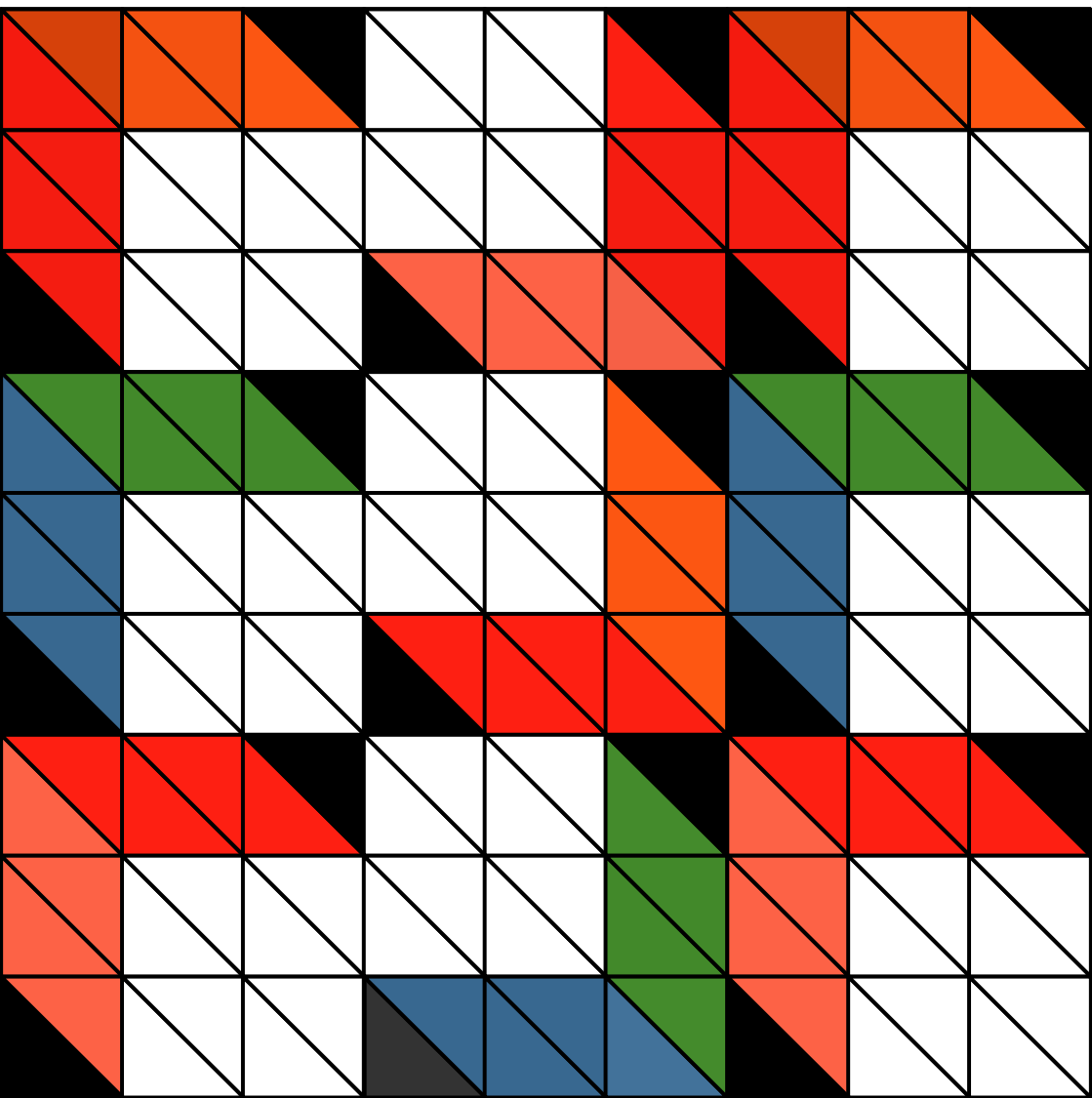
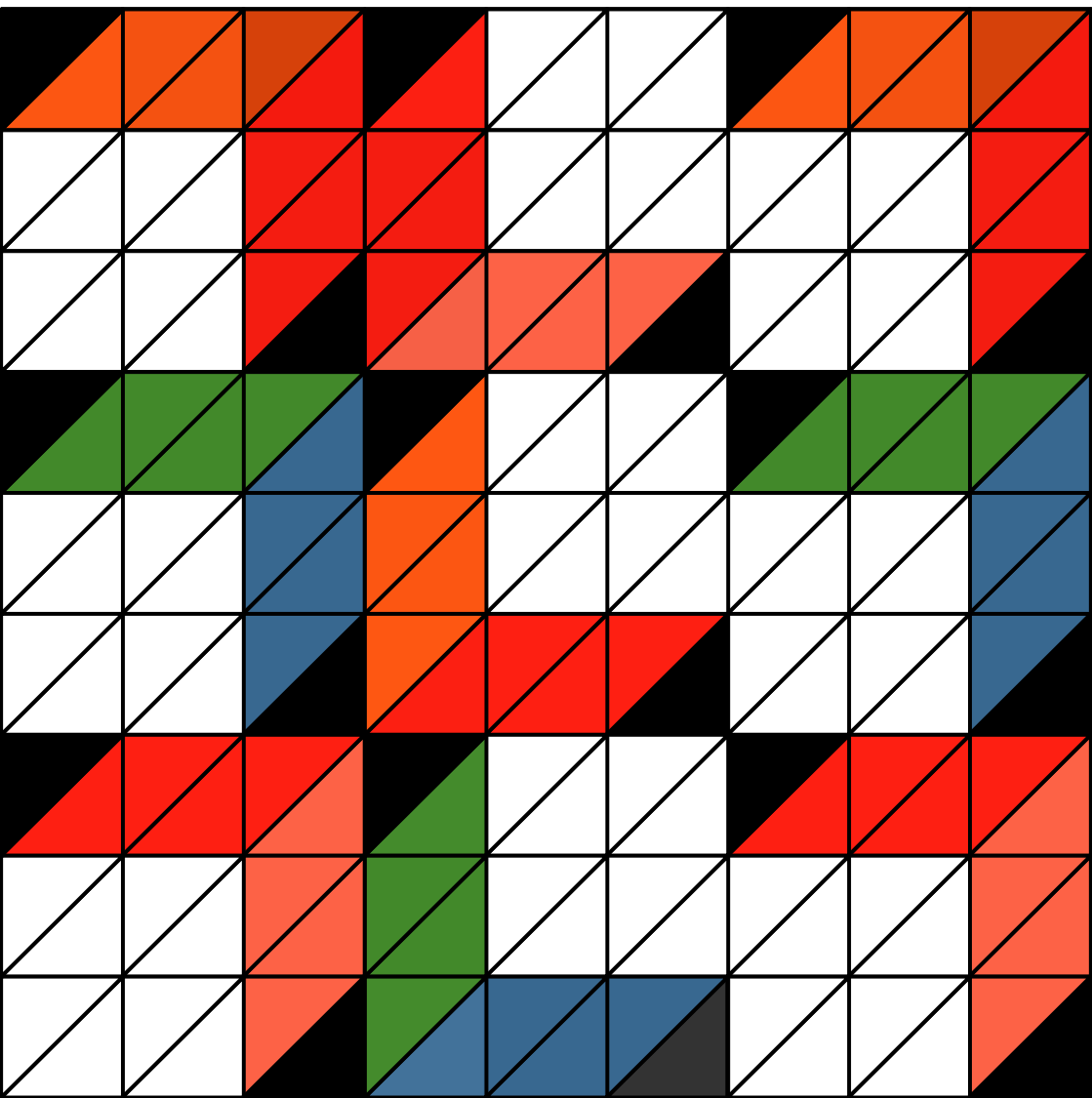


Lev Tarasov

The World Is Built on Probability

Mir Publishers Moscow





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2023

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MIR Publishers Moscow

Translated from the Russian by *Michael Burov*

First published 1988

Revised from the 1984 Russian edition

This completely digital version typeset in using \LaTeX with EB Garamond font by

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Released on the web by <http://mirtitles.org> in 2023.

Access the \LaTeX project files gitlab.com/mirtitles/twibop



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About Mir Titles Project

The Mir Titles project is an attempt to conserve the knowledge in the form of various books that came out during the Soviet era for the future generations. The collection contains books on science, mathematics, philosophy, popular science, history. The collection also has Soviet, Russian and children's literature. The project would not have been possible without the help we received from friends and contributors from across the world.

Foreword

I am very happy to release this completely electronic version of one of my favourite books. In this electronic edition all the figures have been reworked in the SVG format using Inkscape for a clearer presentation. in \LaTeX . – DAMITR

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Preface

...In nature, where chance also seems to reign, we have long ago demonstrated in each particular field the inherent necessity and regularity that asserts itself in this chance.

F. Engels

A vast concourse of events and phenomena occur in the world around us. The events are interrelated: some are effects or outcomes of others which are, in turn, the causes of still others. Gazing into this gigantic whirlpool of interrelated phenomena, we can come to two significant conclusions. One is that there are both completely determined (uniquely defined) outcomes and ambiguous outcomes. While the former can be precisely predicted, the latter can only be treated probabilistically. The second, no less essential conclusion is that ambiguous outcomes occur much more frequently than completely determined ones. Suppose you press a button and the lamp on your desk lights up. The second event (the lamp lights up) is the completely determined result of the first event (the button is pressed). Such an event is called a *completely determined* one. Take another example: a die is tossed. Each face of the die has a different number of dots. The die falls and the face with four dots ends up at the top. The second event in this case (four dots face-up) is not the completely determined outcome of the first event (the die is tossed). The top face may have contained one, two, three, five, or six dots. The event of appearance of the number of dots on the top face after a die is tossed is an example of a *random* event. These examples clearly indicate the difference between random and completely determined events.

We encounter random events (and randomness of various kinds) very often, much more frequently than is commonly thought. The choice of the winning numbers in a lottery is random. The final score of a football match is random. The number of sunny days at a given geographical location varies randomly from year to year. A set of random factors underlies the completion of any service activity: delivery ambulance

arrival, telephone connection, etc.

Maurice Glaymann and Tamas Varga have written an interesting book called *Les Probabilités à l'école* (Probability in Games and Entertainment), in which they make an interesting remark:

“When facing a chance situation, small children think that it is possible to *predict* its outcome. When they are a bit older, they believe that *nothing can be postulated*. Little by little they discover that there are patterns hiding behind the seeming chaos of the random world, and these patterns can be used to get their bearings in reality.”

There are three distinct stages here: lack of understanding of the random at first, then mere confusion, and finally a correct viewpoint. Let us forget small children for a time and try to apply this to ourselves. We shall have to recognize that frequently we stop at the first stage in a simple-minded belief that any outcome can be precisely predicted. The misconception that randomness is simply equal to chaos, or the absence of causality, has lasted a long time. And even now not everybody clearly appreciates that the abundance of random events around us conceal definite (*probabilistic*) patterns.

These ideas prompted me to write this book. I want to help the reader discover for himself the probabilistic nature of the world around us, to introduce random phenomena and processes, and to show that it is possible to orient oneself in this random world and to operate effectively within it.

This book begins with a talk between myself and an imaginary reader about the role of chance, and ends with another talk about the relationship between randomness and symmetry. The text is divided into two major parts. The first is on the concept of probability and considers the various applications of probability in practice, namely, making decisions in complicated situations, organizing queues, participating in games, optimizing the control of various processes, and doing random searches. The basic notions of cybernetics, information theory, and such comparatively new fields as operations research and the theory of games are given. The aim of the first part is to convince the reader that the random world begins directly in his own living room because, in fact, all modern life is based on probabilistic methods. The second part shows how fundamental chance is in Nature using the probabilistic laws of modern physics and biology as examples. Elements of quantum mechanics are also involved, and this allows me to demonstrate how probabilistic laws are basic to microscopic phenomena. The idea was that by passing from the first part of the book to the second one, the reader would see that probability is not only around us but is at the basis of everything.

