q(t). \]  

(B.31)

Consider a stochastic process $q(t)$, whose average properties show invariance over a wide range of scales, compared to the limiting resolution of the data. Consider what happens if we scale the basic definition of the local averaging procedure; the procedure, starting from the basic function, is as follows:

1. The coarse graining parameterization is $t = \Delta t \cdot \bar{t} + \bar{\ell}$, i.e. $q(t) \rightarrow q(\bar{t}, \bar{\ell})$, where $\bar{t}$ is the slowly varying parameter, and $\bar{\ell}$ is the more rapidly varying parameter.

2. Average over the intervals of size $\Delta t$, by integrating or summing over $\bar{\ell}$.

3. Rescale the coarse-graining interval from $\Delta t \rightarrow s\Delta t$.

Beginning with the definition of the averaging procedure:

\[ \langle q(\bar{t}) \rangle_{\Delta t} = \frac{1}{s\Delta t} \int_{\bar{t}-\Delta t/2}^{\bar{t}+\Delta t/2} q(\bar{t}) (sd\bar{\ell}). \]  

(B.32)

one may scale the variable of integration, so that the left hand side of the equation is the same. From the assumption of the scale invariance of $q(t)$, in eqn. (B.31), one may write

\[ \langle q(\bar{t}) \rangle_{\Delta t} = \frac{1}{s\Delta t} \int_{\bar{t}-\Delta t/2}^{\bar{t}+\Delta t/2} \left[ \frac{q(s\bar{\ell})}{s^H} \right] d(s\bar{\ell}). \]  

(B.33)
We now extend the limits of the integral, without amplifying the function itself, in order to perform a further coarse-graining, incorporating $s$ times more points:

$$\langle q(t) \rangle_{\Delta t} = \frac{1}{s \Delta t} \int_{\tau - s \Delta t/2}^{\tau + s \Delta t/2} \left[ \frac{q(s \tilde{t}')}{s^H} \right] d(s \tilde{t}'). \quad (B.34)$$

By rewriting slightly, we now observe that the function has the form of an average of a new quantity, over the larger interval $s \Delta t$:

$$\langle q(t) \rangle_{\Delta t} = \frac{1}{s \Delta t} \int_{\tau - s \Delta t/2}^{\tau + s \Delta t/2} \left[ \frac{q(s \tilde{t}')}{s^H} \right] d(s \tilde{t}'). \quad (B.35)$$

$$\langle q(t) \rangle_{\Delta t} = \frac{\langle q(st) \rangle_{s \Delta t}}{s^{H-1}} \quad (B.36)$$

The same procedure can be applied to the variance, which behaves simply as the square of the average:

$$\sigma^2_{\Delta t} = \int dt dt' \langle q(t)q(t') \rangle$$

$$= \int (st \ sdt') \frac{\langle q(t)q(t') \rangle}{s^{2H}} \quad (B.37)$$

$$\sigma^2_{\Delta t} = \frac{\sigma^2_{2 \Delta t}}{s^{2H-2}}. \quad (B.38)$$
APPENDIX C

PERCOLATION CONDITIONS

C.1 RANDOM GRAPH CONDITION

We reproduce here the argument of ref. [NSW01] to derive the condition for the probable existence of a giant cluster for a uni-partite random graph with degree distribution \( p_k \), and correct it for smaller graphs.

The method of generating functions is a powerful way of encapsulating the properties of a whole graph in a single analytical expression. Let \( k \) represent the degree of each node, and \( p_k \) be the probability distribution for the occurrence of nodes of degree \( k \) within the graph. We have,

\[
\sum_k p_k = \sum_k \frac{n_k}{N} = 1,
\]

where \( n_k \) is the number of nodes of degree \( k \), and \( N \) is the total number of nodes. The generating function for this distribution is simply the polynomial, in a dummy source variable \( J \), whose \( k \)th power coefficient is \( p_k \), i.e.

\[
G(J) = \sum_{k=0}^{k_{\text{max}}} p_k J^k,
\]

so that the probability distribution is recovered by the derivatives:

\[
p_k = \frac{1}{k!} \left. \frac{d^k G(J)}{dJ^k} \right|_{J=0},
\]

and the average degree of nodes in the graph is

\[
z \equiv \langle k \rangle = \left. J \frac{d}{dJ} G(J) \right|_{J=0}.
\]

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Figure C.1: Graphical form of the first three terms in the second power of the generating function. The \( n \)th power of \( G(J) \) generates the probabilities of finding a total degree of \( k \) from a cluster of \( n \) nodes, i.e. the probability that \( n \) nodes have \( k \) outgoing edges.