In conclusion I would like to express my gratitude to everyone who helped me when writing this book. I.I. Gurevich, Corresponding Member of the USSR Academy of

Sciences, gave me the idea of writing this text and gave me a number of other provoking ideas concerning the material and structure of the book. B.V. Gnedenko, Member of the USSR Academy of Sciences, G.Ya. Myakishev, D.Sc. (Philosophy), and O.F. Kabardin. Cand. Sc. (Physics and Mathematics) read the manuscript thoroughly and made valuable remarks. V.A. Ezhiv and A.N. Tarasova rendered me constant advice and unstinting support the whole time I was preparing the text.

Part I

Tamed Chance

Introduction

And chance, inventor God ...

A. S. Pushkin

A Discussion on the Role of Chance

AUTHOR: “You wrote some nice words about chance in the Preface. In spite of them, I still think chance plays a negative role on the whole. Naturally, there is good luck, but everybody knows it is better not to count on it. Chance interferes with our plans, so it’s better not hang on it, we should rather ward it off as much as possible.”

AUTHOR: “That is exactly the traditional attitude towards the random. However, it is an attitude we must clearly review. First of all, is it really possible to get by without the random?”

READER: “I don’t say that it’s possible. I said we should try.”

AUTHOR: “Suppose you work at an ambulance centre. Obviously, you cannot foresee when an ambulance will be needed, where it will be necessary to send it to, and how much time the patient will require. But a practical decision depends on all of these points. How many doctors should be on duty at anyone time? On the one hand, they should not be idle waiting for calls for long periods of time, yet on the other hand, patients should not have to remain without aid for too long. You cannot avoid chance. What I am trying to say is: we cannot *eliminate* chance, and so we must *take* it *into account*.”

READER: “True, we have to make peace with chance in this example. However,

it still is a negative factor.”

AUTHOR: “Thus, we see that sometimes we have to take chance into consideration rather than control it. But we can go further. We can discover situations in which chance becomes a positive factor rather than a negative one, so that it is desirable to raise the level of the random threshold.”

READER: “I don’t understand you.”

AUTHOR: “Of course, chance occasions interfere with our plans. At the same time because it makes us new solutions and improve our ability to create”.

READER: “Do you mean an improvement is obtained by overcoming difficulties?”

AUTHOR: “The main point is that randomness can create new possibilities. An American writer has written an interesting science fiction story. A group of scientists with various disciplines is officially informed that a sensational discovery has been made, but unfortunately the discoverer died in an explosion during a demonstration of the phenomenon and thus the secret was lost. In reality neither the invention nor the inventor ever existed. The scientists were presented with the evidence of a tragedy: indistinct fragments of records, a library, and an equipped laboratory. In other words, the scientists were given a vast quantity of unconnected information with chance data from various fields of science and technology. The evidence could be called informational noise. The scientists were certain a discovery had been made, and therefore the target was achievable. They utilized all the information at their disposal and ‘revealed’ the secret of the non-existing invention. We might say that they succeeded in sifting information from the noise.”

READER: “But that’s only a science fiction story.”

AUTHOR: “True. However, the idea behind the story is far from being fiction. Any discovery is related to the use of random factors.”

READER: “I don’t think anyone can discover anything important unless he or she has a professional grasp of the subject.”

AUTHOR: “I think so too. Moreover, a discovery requires both expertise on the part of the researcher and a certain level of the development within the science as a whole. And yet ..., random factors play a fundamental role in that.”

READER: “As I understand, the word fundamental means something primary, something at the basis. Can you apply the term fundamental to something random? I admit that randomness may be useful. But can it be fundamental? In the last analysis, we deal with random variables when there is something we do not know and cannot take into account. ”

AUTHOR: “By believing that randomness is related to inadequate knowledge, you make it *subjective*. It follows that you believe that randomness appears, as it were, on the surface and that there is nothing random at the basis of phenomena. Is it correct?”

READER: “Precisely. That is why we cannot assert randomness is fundamentality. As science develops, our ability to take different factors into account increases, and the result is that the domain of random variables will gradually recede. There is sense in saying that science is the enemy of chance. ”

AUTHOR: “You’re not quite right. Indeed, the advance of science enhances our ability to make scientific predictions, that is, science is against the random factor. But at the same time, it turns out that while our scientific knowledge becomes deeper, or, more accurately, while we look at the molecular and atomic aspects of phenomena, randomness not only does not become less important, but on the contrary, it reigns supreme. Its existence proves to be independent of the degree of our knowledge. Randomness reveals its *fundamentality* at the level of the microcosm.”

READER: “This is the first time I’ve heard someone say that. Please tell me more.”

AUTHOR: “Let me say at once that this topic has had a long history. It was first formalized in Ancient Greece with two approaches to the random being stated. The two views are associated with the names of Democritus and Epicurus. Democritus identified the random with the *unknown*, believing that Nature is completely deterministic. He said: People have created an idol out of the random as a cover for their inability to think things out. Epicurus considered that the random is inherent in various phenomena, and that it is, therefore, *objective*. Democritus’s point of view was preferred for a long time, but in the 20th century, the progress of science showed that Epicurus was right. In his doctoral thesis *Difference Between the Democritian and Epicurian Philosophy on Nature* (1841), Karl Marx positively evaluated Epicurus’s view of the random and pointed out

the deep philosophical significance of the teachings of Epicurus on the *spontaneous* displacement of atoms. Of course, we should not exaggerate the contribution of Epicurus to our understanding of the random because he could only guess.”

READER: “It turns out that I presented Democritus’s views on the random without knowing it. But I would like to have some concrete examples showing the fundamentality of the random.”

AUTHOR: “Consider, for instance, a nuclear-powered submarine. How is the engine started?”

READER: “As far as I understand it, special neutron-absorbing rods are drawn from the core of the reactor. Then a controlled chain reaction involving the fission of uranium nuclei begins ...”

AUTHOR: “(interrupting) Let us try and see how everything begins.”

READER: “After entering a uranium nucleus, a neutron triggers its disintegration into two fragments and another neutron is released. The neutrons split two more uranium nuclei; four neutrons are then set free, which in turn split four more nuclei. The process develops like an avalanche.”

AUTHOR: “All right. But where does the first neutron come from?”

READER: “Who knows? Say, they come from cosmic rays.”

AUTHOR: “The submarine is deep under water. The thick layer of water protects it from cosmic rays.”

READER: “Well then, I don’t know ...”

AUTHOR: “The fact is that a uranium nucleus may either split because a neutron enters it or it may decay *spontaneously*. *The process of spontaneous nuclear fission is random.*”

READER: “But maybe spontaneous nuclear fission is caused by factors we do not know about yet.”

AUTHOR: “This is something physicists have been trying to solve. Many attempts have been made to find the hidden parameters which govern the processes in the microcosm. It has been concluded that there are no such parameters, and therefore randomness in the microcosm is fundamental. This cornerstone problem is thoroughly treated in *quantum mechanics*,

a theory which appeared in the early 20th century in connection with research on atomic processes.”

READER: “The only thing I know about quantum mechanics is that it describes the laws governing the behaviour of elementary particles.”

AUTHOR: “We shall talk about quantum mechanics in more detail later. Let me only note here that it demonstrates the fundamental role of spontaneous processes and, therefore, demonstrates the fundamentality of the random. The operation of any radiation generator, from a vacuum tube to a laser, would be impossible without spontaneous processes. They are fundamental as the trigger without which the radiation generation would not start.”

READER: “And yet, it is difficult for me to believe that randomness is fundamental. You mentioned a nuclear-powered submarine. When the captain orders that the engines be turned on, he does not rely on a lucky chance. An appropriate button is pressed, and the engines start (if they are in good condition). The same can be said when a vacuum tube is turned on. Where is the randomness here?”

AUTHOR: “Nevertheless, when we consider phenomena in the microcosm, the processes are triggered by random factors.”

READER: “However, we generally deal with processes occurring in the macrocosm.”

AUTHOR: “Firstly, while studying the world around us and trying to comprehend its *cause and effect relations*, we must address the atomic level, i. e., the level of microcosm phenomena. Secondly, the randomness in microcosmic phenomena is essentially reflected in the processes observed at the macrocosmic scale.”

READER: “Can you give me an example when the fundamentality of randomness reveals itself at the macrocosmic scale?”

AUTHOR: “*Evolution*, which is a continuous process in both the plant and animal kingdoms, may serve as an example. Evolution relies on mutation, i.e., random changes in the structure of genes. A random mutation may be rapidly spread by the reproduction of the organisms themselves. It is essential that selection occurs simultaneously with mutation. The organisms which contain the random gene are then selected to that those best

fitted to their environment survive. In consequence, evolution requires the *selection of random gene changes*.”

READER: “I don’t quite understand this business of selection.”

AUTHOR: “Here’s an example. The flowers of a certain orchid look like a female wasp. They are pollinated by male wasps which take the flowers to be females. Suppose a mutation occurs, and the shape and colour of the flower are changed. The flower will then remain unpollinated. The result is that the mutation is not passed on to the new generation. It may be said that selection rejected the mutation which changed the outward appearance of the flower. There was a species of orchid which became a self-pollinator, the flowers of this species rapidly acquired diverse shape and colour owing to the mutation.”

READER: “As far as I know, evolution progresses in the direction of the differentiation of species. Doesn’t this show that the mutations underlying evolution are not, in fact, so random?”

AUTHOR: “That argument doesn’t stand to reason. Evolution selects the fittest organisms rather than the more complex. Sometimes a higher degree of organization is preferable, but sometimes this is not the case. This is why human beings, jelly-fish, and the influenza virus can coexist in today’s world. It is essential that evolution leads to the appearance of new species that are unpredictable in principle. It may be said that *any species is unique because it occurred fundamentally* by chance.”

READER: “I have to admit that the randomness does look to be a fundamental factor indeed.”

AUTHOR: “Since we are discussing the fundamentality of randomness in the picture of evolution, let me draw your attention to one more important point. Modern science demonstrates that *chance* and *selection* are the ‘creator’.”

READER: “Just as Pushkin said, ‘And chance, inventor God ...’”

AUTHOR: “Precisely. This line is strikingly accurate.”

READER: “It appears that when speaking about chance and selection, we should imply the *selection of information from noise*, shouldn’t we? The same selection that we discussed in connection with the science-fiction story.”

AUTHOR: “Absolutely.”

READER: "I have to agree that we should consciously recognize the existence of randomness rather than try and control it."

AUTHOR: "We could say more. Naturally, the randomness which is due to the incompleteness of our knowledge is undesirable. While studying the world, man has fought, is fighting, and will continue to fight it. It should be noted at the same time that there is an *objective* randomness underlying every phenomena along with the *subjective* randomness which is due to lack of data on a phenomenon. We should also take into account the positive, creative role of the random. And in this connection it is really necessary to recognize and control randomness. Man should be able, when necessary, to create special situations, abundant with the random, and utilize the situation to his own ends."

READER: "But is it really possible to treat randomness in such a way? Isn't it like trying *to control the uncontrollable*?"

AUTHOR: "Both science and daily life indicate that it is possible to orient ourselves consciously in very random situations. Special calculation methods have been developed that depend on randomness. Special theories have been produced, such as *queueing theory*, the *theory of games*, and the *theory of random search*, to deal with it."

READER: "It is hard for me to imagine a scientific theory built on randomness."

AUTHOR: "Let me emphasize right away that randomness does not preclude scientific prediction. The fundamentality of randomness does not mean that the world around us is chaotic and devoid of order. Randomness does not imply there are no causal relations. But we shall deal with all that later. It is interesting to try and imagine a world in which randomness as an objective factor is *completely absent*."

READER: "This would be an ideally ordered world."

AUTHOR: "In such a world, the state of any object at a given time would be unambiguously determined by its past states and, in its turn, would determine the future states just as definitely. The past would be strictly connected with the present, as would the present with the future."

READER: "Anything occurring in such a world would be predetermined."

AUTHOR: "Pierre Laplace, a great French scientist of the 17th century, suggested in this connection that we imagine a superbeing who knew the past

and the future of such a world in every detail. Laplace wrote:”

‘The intellect who could know, at a given moment, every force that animates the nature and the relative positions of its every component, and would, in addition, be vast enough to analyse these data, would describe by a single formula the motions of the greatest bodies in the universe and the motions of the lightest atoms. There would be nothing uncertain for this being, and the future, like the past, be open to his gaze.’

READER: “An ideally ordered world is therefore unreal.”

AUTHOR: “As you see, it isn’t hard to feel that the real world should admit the existence of objective randomness. Now let us return to the problem of causal relations. These relations are *probabilistic* in the real world. It is only in particular cases (for example, when solving maths problems at school) that we deal with unambiguous, strictly determined relations. Here we approach one of the most essential notions of the modern science, the notion of *probability*.”

READER: “I’m familiar with it. If I throw a die, I can equally expect any number of dots from one to six. The probability of each number is the same and equal to $1/6$.”

AUTHOR: “Suppose you stand at the side of a road, with motor-cars passing by. What is the probability of the first two digits in their four digit number being equal?”

READER: “The probability equals $1/10$.”

AUTHOR: “Therefore, if you’re patient and observe enough cars, about one tenth of them will have number-plates with the same first two digits, would they? Say, about thirty cars out of 300 will have such plates. Maybe, 27 or 32, but not 10 or 100.”

READER: “I think so.”

AUTHOR: “But then there would be no need to stand at the roadside. The result could be predicted. This is an example of *probabilistic prediction*. Look at how many random factors are involved in this situation. A car could turn off the road before reaching the observer, or another car could stop or even turn back. And nonetheless, both today and tomorrow, about 30 cars out of 300 would have plates with the same first two digits.”

READER: “So, in spite of numerous random factors, the situation has a certain constancy.”

AUTHOR: “This constancy is commonly called *statistical stability*. It is essential that statistical stability is observed because of random factors rather than despite them.”

READER: “I hadn’t thought that we deal with probabilistic predictions everywhere. They include, for instance, sports predictions and weather forecasts.”

AUTHOR: “You’re absolutely right. An important point is that probabilistic (statistical) causal relations are common, while those leading to unambiguous predictions are just a special case. While definite predictions only presuppose the necessity of a phenomenon, probabilistic predictions are related simultaneously both with necessity and randomness. Thus, mutations are random, but the process of selection is governed by laws, that is, it is a necessary prerequisite.”

READER: “I see. The individual acts of the spontaneous fission of uranium nuclei are random, but the development of the chain reaction is unavoidable.”

AUTHOR: “Taken separately, any discovery is random. However, a situation which is favourable for the appearance of such a chance should exist. This chance is determined by the advance of science, the expertise of the researchers, and the level of measurement technology. A discovery is random, but the logic of the progress leading to the discovery in the long run is regular, unavoidable, and necessary.”

READER: “Now I see why the fundamentality of randomness does not result in the disorder of our world. Randomness and necessity are always combined.”

AUTHOR: “Correct. Friedrich Engels wrote in *The Origin of the Family, Private Property, and the State* (1884): ‘In Nature, where chance also seems to reign, we have long ago demonstrated in each particular field the inherent necessity and regularity that asserts itself in this chance.’ The Hungarian mathematician Á. Rényi wrote about the same thing in an interesting book *Letters on Probability*”:

‘I came across *Contemplations* by Aurelius and accidentally opened the page where he wrote about two possibilities: the world is either

in vast chaos or, otherwise, order and regularity reign supreme. And although I had read these lines many times, it was the first time that I thought over why Marcus Aurelius believed that the world should be dominated by either chance or order. Why did he believe that these two possibilities are contradictory? The world is dominated by randomness, but order and regularity operate at the same time, being shaped out of the mass of random events according to the laws of the random.'

READER: "As far as I understand, order and regularity are produced from a mass of random events, and this leads to the concept probability. "

AUTHOR: "You're absolutely right. *Individual factors* vary from case to case. At the same time, the *picture as a whole* remains stable. This stability is expressed in terms of *probability*. This is why our world proves to be flexible, dynamic, and capable of advancing."

READER: "It follows that the world around us may justly be said to be a world of probability."

AUTHOR: "It is better to speak of the *world as being built on probability*. When we examine this world, we shall concentrate on two groups of questions. Firstly, I shall show how man, owing to his use of probability in science and technology, was able to tame randomness and thus turn it from being his enemy into an ally and friend. Secondly, using the achievements of modern physics and biology, I shall demonstrate the probabilistic features of the laws of nature. In consequence, I shall show that the world around us (including both the natural and artificial world) is really built on probability."

Chapter 1

Mathematics of Randomness

This doctrine, combining the accuracy of mathematical proofs and the uncertainty of chance occasions and making peace between these seemingly contradictory elements has a full right to contend for the title of the mathematics of the random.

Blaise Pascal

Probability

Classical definition of probability. When we toss a coin, we do not know which will land face up, heads or tails. However, there is something we do know. We know that the chances of both heads and tails are equal. We also know that the chances of any of the faces of a die landing face up are equal. That the chances are equal in both examples is due to *symmetry*. Both the coin and the die are symmetrical. When two or more events have equal chances of occurring, we call them equally possible outcomes. Heads or tails are *equally possible* outcomes. Suppose we are interested in a certain result while throwing a die, for instance, a face with a number of dots exactly divisible by three. Let us call outcomes satisfying such a requirement *favourable*. There are two favourable outcomes in our example, namely, a three or a six. Now let us call outcomes *exclusive* if the appearance of one in single trial makes it impossible for the others to appear at the same trial. A die cannot land with several faces up, so they are exclusive outcomes.

We can now formulate the classical definition of probability:

The probability of an event is the ratio of the number of favourable outcomes to the total number of equally possible exclusive outcomes.

Suppose P_A is the probability of an even A , m_A is the number of favourable outcomes, and n the total number of equally possible and exclusive outcomes. According to the classical definition of probability

$$P_A = \frac{m_A}{n}. \quad (1.1)$$

If $m_A = n$, then $P_A = 1$ and the event A is a **certain** event (it always occurs in every outcome). If $m_A = 0$, then $P_A = 0$, and the event A is an **impossible** event (it never occurs). The probability of a **random** event lies between 0 and 1.

Let an event A be throwing a die and getting a number exactly divisible by three. Here $m_A = 2$ and so the probability of the event is $1/3$, because $n = 6$. Consider one more example. We have a bag with 15 identical but differently coloured balls (seven white, two green, and six red). You draw a ball at random. What is the probability of drawing a white (red or green) ball? Drawing a white ball can be regarded as an event A , drawing a red ball is an event B , and drawing a green ball is an event C . The number of favourable outcomes of drawing a ball of a certain colour equals the number of balls of this colour, i. e., $m_A = 7$, $m_B = 6$, and $m_C = 2$. Using (1.1) and given $n = 15$, we can find the probabilities:

$$P_A = \frac{m_A}{n} = \frac{7}{15}, \quad P_B = \frac{m_B}{n} = \frac{6}{15}, \quad P_C = \frac{m_C}{n} = \frac{2}{15}.$$

Addition and multiplication of probabilities. What is the probability that a randomly drawn ball will be either red or green? The number of favourable outcomes is $m_B + m_C = 6 + 2 = 8$, and therefore the probability will be

$$P_{B+C} = \frac{m_B + m_C}{n} = \frac{8}{15}.$$

We see that $P_{B+C} = P_B + P_C$. The probability of drawing either a red or a green ball is the sum of two probabilities: the probability of drawing a red ball and that of drawing a green ball. The probability of drawing a ball which is either red or green or white is the sum of three probabilities, $P_A + P_B + P_C$. It is equal to unity ($7/15 + 6/15 + 2/15 = 1$). This stands to reason because the event in question will always occur.

The rule for adding probabilities can be formulated as follows:

The probability of one event of several exclusive events occurring is the sum of the probabilities of each separate event.

Suppose that two dice are thrown. What is the probability of getting two fours at the same time? The total number of equally possible exclusive outcomes is $n = 6 \times 6 = 36$. Each one is listed in Figure 1.1, where the left figure in the parentheses is the number on one die, and the right figure is the number on the other. There is only one favourable outcome, and it is indicated in Figure 1.1 as (4,4). Hence, the probability of the event is $1/36$. This probability is the product of two probabilities: the probability of a four appearing on one die and that of a four on the other i.e.

(1,1)	(2,1)	(3,1)	(4,1)	(5,1)	(6,1)
(1,2)	(2,2)	(3,2)	(4,2)	(5,2)	(6,2)
(1,3)	(2,3)	(3,3)	(4,3)	(5,3)	(6,3)
(1,4)	(2,4)	(3,4)	(4,4)	(5,4)	(6,4)
(1,5)	(2,5)	(3,5)	(4,5)	(5,5)	(6,5)
(1,6)	(2,6)	(3,6)	(4,6)	(5,6)	(6,6)

Figure 1.1: Possible outcomes of rolling a die.

$$P_{44} = P_4 \times P_4 = \frac{1}{6} \times \frac{1}{6} = \frac{1}{36}.$$

The rule for multiplication of probabilities can be formulated as follows:

The probability of several events occurring simultaneously equals the product of the probabilities of each separate event.

By the way, it is not necessary for the events to be simultaneous. Instead of throwing two dice at the same time, we could throw a single die twice. The probability of getting two fours at the same time when two dice are thrown is the same as the probability of getting two fours when one die is thrown twice.

In many cases both rules (addition and multiplication of probabilities) are used jointly to calculate the probability of an event. Suppose we are interested in the probability P of the *same* number coming up on two dice. Since it is only essential that the numbers be equal, we can apply the rule for adding probabilities,

$$P = P_{11} + P_{22} + P_{33} + P_{44} + P_{55} + P_{66}.$$

Each of the probabilities P_{ii} is, in turn, a product $P_i \times P_i$. *Hence*

$$P = (P_1 \times P_1) + (P_2 \times P_2) + \dots + (P_6 \times P_6) = 6 \left(\frac{1}{6} \times \frac{1}{6} \right) = \frac{1}{6}.$$

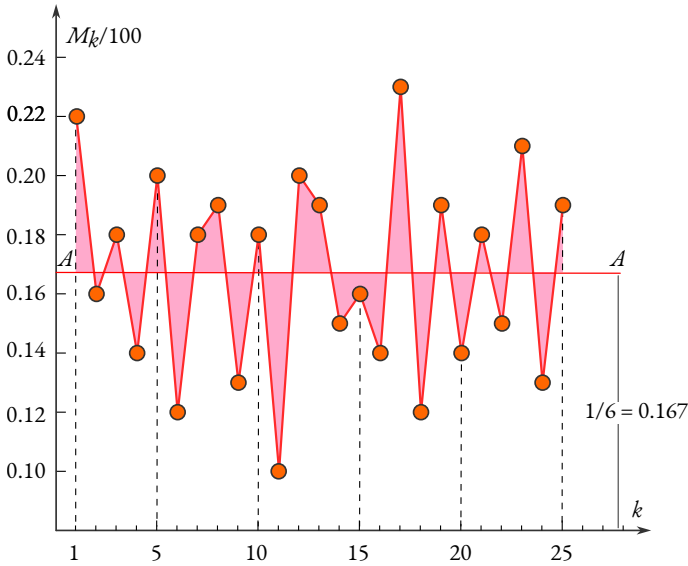


Figure 1.2: Outcomes of rolling a dice many times.

This result can be obtained right away from Figure 1.1, where the favourable outcomes are shown in the gray, (1,1), (2,2), (3,3), (4,4), (5,5), and (6,6). The total number of such outcomes is six. Consequently, $P = 6/36 = 1/6$.

Frequency and probability. The classical definition of probability and the rules for addition and multiplication of probabilities can be used to calculate the probability of a random event. However, what is the practical value of such calculations? For instance, what does it mean in practice that the probability of getting a four when a die is thrown equals $1/6$? Naturally, the assertion does not imply that a four will appear once and only once in any six trials. It is possible that it will appear once, but it is also possible that it will appear two (or more) times, or that it will not appear at all. In order to discover the probability of an event in practice we should perform a large number of trials and calculate how frequently a four appears.