Note that \( k_{\text{max}} \) is normally taken to be infinite to approximate large graphs. We can use this generating function to evaluate average (probabilistic) properties of the graph.

If we pick an arbitrary node and follow one of the edges (links) of the graph to another node, the probability of arriving at a node of degree \( k' \) is proportional to \( k' \), since a highly connected node is proportionally more likely to be arrived at than a poorly connected node (there are more ways for it to occur). Thus, in our average picture, the probability of getting to a node of degree \( k \) is

\[
P_k = \frac{k p_k}{\sum_k k p_k} = \frac{k}{\langle k \rangle} p_k. \tag{C.5}
\]

This distribution is generated by the normalized derivative of \( G(J) \), like this:

\[
G_1(J) \equiv \sum_k k p_k J^{k-1} = \frac{1}{\langle k \rangle} \frac{d}{dJ} G(J). \tag{C.6}
\]

Following ref. [NSW01] we note that, if a distribution \( p_k \) is generated by \( G(J) \), then a number of related generating functions are obtained by taking powers of \( G(J) \). Suppose that there are \( m \) independent ways of obtaining the probability \( P_{k'} \), from different, but equivalent, graph configurations \( p_k \), then the function that generates the right combinatorics for \( p_k \) is the \( m \)th power of \( G(J) \).

\[
\gamma_m(J) \equiv [G(J)]^m = \sum_k \pi_k J^k. \tag{C.7}
\]

This is easy to see when \( m = 2 \) (see fig. C.1):

\[
\gamma_2(J) = [G(J)]^2 = \left[ \sum_k p_k J^k \right]^2 = \sum_{i,j} p_i p_j J^{i+j} = p_0 p_0 J^0 + (p_0 p_1 + p_1 p_0) J^1 + (p_0 p_2 + p_1 p_1 + p_2 p_0) J^2 + \ldots
\]
If we compare the coefficients of $J^k$ in eqns. (C.7) and (C.8), the we see that

\[
\begin{align*}
\pi_0 &= p_0 p_0 \\
\pi_1 &= p_0 p_1 + p_1 p_0 \\
\pi_2 &= p_0 p_2 + p_1 p_1 + p_2 p_0 \\
\pi_k &= C(i, j, k),
\end{align*}
\]

(C.8)

where $C(p_m)$ is the sum of all combinations such that $i + j = k$.

Thus, suppose now that we wish to calculate the average number of nodes within a connected cluster, i.e. the size of the cluster. We can obtain this result by summing the nodes that, themselves, have connected neighbours. This can be achieved by using an effective generating function, of the form:

\[
W_c[J] = J \sum_k p_k [\chi_c(J)]^k.
\]

(C.9)

Here we postulate the existence of a constrained function $\chi_c(J)$, that pertains to a given cluster $c$, within the graph, and generates the distribution of degrees recursively at all connected sub-nodes of a cluster, starting from some arbitrary point. An additional power of $J$ is added here, by convention, so that the counting starts from 1. The constraint is derived using a recursive definition that sums over clusters of connected nodes. Suppose we define the normalized distribution

\[
\chi_c(J) = \sum_k c_k J^k
\]

(C.10)

\[
= \sum_k k p_k [\chi_c(J)]^k
\]

(C.11)

\[
= J G_1(\chi_c(J)).
\]

(C.12)

Equation (C.11) is a constraint equation; its right hand side is interpreted as a sum of probabilities for arriving at a node of degree $k$, from some arbitrary starting point, that has a number of $k$ nearest neighbours each with degree distributions generated by $\chi_c(J)^k$, i.e.