Let us perform several sets of trials, for instance, throwing the die 100 times in each set. Let us designate M_1 to be the number of times a four appears in the first set, M_2 to be the number of fours in the second set, etc. The ratios $M_1/100$, $M_2/100$, $M_3/100$, ... are the frequencies with which a four appeared in each set. Having performed several sets of trials, we can see that the frequency of the appearance of a four *varies from set to*

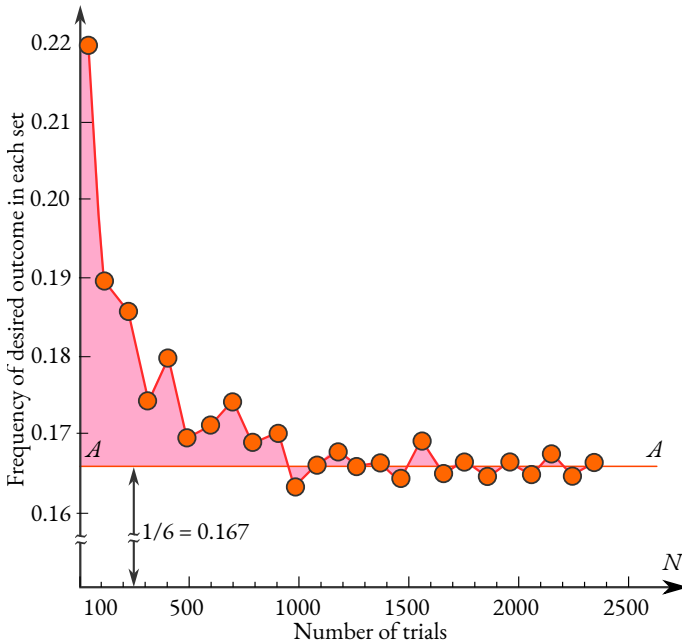


Figure 1.3: Frequencies of outcomes of trials of a die as a function of number of trials. Note how the deviation of the frequency of the occurrence of an event from its probability decreases as the number of trials increases.

set in a random fashion in the vicinity of the probability of the trials, we can see that the frequency of the appearance of a four *varies* from set to set *in a random fashion in the vicinity of the probability* of the given event, i.e. in the vicinity of $1/6$. This is clear from Figure 1.2, where the number k of sets of trials is plotted along the abscissa axis and the frequencies with which a four appears along the axis of ordinates.

Naturally, if we perform the experiment again, we will get other values of the frequencies $M_k/100$. However, the pattern of oscillations of the frequencies of the event under consideration will be stable: the deviations upwards and downwards from the straight line AA , which is associated with the probability of the event, will balance. Even though the amplitudes of the deviations will vary from set to set, they will not tend to grow or decrease. This is a consequence of the *equivalence* of each set of trials. The number of trials in each set is the same, and the results obtained in a given set do not depend on the results in any other set.

Let us make an important change in that we gradually increase the number of trials in each set. Using the results of our previous experiment, as presented in Figure 1.2, let us obtain a new result by *adding* the value of a set of trials to the result of the preceding sets. In other words, we calculate the number of fours in the first 100 trials (in our case, $M_1 = 22$), then the number of fours in the first 200 trials ($M_1 + M_2 = 22 + 16 = 38$), then in the first 300 trials ($M_1 + M_2 + M_3 = 22 + 16 + 18 = 56$) etc. We then find the frequencies of getting a four in the each new set: $M_1/100 = 0.22$, $(M_1 + M_2)/200 = 0.19$, $(M_1 + M_2 + M_3)/300 = 0.187$, etc. These frequencies are plotted in Figure 1.3 against the number of trials in each set (100, 200, ..., 2500). The figure demonstrates a crucial fact: the deviation of the frequency of the occurrence of an event from its probability decreases as the number of trials increases. In other words,

frequency of the occurrence of a random event tends to its probability with increasing number of trials.

Is it possible to give a definition of probability based on frequency? Since the frequency of the occurrence of a random event tends to its probability as the number of trials increases, we might well ask whether we can define the probability of an event as the limit of the ratio of the number of its occurrence to the number of trials as the number of trials tends to infinity. Suppose N is the number of trials and $M_A(N)$ is the number of occurrence of an event A . We want to know whether we can define the probability P_A of the event A as

$$P_A = \lim_{N \rightarrow \infty} \left[\frac{M_A(N)}{N} \right]. \quad (1.2)$$

Richard von Mises (1883-1953), a German mathematician of the early 20th century, believed that equation (1.2) could be considered a definition of the probability of a random event, and he called it the *frequency* definition of probability. Von Mises pointed out that the classical definition of probability (1.1) only “works” when there is a *finite* number of *equally* possible outcomes. For instance, situations involving the throwing of coins or dice.

However, we often encounter situations without the *symmetry* that determines whether the outcomes are equally possible. These are the cases when we cannot apply the classical definition of probability. Von Mises assumed that then the frequency definition can be used because it does not require a finite number of equally possible outcomes and, moreover, does not require any calculation of probability at all. A probability using the frequency approach is determined by experiment rather than being calculated.

However, is it possible to determine the probability of a random event in practice using (1.2)? The relationship presupposes an *infinite* number of identical trials. In practice, we must stop at a *finite* number of trials, and it is debatable what number to stop at. Should we stop after a hundred trials, or is it necessary for there to be a thousand, a million, or a hundred million? And what is the accuracy of the probability determined in such a way? There are no answers to these questions. Besides, it is not practicable to provide the same conditions while performing a very large number of trials, to say nothing of the fact that the trials may be impossible to repeat.

Consequently, relationship (1.2) is practically useless, moreover it is possible to prove (though I shall not do so) that the limit in (1.2) *does not* strictly speaking *exist*. This means that the Von Mises's error was that he made an unwarranted generalization from a correct proposition: he concluded that the probability of a random even is the limit of the frequency of its occurrence when the number of trials tends to infinity from the correct observation that the frequency of the occurrence of a random even approaches its probability as the number of trials increases.

Geometrical definition of probability. Suppose that two people have agreed to meet at a certain place between nine and ten o'clock. They also agreed that each would wait for a quarter of an hour and, if the other didn't arrive, would leave. What is the probability that they meet? Suppose x is the moment one person arrives at the appointed place, and y is the moment the other arrives. Let us consider a point with coordinates (x, y) on a plane as an outcome of the rendezvous. Every possible outcome is within the area of a square each side of which corresponds to one hour (Figure 1.4). The outcome is favourable (the two meet) for all points (x, y) such that $|x - y| \leq 1/4$. These points are within the blue part of the square in the Figure 1.4.

All the outcomes are exclusive and equally possible, and therefore the probability of the rendezvous equals the ratio of the blue area to the area of the square. This is reminiscent of the ratio of favourable outcomes to the total number of equally possible outcomes in the classical definition of probability. It should be borne in mind that this is a case where the number of outcomes (both favourable and unfavourable) is infinite. Therefore, instead of calculating the ratio of the number of favourable outcomes to the total number of outcomes, it is better to consider here the ratio of the area containing favourable outcomes to the total area of the random events.

It is not difficult to use Figure 1.4 and find the favourable area; it is the difference between the area of the whole square and the unhatched area, i.e. $1 - (3/4)^2 = 7/16 b^2$. Dividing $7/16 b^2$ by $1 b^2$, we find the probability of the rendezvous to be $7/16$.

This example illustrates the geometrical definition of probability:

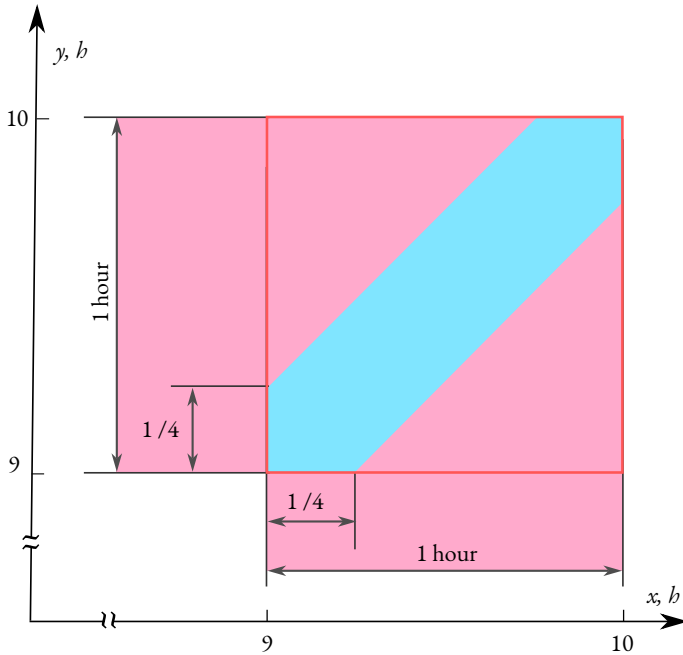


Figure 1.4: Finding the probability using the geometrical method.

The probability of a random event is the ratio of the area favourable for an event to the total area of events.

The geometrical definition of probability is a generalization of the classical definition for the case when the number of equally possible outcomes is infinite.

The development of the concept of probability. Although probabilistic notions were used by ancient Greek philosophers (such as Democritus, Epicurus, and Carus Lucretius), the theory of probability as a science began in the mid-17th century, with the work of the French scientists Blaise Pascal and Pierre Fermat and the Dutch scientist Christian Huygens. The classical definition for the probability of a random event was formulated by the Swiss mathematician Jacob Bernoulli in *Ars conjectandi* (The Art of Conjectures). The definition was given its final shape later by Pierre Laplace. The geometrical definition of probability was first applied in the 18th century. Important contributions to probability theory were made by the Russian mathematical school in

the 19th century (P.L. Chebyshev, A.A. Markov, and A.M. Lyapunov).

The extensive employment of probabilistic concepts in physics and technology demonstrated, by the early 20th century, that there was a need for a more refined definition of probability. It was necessary, in particular, in order to eliminate the reliance of probability on “common sense”. An unsuccessful attempt to give a general definition for the probability of a random event on the basis of the limit of its frequency of occurrence was made, as we have seen, by Richard von Mises. However, an *axiomatic* approach rather than a frequency one resulted in more refined definition of probability. The new approach was based on a set of certain assumptions (axioms) from which all the other propositions are deduced using clearly formulated rules.

The *axiomatic* definition of probability now generally accepted was elaborated by the Soviet mathematician A.N. Kolmogorov, Member of the USSR Academy of Sciences, in *The Basic Notions of the Probability Theory* (1936, in Russian). I shall not discuss the axiomatic definition of probability because it would require set theory. Let me only remark that Kolmogorov’s axioms gave a strict mathematical substantiation to the concept of probability and made probability theory a fully fledged mathematical discipline.

The existence of several definitions for the same notion (probability) should not worry the reader.

As L.E. Maistrov put it in *The Development of the Notion of Probability* (Nauka, Moscow, 1980):

“There are many definitions of notions, and this is an essential feature of modern science. Hence the notion of probability is no exception. Modern definitions in science represent diverse viewpoints, of which there may be very many for a fundamental notion, and each view reflects a property of the defined notion. This includes the notion of probability.”

Let me add that new definitions for a notion appear as our understanding of it becomes deeper and its properties are made clearer.

Random Numbers

Random Number Generators. Let us put ten identical balls numbered from 0 to 9 into a box. We take out a ball at random and write down its number. Suppose it is five. Then we put the ball back into the box, stir the balls well, and take out a ball at random. Suppose this time we get a one. We write it down, put the ball back into the box, stir

the balls, and take out a ball at random again. This time we get a two. Repeating this procedure many times, we obtain a disordered set of numbers, for instance: 5, 1, 2, 7, 2, 3, 0, 2, 1, 3, 9, 2, 4, 4, 1, 3, ... This sequence is disordered because each number appeared *at random*, since each time a ball was taken out at random from a well-stirred set of identical balls.

Having obtained a set of random digits, we can compile a set of random numbers. Let us consider, for instance, four-digit numbers. We need only separate our series of random numbers into groups of four digits and consider each group to be a random number: 5127, 2302, 1392, 4413, ...

Any device that yields random numbers is called a *random number generator*. There are three types of generators: *urns*, *dice*, and *roulettes* (Figure 1.5). Our box with balls is an urn.

Dice are the simplest random number generators. An example of such a generator is a cube each of whose faces is marked with a different number. Another example is a coin (or a token). Suppose five of the faces of a cube are marked with the numbers 0, 1, 2, 3, 4, while the sixth face is unmarked. Now suppose we have a token one side of which is labelled with 0 and the other with 5. Let us throw the cube and token simultaneously and add together the numbers that appear face up, the trial being discounted when the unmarked face lands face up. This generator allows us to obtain a disordered set of numbers from 0 to 9, which can then be easily used to produce sets of random numbers. A roulette is a circle marked in sectors, each of which is marked with a different number. A roulette has a rotating arrow or rolling ball. A trial involves spinning the arrow and recording the number

A roulette is a circle marked in sectors, each of which is marked with a different number. A roulette has a rotating arrow or rolling ball. A trial involves spinning the arrow and recording the number corresponding to the sector of the roulette circle within which the arrow stops.

Note that a roulette may have any number of sectors. For instance, we could divide a circle into ten sectors and label them from 0 to 9. As a random number generator, our roulette in this case is equivalent to the two generators discussed above:

- (1) an urn with ten balls and
- (2) a die and a token thrown at the same time.

A diagram of these equivalent random number generators is shown in Figure 1.5.

Tables of Random Numbers. An example of a random number table is shown in Figure 1.6. The table consists of three hundred four-digit numbers. Each digit in

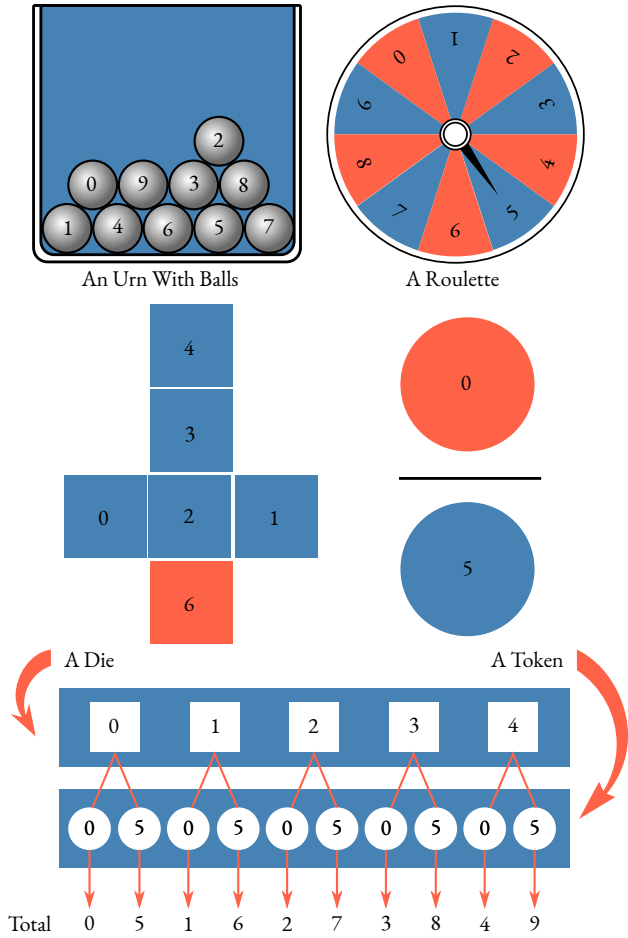


Figure 1.5: Three types of random number generators: urns, dice and roulettes.

the table was chosen randomly, as a result of a trial, e.g. throwing a die and a token. Therefore, it is understandable that there is no order in the numbers, and there is no way of predicting which digit will follow a given one. You could compile many tables after many trials. Nevertheless, there will not be even the shadow of order in the sequence of digits.

This is not amazing. A chance is a chance. But a chance has a reverse aspect. For

0655	8453	4467	3234	5320	0709	2523	9224	6271	2607
5255	5161	4889	7429	4647	4331	0010	8144	8638	0307
6314	8951	2335	0174	6993	6157	0063	6006	1736	3775
3157	9764	4862	5848	6919	3135	2837	9910	7791	8941
9052	9565	4635	0653	2254	5704	8865	2627	7959	3682
4105	4105	3187	4312	1596	9403	6859	7802	3180	4499
1437	2851	6727	5580	0368	4746	0604	7956	2304	8417
4064	4171	7013	4631	8288	4785	6560	8851	9928	2439
1037	5765	1562	9869	0756	5761	6346	5392	2986	2018
5718	8791	0754	2222	2013	0830	0927	0466	7526	6610
5127	2302	1392	4413	9651	8922	1023	6265	7877	4733
9401	2423	6301	2611	0650	0400	5998	1863	9182	9032
4064	5228	4153	2544	4125	9654	6380	6650	8567	5045
5458	1402	9849	9886	5579	4171	9844	0159	2260	1314
2461	3497	9785	5678	4471	2873	3724	8900	7852	5843
4320	4553	2545	4436	9265	6675	7989	5592	3759	3431
3466	8269	9926	7429	7516	1126	6345	4576	5059	7746
9313	7489	2464	2575	9284	1787	2391	4245	5618	0146
5179	8081	3361	0109	7730	6256	1303	6503	4081	4754
3010	5081	3300	9979	1970	6279	6307	7935	4977	0501
9599	9828	8740	6666	6692	5590	2455	3963	6463	1609
4242	3961	6247	4911	7264	0247	0583	7679	7942	2482
3585	9123	5014	6328	9659	1863	0532	6313	3199	7619
5950	3384	0276	4503	3333	8967	3382	3016	0639	2007
8462	3145	6582	8605	7300	6298	6673	6406	5951	7427
0456	0944	3058	2545	3756	2436	2408	4477	5707	5441
0672	1281	8897	5409	0653	5519	9720	0111	4745	7979
5163	9690	0413	3043	1014	0226	5460	2835	3294	3674
4995	9115	5273	1293	7894	9050	1378	2220	3756	9795
6751	6447	4991	6458	9307	3371	3243	2958	4738	3996

Figure 1.6: A table of random numbers.

instance, try and count how many times each digit occurs in Figure 1.6. You will find that digit 0 occurs 118 times (the frequency it appears is $118/1200 = 0.099$), digit 1 occurs 110 times (the frequency it appears is 0.090), digit 2 occurs 114 times (0.095), digit 3 occurs 125 times (0.104), digit 4 occurs 135 times (0.113), digit 5 occurs 135 times (0.113), digit 6 occurs 132 times (0.110), digit 7 occurs 116 times (0.097), digit 8 occurs 93 times (0.078), and digit 9 occurs 122 times (0.102). We can see that the appearance frequency for each digit is about the same, i. e. close to 0.1. Naturally, the reader has come to a conclusion that 0.1 is the *probability* that a digit appears. The reader may say that the appearance frequency of a digit is close to the probability of its appearance over a long series of trials (there are 1200 trials here).

Although this is natural, we should wonder once (again how an unordered set of *random* digits can have an *inherent stability*). This is a demonstration of the reverse aspect of chance and illustrates the determinism of *probability*.

I advise the reader to “work” a little with a random number table (see Figure 1.6). For instance, 32 numbers out of the three hundred ones in the table begin with zero, 20 begin with 1, 33 begin with 2, 33 begin with 3, 38 begin with 4, 34 begin with 5, 34 begin with 6, 24 begin with 7, 20 begin with 8, and 32 begin with 9. The probability that a number begins with a certain digit equals 0.1. It is easy to see that the results of our count are in a rather good keeping with this probability (one tenth of three hundred is thirty). However, the deviations are more noticeable than in the example considered earlier. But this is natural because the number of trials above was 1200 while here it is much less, only 300.

It is also interesting to count how many times a digit occurs in the second place (the number of hundreds), in the third place (tens), and the fourth place (units). It is easy to see that in every case the frequency with which a given digit appears is close to the probability, i.e. close to 0.1. Thus, zero occurs in the second place 25 times, in the third place 33 times, and in the fourth place 28 times.

An example with the number-plates of motor-cars randomly passing the observer was cited in the introduction. It was noted that the probability that the first two digits in the licence number were identical is 0.1. The probability that the two last digits of the number or two middle digits or the first and the last digit are identical is the same.

In order to see this, we need not observe a sequence of cars passing by. We can simply use a random number table (see Figure 1.6). The four-digit random numbers in the table can be taken as the license numbers of cars randomly passing the observer. We can see that 40 of the 300 number's have the same two first digits, 28 numbers have the same two last digits, 24 numbers have the same two middle digits, and 32 numbers have the same first and last digits. In other words, the frequencies with which a pair of identical digits appears actually varies around the probability, i.e. in the neighbourhood of 0.1.

Random Events

When we throw a die or take a ball out of an urn we deal with a *random event*. There are several interesting problems where the probability of a random event is required to be found.

A problem with coloured balls. There are three blue balls and a red ball in a box. You take two balls out of the box at random. Which is more probable: that the two balls are blue or that one is blue and one is red?

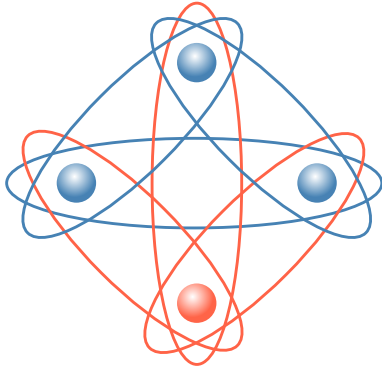


Figure 1.7: Different ways of taking out two out of three blue and one red balls.

People often answer that it is more probable that two blue balls are taken out because the number of blue balls in the box is three times greater than the number of red ones. However, the probability of taking out two blue balls is *equal* to the probability of taking out a blue and a red ball. You can see this by considering Figure 1.7. Clearly there are three ways in which two blue balls may be chosen and three ways of choosing a blue and a red ball at the same time. Therefore, the outcomes are equally probable.

We can also calculate the probability of the outcomes. The probability of taking out two blue balls equals the product of two probabilities. The first one is the probability of taking out a blue ball from a set of four balls (three blue ones plus a red one), which is $3/4$. The second probability is that of taking out a blue ball from a set of three balls (two blue ones plus a red one) which is $2/3$. Consequently, the probability of taking out two blue balls simultaneously is $3/4 \times 2/3 = 1/2$.

The probability of taking out a blue and a red ball is the sum $P_{br} + P_{rb}$, where P_{br} is the probability of taking out a blue ball from a set of four balls (three blue ones plus a red one) multiplied by the probability of taking out a red ball from a set of three balls (two blue ones plus a red one) and P_{rb} is the probability of taking out a red ball from a set of four balls (the second all in this case must then be a blue one). In other words, P_{br} is the probability of taking out a blue ball first and then a red ball while P_{rb} is the probability of taking out a red ball first and then a blue ball. Inasmuch as $P_{br} = 3/4 \times 1/3 = 1/4$ and $P_{rb} = 1/4$, the probability of taking out a pair of differently coloured balls equals $1/4 + 1/4 = 1/2$.

Throwing A Die: A Game. There are two players in this game, player *A* and player *B*. The die is thrown three times in succession during each turn. If a certain face turns up at least once during a turn (let it be a 5), player *A* scores a point. But if the five does not turn up, a point is scored by player *B*. The game is played until one of them scores, say, a hundred points. Who has the chance of winning greater? Player *A* or player *B*?