\[
\chi(J) \propto \sum_k \text{Probability of picking a node of degree } k \times \text{Probable ways of connecting to } k \text{ nodes from a random node}
\]

The recursive definition indicates that the same average probabilities exist at each node of the graph; only the limit of total nodes in the cluster stops the iteration. Substituting in the generic form with coefficients $c_k$ leads to an eigenvalue equation for the vector $c_k$, with a matrix of probabilities. The principal eigenvector gives the appropriate solution for the largest cluster. Remarkably, we do not need to know the solution of $\chi_c(J)$ in order to find out when the size of connected clusters becomes dangerously large for system security. Instead, the constraint can be eliminated.
Differentiation of $W_c(J)$ with respect to the source $J$ gives a quantity that is analogous to the result in eqn. (C.4), but with a new kind of average that includes both nearest neighbour degrees, next-nearest neighbour degrees, and so on:

$$\langle\langle k \rangle\rangle \equiv \frac{d}{dJ} W_c[J] = 1 + J \frac{d}{dJ} G(\chi_c(J)) = 1 + J \frac{dG(\chi_c)}{d\chi_c} \cdot \frac{d\chi_c}{dJ} \bigg|_{J=1}.$$  \hspace{1cm} (C.13)

The result is an average estimate of the size of a connected cluster. Using eqn. (C.12), we find that

$$\frac{d\chi_c}{dJ} = \left(1 - \frac{dG_1(J)}{dJ} \right)^{-1},$$  \hspace{1cm} (C.14)

thus the average size of a randomly picked cluster is

$$\langle\langle k \rangle\rangle = 1 + J \frac{dG_1(J)}{dJ} \left(1 - \frac{dG_1(J)}{dJ} \right) \bigg|_{J=1}.$$  \hspace{1cm} (C.15)

Here we note that, self-consistently $W[1] = 1$, as long as $W[J]$ has no singularities. In the general case we must define $\Gamma_c = \ln W_c$ and $\langle\langle k \rangle\rangle = d\Gamma_c/dJ$ at $J = 1$. A giant component or cluster is defined to be a cluster that is of order $N$ nodes. If such a cluster exists, then other smaller clusters of order $\log N$ might also exist[MR98]. The condition for a giant cluster is thus that the denominator in this fraction becomes small, or

$$\frac{dG_1(J)}{dJ} (1) = 1.$$  \hspace{1cm} (C.16)

Using eqn. (C.6), we find the critical point for the emergence of a giant cluster. The large-graph condition for the existence of a giant cluster (of infinite size) is simply

$$\sum k(k-2) p_k \geq 0.$$  \hspace{1cm} (C.17)

This provides a simple test that can be applied to a human-computer system, in order to estimate the possibility of complete failure via percolating damage. If we only determine the $p_k$, then we have an immediate machine-testable criterion for the possibility of a systemwide security breach. The condition is only slightly more complicated than the simple Cayley tree approximation; but (as we will see below) it tends to give more realistic answers.

### C.2 Bi-partite Form

Random bi-partite graphs are also discussed in [NSW01] and a corresponding expression is derived for giant clusters. Here we can let $p_k$ be the fraction of users with degree $k$ (ie, having access to $k$
files), and $q_k$ be the fraction of files to which $k$ users have access. Then, from Ref. [NSW01], the large-graph condition for the appearance of a giant bi-partite cluster is:

$$\sum_{jk} jk(jk - j - k) p_j q_k > 0.$$  \hfill (C.18)

This result is still relatively simple, and provides a useful guideline for avoiding the possibility of systemwide infection—in those cases where such is practical, one seeks to hold the whole system below the percolation threshold, by not satisfying the inequality in (C.18). The left hand side of (C.18) can be viewed as a weighted scalar product of the two vectors of degree distributions:

$$q^T W p = p^T W q > 0 ,$$  \hfill (C.19)

with $W_{jk} = jk(jk - j - k)$ forming a symmetric, graph-independent weighting matrix.