In order to answer, we first calculate the probability of player *A* scoring a point in a turn (the die is thrown three times in succession). He receives a point in any of the following three cases: if five turns up in the first trial, if five does not turn up in the first

trial but turns up in the second one, and if five does not turn up in the first two trials but turns up in the third one. Let us designate the probability of these three events as P_1 , P_2 , and P_3 , respectively. The sought probability is $P = P_1 + P_2 + P_3$. Note that the probability of five appearing when the die is thrown is $1/6$, and the probability that five does not appear is $5/6$. It is clear that $P_1 = 1/6$. To find P_2 , we should multiply the probability of the absence of a five in the first trial by the probability of its presence in the second trial, $P_2 = 5/6 \times 1/6 = 5/36$. The probability P_3 is the product of the probability of the absence of a five in two trials (the first and the second) and the probability of a five in the third trial, $P_3 = (5/6)^2 \times 1/6 = 25/216$. Consequently, $P = P_1 + P_2 + P_3 = 1/6 + 5/36 + 25/216 = 91/216$. Since $P < 1/2$, player B has more chance of winning this game. We could have reached the same conclusion in a simpler way by considering the probability of player B scoring a point after three trials. This is the probability of the absence of five in three trials: $p = 5/6 \times 5/6 \times 5/6 = 125/216$. Since $p > 1/2$, player B 's chances are better. Note that $P + p = 91/216 + 125/216 = 1$. This is natural because one of the players, A or B , must score a point in each turn.

Let us change the rules of the game a little: the die is thrown four times rather than three times in each turn. The other conditions remain the same. The probability of player B scoring a point in a turn is $5/6 \times 5/6 \times 5/6 \times 5/6 = 625/1296$. This is less than $1/2$, and therefore now A has a better chance of winning a game.

The Problem Of An Astrologer. A tyrant got angry with an astrologer and ordered his execution. However, at the last moment the tyrant made up his mind to give the astrologer a chance to save himself. He took two black and two white balls and told the astrologer to put them into two urns at random. The executioner was to choose an urn and pick a ball out of it at random. If the ball was white, the astrologer would be pardoned, and if the ball was black, he would be executed. How should the astrologer distribute the balls between the two urns in order to give himself the greatest chance of being saved?

Suppose the astrologer puts a white and a black ball into each urn (Figure 1.8 (a)). In this case, no matter which urn the executioner chooses, he will draw a white ball out of it with a probability of $1/2$. Therefore, the probability the astrologer would be saved is $1/2$.

The probability of the astrologer being saved will be the same if he puts the two white balls into one urn and the two black balls into the other (Figure 1.8 (b)). His destiny will be decided by the executioner when he chooses an urn. The executioner may choose either urn with equal probability.

The best solution for the astrologer is to put a white ball into one urn and a white

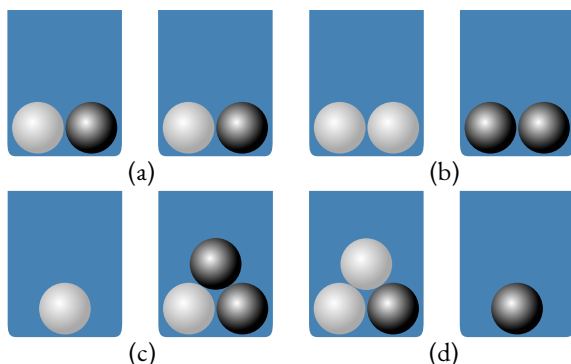


Figure 1.8: Different ways of arranging two white and two black balls for different probabilities of drawing out a ball of a given colour.

ball and two black ones into the other urn (Figure 1.8 (c)). If the executioner chooses the first urn, the astrologer will certainly be saved, but if the executioner picks the second urn, the astrologer will be saved with a probability of $1/3$. Since the executioner chooses either urn with probability $1/2$, the overall probability that the astrologer will be saved is $(1/2 \times 1) + (1/2 \times 1/3) = 2/3$.

By contrast, if the astrologer puts a black ball into one urn and a black ball and two white balls into the other (Figure 1.8 (d)), the probability of him being saved will be smallest: $(1/2 \times 0) + (1/2 \times 2/3) = 1/3$.

Thus, in order to have the greatest chance of being saved, the astrologer should distribute the balls between the urns as shown in (Figure 1.8 (c)). This is the best strategy. The worst strategy is to distribute the balls as shown in (Figure 1.8 (d)). Of course, the selection of the best strategy does not guarantee the desired outcome. Although the risk is decreased, it still remains.

Wandering In A Labyrinth. A labyrinth with treasure has a death trap, as shown in Figure 1.9. Unlucky treasure-hunters die in the trap. What is the Probability that they will avoid the trap and reach the treasure?

After walking away from the entrance A to point 1 (see Figure 1.9) a treasure-hunter may either go straight ahead (in which case he walks directly into the trap) or turn to the left (in which case he arrives at point 2) We shall suppose he picks either path at random, with equal probability, i.e. with probability $1/2$. After arriving at point 2, the treasure-hunter may either go straight ahead or turn right or turn left with probability $1/3$. The

first two paths lead to the trap, while the third path leads to point 3. The probability of someone getting from the entrance A to point 3 is the product of the probability of turning left at point 1 and the probability of turning left at point 2, i.e., $1/2 \times 1/3$. It is easy to see now that the probability of reaching point 4 from A is $1/2 \times 1/3 \times 1/2$; the probability of reaching point 5 from A is $1/2 \times 1/3 \times 1/2 \times 1/3$; and finally, the probability of reaching the treasure from A is $P^+ = 1/2 \times 1/3 \times 1/2 \times 1/3 \times 1/2 = 1/72$. The only way of getting from the entrance of the labyrinth to the treasure is shown in the figure by the dash line. The probability that a person will follow it is thus $P^+ = 1/72$, while the probability of walking into the trap is $P^- = 71/72$.

The probability P^- was calculated from the fact that $P^+ + P^- = 1$. However, we can calculate P^- directly. Let us expand P^- as the sum $P^- = P_1 + P_2 + P_3 + P_4 + P_5$ where the P_i are the probabilities of arriving at point i from A multiplied by the probability of walking into the trap from point i ($i = 1, 2, 3, 4, 5$).

$$P_1 = 1/2,$$

$$P_2 = 1/2 \times 2/3,$$

$$P_3 = 1/2 \times 1/3 \times 1/2,$$

$$P_4 = 1/2 \times 1/3 \times 1/2 \times 2/3,$$

$$P_5 = 1/2 \times 1/3 \times 1/2 \times 1/3 \times 1/2.$$

You can then find that $P_1 + P_2 + P_3 + P_4 + P_5 = 71/72$.

Discrete Random Variables

Random Variables. Suppose there is a batch of 100 manufactured articles and 11 articles are rejected as defective, 9 articles are rejected in another batch of the same size, 10 articles are rejected in the third one, 12 articles are rejected in the fourth one, etc. We use n to denote the overall number of manufactured articles in a batch and m to denote the number of rejected articles. The number n is constant (here $n =$

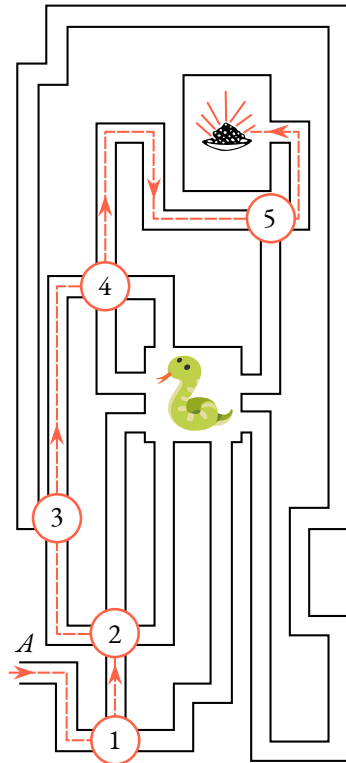


Figure 1.9: The probability of finding the treasure or a trap in a labyrinth.

100) while the value of m varies from batch to batch in a random manner. Suppose there is a *definite probability* that there will be m rejected articles in a randomly selected batch of n articles.

The number of rejected articles (the variable m) is an example of a *random variable*. *It varies randomly from one trial to another, and a certain probability is associated with the occurrence of each value of the variable*. Note that we are dealing with a discrete random variable here, i.e. it may only take a discrete set of values (the integers from 0 to 100 in this case).

There are also *continuous* random variables. For instance, the length and weight of newborn babies vary randomly from child to child and may take any value within a particular interval. There are some special features of continuous random variables which we shall discuss later; we shall first consider discrete variables.

Expected values and variance of a discrete random variable. Let x be a discrete random variable which may assume s values: $x_1, x_2, \dots, x_m, \dots, x_s$. These values are associated with the probabilities $p_1, p_2, \dots, p_m, \dots, p_s$. For instance, p_m is the probability that a variable is x_m . The sum of all the probabilities ($p_1 + p_2 + \dots + p_s$) is the probability that a trial will give one of the values x_1, x_2, \dots, x_s , (without saying which one). This probability is unity. Consequently,

$$\sum_{m=1}^s p_m = 1, \quad (1.3)$$

¹ The set of probabilities $p_1 + p_2 + \dots + p_s$ (also called the distribution of the probabilities) contains all the information needed about the random variable. However, we do not need all the probabilities for many practical purposes. It is sufficient to know two most important characteristics of a random variable: its expected value (its mathematical expectation) and its variance.

The *expected value* is an average value of the random variable taken over a large number of trials. We shall use the letter E to denote the expected value. The expected value of a random variable x is the sum of the products of each variable and its probability, i.e.

$$E(x) = p_1 x_1 + p_2 x_2 + \dots + p_s x_s,$$

or using the summation sign,

$$E(x) = \sum_{m=1}^s p_m x_m. \quad (1.4)$$

¹The notation $\sum_{m=1}^s$ means that the summation is performed over all m from 1 to s .

We also need to know how a variable deviates from the expected value, or, in other words, how much the random variable is *scattered*. The expected value of the deviation from the expected value (that is the difference $x - E(x)$) cannot be used because it is equal to zero. We can show this as follows:

$$\begin{aligned} E(x - E(x)) &= \sum_{m=1}^s p_m (x_m - E(x)), \\ &= \sum_{m=1}^s p_m x_m - E(x) \sum_{m=1}^s p_m, \\ &= E(x) - E(x), \\ &= 0. \end{aligned}$$

This is why the expected value of the *squared* deviation (rather than the expected value of the deviation itself) is used, i.e.

$$\text{var} = \sigma^2 = E(x - E(x))^2 = \sum_{m=1}^s p_m (x_m - E(x))^2. \quad (1.5)$$

This is the variance of a random variable and we shall use var to denote it. The square root of the variable $\sqrt{\text{var}}$ is called the *standard* (or *root-mean-square*) *deviation* σ of the random variable. It is easy to show that

$$\text{var} = E(x^2) - (E(x))^2. \quad (1.6)$$

Indeed,

$$\begin{aligned} \sum_{m=1}^s p_m (x_m - E(x))^2 &= \sum_{m=1}^s p_m (x_m^2 - 2x_m E(x) + E(x)^2), \\ &= \sum_{m=1}^s p_m x_m^2 - 2E(x) \sum_{m=1}^s p_m x_m + (E(x))^2 \sum_{m=1}^s p_m, \\ &= E(x^2) - 2E(x)E(x) + (E(x))^2, \\ &= E(x^2) - (E(x))^2. \end{aligned}$$

Two probability distributions are shown in Figure 1.10 (a). The two random variables possess different expected values while having the same variance. Looking at Figure 1.10 (b), we can see a different picture: the random variables possess different variances while having the same expected values.

Bernoulli's Binomial Distribution. Suppose a series of n independent identical trials is performed. The trials are independent in the sense that the results of any trial

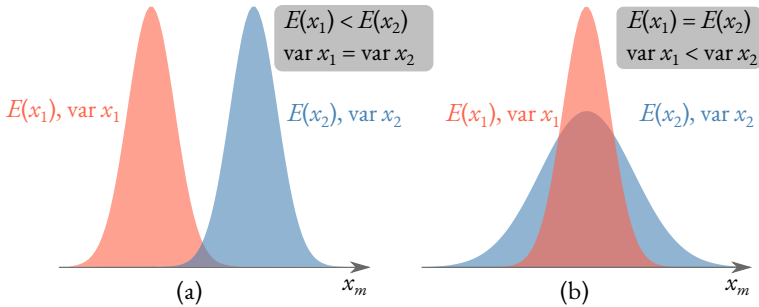


Figure 1.10: Distributions of random variables with different parameters. (a) shows two distributions with different expected values but the same variance, while (b) shows two distributions with different variance but the same expected values.

do not influence the results of any other trial. Some trials produce a desired outcome while the rest do not. Let us call the desired outcome, “event U ”. This is a random event. Suppose event U occurs in m trials. This is a random variable. Let us consider the probability $P_n(m)$ that event U will occur m times in a series of n trials.

This is a commonly occurring situation. Suppose n manufactured articles are checked. Then event U is a rejection, and $P_n(m)$ is the probability of m articles being rejected out of a set of n articles. Suppose a hospital registers n newborn babies and the event U is the birth of a girl. Hence $P_n(m)$ is the probability that there will be m girls in a set of n newborn babies. Suppose in a lottery, n tickets are checked, event U is the discovery of a prize-winning ticket, and $P_n(m)$ is the probability that m prize-winning tickets will be found out of a total of n tickets. Suppose in a physics experiment n neutrons are recorded, the event U is the occurrence of a neutron with an energy within a certain range, and $P_n(m)$ is the probability that m of the n neutrons will possess energies in the range. In all these examples, the probability $P_n(m)$ is described by the same formula which is the *binomial distribution* (sometimes named after a 17th century Swiss mathematician called Jacob Bernoulli).

The binomial distribution is derived by assuming that the probability that event U will occur in a single trial is known and does not vary from trial to trial. Let us call this probability p . The probability that event U does not occur in a single trial is $q = 1 - p$. It is important that the probability that an article is rejected does not depend in any way on how many rejected articles there are in the given batch. The probability that a girl is born in any actual delivery does not depend on whether a girl or a boy was born in the previous birth (nor on how many girls have so far been born). The probability of

winning a prize neither increases nor decreases as the lottery tickets are checked. The probability that a neutron has an energy in a given range does not change during the experiment.

Now, *once the probability p that a certain random event will occur in a single trial is known, we find the probability $P_n(m)$ of m occurrences in a series of n independent identical trials.*

Suppose the event U occurred in the first m trials but did not occur in $n - m$ trials, then the probability of the situation would be $p^m q^{n-m}$. Naturally, other orders are possible. For instance, event U may not occur in the first $n - m$ trials and occur in the rest in trials. The probability of this situation is also $p^m q^{n-m}$. There are also other possible situations. There are as many situations as there are ways choosing n elements taken in at a time (this is written $\binom{n}{m}$). The probability of each situation is identical and equals $p^m q^{n-m}$. The order in which event U occurs is inessential. It is only essential that it occurs in m trials and does not occur in the remaining $n - m$ trials. The sought probability $P_n(m)$ is the sum of the probabilities of each $\binom{n}{m}$ situation, i.e. the product of $p^m q^{n-m}$ and $\binom{n}{m}$:

$$P_n(m) = \binom{n}{m} p^m q^{n-m}. \quad (1.7)$$

There is a formula for the number of combinations of n elements taken m at a time:

$$\binom{n}{m} = \frac{n!}{m!(n-m)!} = \frac{n(n-1)(n-2) \dots (n-m+1)}{m!}. \quad (1.8)$$

Here $n! = 1 \cdot 2 \cdot 3 \cdot \dots \cdot n$ (read $n!$ as “en factorial”), by convention $0! = 1$.

Substituting (1.8) into (1.7), we can find

$$P_n(m) = \frac{n!}{m!(n-m)!} p^m q^{n-m}. \quad (1.9)$$

This is the *binomial distribution*, or the distribution of a binomial random variable. I shall explain this term below, and we shall see that

$$\sum_{m=0}^n P_n(m) = 1. \quad (1.10)$$

By way of example, let us calculate the probability that m girls are born in a group of 20 babies. Assume that the probability of delivering a girl is $1/2$. We set $p = 1/2$ and $n = 20$ in expression (1.9) and consider the integer values of variable m within the range from 0 to 20. The result can be conveniently presented as a diagram (Figure 1.11).

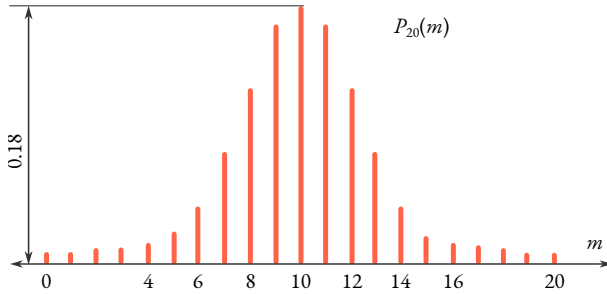


Figure 1.11: The binomial distribution.

We see that the birth of 10 girls is the most probable; the probability of delivering, for instance, 6 or 14 girls is six times smaller.

If a random variable has a binomial distribution, then its expected value is

$$E(m) = \sum_{m=0}^n m P_n(m).$$

or the product of the number of trials and the probability of the event in a single trial,

$$E(m) = np. \quad (1.11)$$

The variance of such a random variable is the product of the number of trials, the probability of the occurrence of the event in a single trial, and the probability it does not occur:

$$\text{var} = E(m^2) - (E(m))^2 = npq. \quad (1.12)$$

The normal (Gaussian) Distribution. Probability calculations using the binomial distribution are difficult for large n . For instance, in order to find the probability that 30 girls were delivered from 50 births, you have to calculate

$$P_{50}(30) = \frac{50!}{30!20!} (0.5)^{50}.$$

Note that even $20!$ is a 19—digit number. In such cases one can use a formula which is the limit of the binomial distribution at large n :

$$P_n(m) = \frac{1}{\sqrt{2\pi \text{var}}} \exp\left(-\frac{(m - E(m))^2}{2 \text{var}}\right), \quad (1.13)$$

where $E(m) = np$ and $\text{var} = npq$, and $\exp = 2.718 \dots$ is the base of natural logarithms. The distribution defined in (1.13) is called the *normal* or *Gaussian distribution*.

The Poisson Distribution. If the probability that an event will occur in a single trial is very small ($p \ll 1$), the binomial distribution at large n becomes the *Poisson* (rather than the normal) distribution, and is defined as

$$P_n(m) = \frac{(np)^m}{m!} \exp(-np). \quad (1.14)$$

This distribution is also sometimes called the *law of rare events*. It is interesting to note that the variance of a random variable with the Poisson distribution equals its expected value.

Two distributions are compared in Figure 1.12. The parameters of the first distribution are $n = 30$ and $p = 0.3$, and it is close to the normal distribution with the expected value $E(m) = 9$. The second distribution's parameters are $n = 30$ and $p = 0.05$, and it is close to the Poisson distribution with $E(m) = 1.5$.

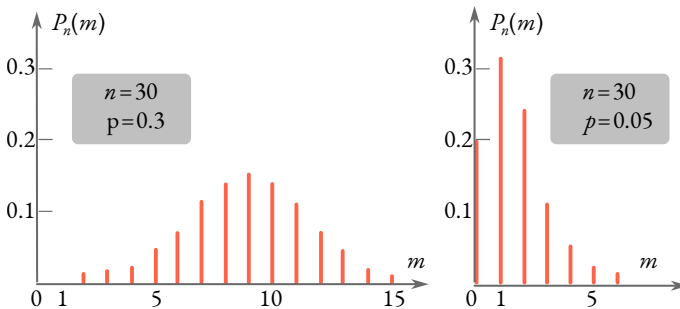


Figure 1.12: The Poisson (right) and Gaussian (left) distributions.

A Little of Mathematics. The expression $(q + p)^n$, where n is a positive integer, is called a binomial (two-term) expression of degree n . You should know about the binomial expansions of second and third degrees:

$$\begin{aligned} (q + p)^2 &= q^2 + 2qp + p^2, \\ (q + p)^3 &= q^3 + 3q^2p + 3qp^2 + p^3. \end{aligned}$$

In general (for a random integer n) the binomial expansion is

$$(q + p)^n = q^n + nq^{n-1}p + \dots + \frac{n(n-1) \dots (n-m+1)}{m!} q^{n-m} p^m + \dots + np^{n-1} + p^n.$$

Using the notation given in (1.8), we can rewrite this formula as

$$(q + p)^n = \binom{n}{0} q^n + \binom{n}{1} q^{n-1} p + \dots + \binom{n}{m} q^{n-m} p^m + \dots + \binom{n}{n-1} q p^{n-1} + \binom{n}{n} p^n.$$

Thus from (1.9), we can conclude that

$$(q + p)^n = \sum_{m=0}^n \binom{n}{m} q^{n-m} p^m = \sum_{m=0}^n P_n(m).$$

Consequently, the probabilities $P_n(m)$ coincide with the coefficients of the binomial expansion, and this is why the *binomial distribution* is so called. The probabilities q and p in a binomial distribution are such that $q + p = 1$. Therefore, $(q + p)^n = 1$. On the other hand,

$$(q + p)^n = \sum_{m=0}^n P_n(m).$$

Hence (1.10).

Continuous Random Variables

Continuous random variables are very unlike discrete ones. A continuous variable can assume any of infinite set of values, which continuously fill a certain interval. It is impossible in principle to list every value or such a variable at the very least because there is no such thing as two neighbouring values (just as it is impossible to mark two neighbouring points on the number axis). Besides, the probability of a concrete value of a continuous random variable is zero.

Can Probability Of A Possible Event Equal To Zero? You know now that an impossible event has a zero probability. However, a possible event can also have a zero probability.

Suppose a thin needle is thrown many times at random onto a strip of paper on which a number axis is marked. We can regard the x -coordinate of the point where the needle crosses the number axis (Figure 1.13 (a)) to be a continuous random variable. This coordinate varies in a random fashion from one trial to another.

We could also use a roulette instead of throwing a needle. A strip of paper with a numbered line could be pasted to the circumference of the roulette circle, as shown in Figure 1.13 (b). Wherever the freely rotating arrow of the roulette is pointing when it stops, it yields a number that will be a continuous random variable.

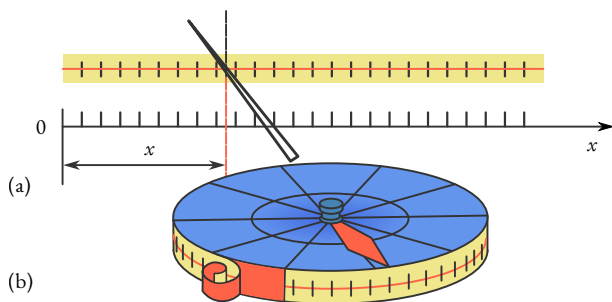


Figure 1.13: The probability that a continuous random variable will take a certain value is zero.

What is the probability of the arrow stopping at a certain point x ? In other words, what is the probability that a concrete value x of a continuous random variable is chosen? Suppose the roulette circle's radius R is divided into a finite number of identical sectors, e.g. 10 sectors (Figure 1.14). The length of the arc corresponding to the sector equals $\Delta x = 2\pi R/10$. The probability that the arrow will stop within the sector hatched in the figure is $\Delta x/2\pi R = 1/10$. Thus, the probability that the random variable will take a value from x to $x + \Delta x$ is $\Delta x/2\pi R$. Let us gradually narrow the range of numbers, i. e. divide the circle into larger numbers of sectors. The probability $\Delta x/2\pi R$ that any value is in the range from x to $x + \Delta x$ also will fall. In order to obtain the probability that the variable will take the value x exactly, we must find the limit as $\Delta x \rightarrow 0$. In this case, the probability $\Delta x/2\pi R$ becomes zero. Thus we can see that the probability that a continuous random variable will take a certain value is indeed zero.

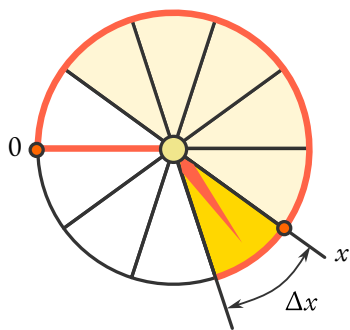


Figure 1.14: A roulette to generate continuous random variables.