### C.3 Small Graph Corrections

The problem with the above expressions is clearly that they are derived under the assumption of there being a smooth differentiable structure to the average properties of the graphs. For a small graph with $N$ nodes (either uni-partite or bi-partite), the criterion for a giant cluster becomes inaccurate. Clusters do not grow to infinity, they can only grow to size $N$ at the most, hence we must be more precise and use a dimensionful scale rather than infinity as a reference point. The correction is not hard to identify; we require

$$\left. \frac{J dG(J)}{dJ} \right|_{J=1} \gg 1,$$

for the uni-partite case. This more precise percolation criterion states that, at percolation, the average size of clusters is of the same order of magnitude as the number of nodes. However for a small graph the size of a giant cluster and of below-threshold clusters [$N$ and $\log(N)$, respectively] are not that different[MR98]. The above criterion translates into:

$$\langle k \rangle^2 - \sum_k k(k - 2) p_k \gg 1.$$  \hfill (C.21)

Thus the threshold point can be taken to be as follows. The small-graph condition for widespread percolation in a uni-partite graph of order $N$ is:

$$\langle k \rangle^2 + \sum_k k(k - 2) p_k > \log(N).$$  \hfill (C.22)

This can be understood as follows. If a graph contains a giant component, it is of order $N$ and the size of the next largest component is typically $O(\log N)$; thus, according to the theory of random
graphs the margin for error in estimating a giant component is of order $\pm \log N$. In the criterion above, the criterion for a cluster that is much greater than unity is that the right hand side is greater than zero. To this we now add the magnitude of the uncertainty in order to reduce the likelihood of an incorrect conclusion.

Similarly, for the bi-partite graph, one has the small-graph condition for widespread percolation in a bi-partite graph of order $N$:

$$\langle k \rangle^2 \langle j \rangle^2 + \sum_{jk} jk(jk - j - k)p_j p_k > \log(N/2). \quad (C.23)$$

These expressions are not much more complex than the large-graph criteria. Moreover, they remain true in the limit of large $N$. Hence we expect these small-graph criteria to be the most reliable choice for testing percolation in small systems. This expectation is borne out in the examples below. In particular, we find that, since the average coordination number $\langle k \rangle$ enters into the small-graph percolation criteria, the earlier problem of ignoring isolated nodes in the uni-partite case is now largely remedied.
Define:

\[ L_\alpha(q) = \int_{-\infty}^{+\infty} \frac{dk}{2\pi} e^{-i\alpha k} L_\alpha(k). \]  

(D.1)

Then, \( L_1 \) is:

\[
L_1 = \int_{-\infty}^{+\infty} \frac{dk}{2\pi} e^{-i\alpha k - c_1 |k|} \]  

(D.2)

\[
= \int_{0}^{+\infty} \frac{dk}{2\pi} e^{-i\alpha k + c_1 k} - \int_{0}^{+\infty} \frac{dk}{2\pi} e^{-i\alpha k - c_1 k} \]  

(D.3)

\[
= \left( \frac{-1}{iq + c_1} + \frac{1}{iq - c_1} \right) [0 - \frac{1}{2\pi}] \]  

(D.4)

\[
= \frac{c/\pi}{q^2 + c_1^2}, \]  

(D.5)

which is the Lorentz-Cauchy distribution, commonly representing signal propagation. For \( L_2 \), we have:

\[
L_2(q) = \int_{-\infty}^{+\infty} \frac{dk}{2\pi} e^{-i\alpha k - c_2 k^2} \]  

(D.6)

\[
= \int_{-\infty}^{+\infty} \frac{dk}{2\pi} e^{-i(k - iq/2c_2)^2 c_2 - q^2/4c_2^2} \]  

(D.7)

\[
= \sqrt{\frac{\pi}{c_2}} \frac{e^{-q^2/4c_2^2}}{2\pi}, \]  

(D.8)

which is the Gaussian distribution, also a maximum entropy distribution. The stable distributions satisfy a central limit theorem.


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