That event may be both possible and possess a zero probability may seem paradoxical, but it is not. In fact there are parallels you are surely well aware of. Consider a body of volume V with a mass M . Let us select a point A within the body and consider a smaller volume V_1 which contains the point (Figure 1.15) and assign a mass M_1 to it. Let us gradually shrink the smaller volume around point A . We obtain a sequence of volumes containing A , i.e. V, V_1, V_2, V_3, \dots , and a corresponding sequence of decreasing masses:

M, M_1, M_2, M_3, \dots . The limit of the mass vanishes as the volume around A contracts to zero. We can see that a body which has a finite mass consists of points which have zero masses. In other words, the nonzero mass of the body is the *sum of an infinite number of zero masses* of its separate points. In the same way, the nonzero probability that a roulette arrow stops within a given range Δx is the *sum of an infinite number of zero probabilities* that the arrow will stop at each individual value within the considered range.

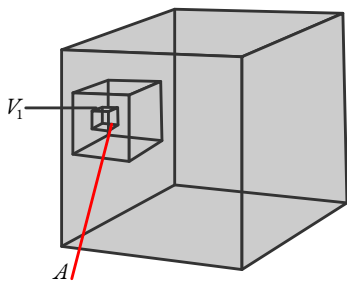


Figure 1.15: A finite non-zero mass can be generated from the sum of an infinite number of zero masses.

The Density Of A Probability. This conceptual difficulty can be avoided by using the idea *density*. Although the mass of a point within a body is zero, the body's density at the point is non-zero. If ΔM is the mass of a volume ΔV within which the point in question is located (we shall describe the point in terms of its position vector \mathbf{r} , then the density $\rho(\mathbf{r})$ at this point is the limit of the ratio $\Delta M / \Delta V$ as ΔV converges to the point at \mathbf{r} , i.e.,

$$\rho(\mathbf{r}) = \lim_{\Delta V \rightarrow 0} \frac{\Delta M}{\Delta V}.$$

If the volume ΔV is small enough, we can say that $\Delta M \approx \rho(\mathbf{r})\Delta V$. Using a strict approach, we should substitute ΔV by the differential dV .

The mass M of a body occupying volume V is then expressed by the *integral*:

$$M = \int_V \rho(\mathbf{r}) dV,$$

over the volume in question.

Probability theory uses a similar approach. When dealing with *continuous* random variables, the *probability density* is used rather than the probability itself. Let $f(x)$ be the probability density of a random variable x , and so by analogy with the mass density we have

$$f(x) = \lim_{\Delta x \rightarrow 0} \frac{\Delta p_x}{\Delta x}.$$

Here Δp_x is the probability that a random variable will take a value between x and $x + \Delta x$. The probability p that a random variable will have a value between x_1 and x_2 is, in terms

of probability density, as follows:

$$p = \int_{x_1}^{x_2} f(x) dx. \quad (1.15)$$

If the integration is over the whole range of values a random variable may take, the integral (1.15) will evaluate to unity (this is the probability of a certain event). In the example with a roulette mentioned above, the whole interval is from $x = 0$ to $x = 2\pi R$. In general, we assume the interval is infinite, when

$$\int_{-\infty}^{+\infty} f(x) dx = 1. \quad (1.16)$$

The integral is very simple in the roulette example because the probability the roulette arrow stops within an interval from x to $x + \Delta x$ *does not depend on* x . Therefore, the probability density does not depend on x , and hence,

A similar situation is encountered when the density of a body is the same at every point, i.e. when the body is *uniform* ($\rho = M/V$). More generally, density $\rho(\mathbf{r})$ varies from point to point, and so does the probability density $f(x)$.

The Expected Value And The Variance Of A Continuous Random Variable.

The *expected value* and *variance* of a discrete random variable are expressed as sums over the probability distribution (see equations (1.4) to (1.6)). When the random variable is continuous, integrals are used instead of sums and the probability density distribution is used rather than the probability distribution:

$$E(x) = \int_{-\infty}^{+\infty} x f(x) dx, \quad (1.17)$$

$$\text{var} = \int_{-\infty}^{+\infty} (x - E(x))^2 f(x) dx. \quad (1.18)$$

The Normal Distribution Of Probability Density. The normal distribution of probability density. When dealing with continuous random variables, we often encounter the *normal* distribution of probability density. This distribution is defined by the following expression (compare it with (1.13)):

$$f(x) = \frac{1}{\sigma\sqrt{2\pi}} \exp\left(-\frac{(x - E(x))^2}{2\sigma^2}\right). \quad (1.19)$$

Here σ is the standard deviation ($\sigma = \sqrt{\text{var}}$) the function (1.19) is called the *normal* or *Gaussian distribution*.

The probability density of a continuous random variable is always normal if the variance of its values is due to many different equally strong factors. It has been proved in probability theory that the sum of a large enough number of independent random variables obeying any distributions tends to the normal distribution, and the larger the number of sums the more accurately the normal distribution is.

For instance, suppose we are dealing with the production of nuts and bolts. The scatter of the inside diameter of the nut is due to random deviations in the properties of the metal, the temperature, vibration of the machine tool, changes in the voltage, wear of the cutter, etc. All of these effects act independently and approximately with the same strength. They are superimposed, and the result is that the inside diameter of the nuts is a continuous random variable with a normal distribution. The expected value of this variable should evidently be the desired inside diameter of the nuts, while the variance characterizes the scatter of the obtained diameters around the desired value.

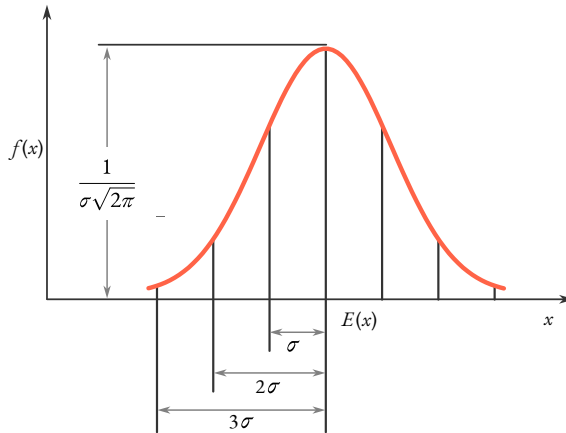


Figure 1.16: The three-sigma rule for a Gaussian distribution.

The Three-Sigma Rule. A normal distribution is shown in Figure 1.16. It has a maximum at the expected value $E(x)$. The curve (the *Gaussian curve*) is bell-shaped and is symmetric about $E(x)$. The area under the entire curve, i. e. for the interval $(-\infty < x < +\infty)$, is given by the integral $\int_{-\infty}^{+\infty} f(x)dx$. Substituting (1.19) here, it can be shown that the area is equal to unity. This agrees with (1.16), whose meaning is that the probability of a certain event is unity. Let us divide the area under the Gaussian curve

using vertical lines (see Figure 1.16). Let us first consider the section corresponding to the interval $E(x) - \sigma \leq x \leq E(x) + \sigma$. It can be shown (please believe me) that

$$\int_{E(x)-\sigma}^{E(x)+\sigma} f(x)dx = 0.683.$$

This means that the probability of x taking a value in the interval from $E(x) - \sigma$ to $E(x) + \sigma$ equals 0.683. It can also be calculated that the probability of x taking a value from $E(x) - 2\sigma$ to $E(x) + 2\sigma$ is 0.954, and the probability of x taking a value in the range of $E(x) - 3\sigma$ to $E(x) + 3\sigma$ is 0.997. Consequently, a continuous random variable with a normal distribution takes a value in the interval $E(x) - 3\sigma$ to $E(x) + 3\sigma$ with probability 0.997. This probability is practically equal to unity. Therefore, it is natural to assume for all practical purposes that a random variable will always take a value in the interval from 3σ on the right to 3σ on the left of $E(x)$. This is called the *three-sigma rule*.

Chapter 2

Decision Making

Practical demands brought forth special scientific methods that can be collected under the heading “operations research”. We shall use this term to mean the application of quantitative mathematical methods to justify decisions in every area of goal-oriented human activity.

E. S. Wentzel

These Difficult Decisions

Decision making under uncertain conditions. We often have to make decisions when not all the information is available and this uncertainty always decreases to some extent our ability to decide. For example, where to go for a vacation or holiday? This has worried me many times, since various uncertainties concerning the weather, the hotel, the entertainment at the resort, and so on, must be foreseen. We try and decide on the best variant from our experience and the advice of our friends, and we often act “by inspiration”. This *subjective* approach to decision making is justifiable when the consequences involve ourselves and relatives. However, there are many situations when a decision can affect a large number of people and therefore requires a *scientific* and *mathematically justifiable* approach rather than a subjective one.

For instance, modern society cannot function without electricity, stores of food, raw materials, etc. The stores are kept everywhere: at factories, shops, hospitals, and garages. But how large should the stores be in a particular case? It is clear that they

should not be too small, otherwise the function of the enterprise would be interrupted. Neither should they also be too large because they cost money to build and maintain: they would be dead stock. *Store-keeping* is a problem of exceptional importance. It is so complicated because a decision must always be made in conditions of uncertainty.

Two kinds of uncertainty. How should we make decisions under conditions of uncertainty? First of all, we should discover which factors are causing the uncertainty and evaluate their nature. There are two kinds of uncertainty. The first kind is due to factors which can be treated using the theory of probability. These are either *random variables* or *random functions*, and they have statistical properties (for instance, the expected value and variance), which are either known or can be obtained over time. Uncertainty of this kind is called *probabilistic* or *stochastic*. The second kind of uncertainty is caused by unknown factors which are not random variables (random functions) because the set of realizations of these factors does not possess statistical stability and therefore the notion of probability cannot be used. We shall call this uncertainty “bad”.

“So”, the reader may say, “it would seem that not every event that cannot be predicted accurately is a random event.”

“Well, yes, in a way.” Let me explain. In the preceding chapter we discussed random events, random variables, and random functions. I repeatedly emphasized that there should always be *statistical stability*, which is expressed in terms of probability. However, there are events, which occur from time to time, that do not have any statistical stability. The notion of probability is inapplicable to such events, and therefore, the term “random” cannot be used here too. For instance, we cannot assign a probability to the event of an individual pupil getting an unsatisfactory mark in a concrete subject. We cannot, even hypothetically, devise a set of uniform trials that might yield the event as one outcome. There would be no sense in conducting such a trial with a group of pupils because each pupil has his or her own individual abilities and level of preparation for the exam. The trials cannot be repeated with the same pupil because he will obviously get better and better in the subject from trial to trial. Similarly there is no way we can discuss the probability of the outcome of a game between two equally matched chess players. In all such situations, there can be no set of uniform trials, and so there is no stability which can be expressed in terms of a probability. We have “bad” uncertainty in all such situations.

I am afraid we do not consider the notion “statistical stability” and often use expressions such as “improbable”, “probable”, “most probable”, and “in all probability” to refer to events that cannot be assigned by any probability. We are apt to ascribe a probability to every event even though it might not be predictable. This is why it became necessary to refine the notion of probability early this century. This was done by A.N.

Kolmogorov when he developed an axiomatic definition of probability.

Options and the measure of effectiveness. When we speak of decision making, we assume that different patterns of behaviour are possible. They are called *options*. Let me emphasize that in the more important problems the number of options is very great. Let X be the set of options in a particular situation. A decision is made when we select one option x from this set. How do we determine which option is the most preferable or the most efficient? A quantitative criterion is needed to allow us to compare different options in terms of their effectiveness. Let us call this criterion the *measure of effectiveness*. This measure is selected for each particular *purpose*, e.g., not to be late for school, to solve a problem correctly and quickly, or to reach the cinema. A doctor wants to find an efficient method of treating his patient. A factory manager is responsible for the fulfilment of a production plan. The most efficient option is the one that suits its purpose best.

Suppose we work in a shop and our target is to maximize the receipts. We could choose profit as the measure of effectiveness and strive to maximize this measure. The selection of the measure in this example is evident. However, there are more complicated situations, when several goals are pursued simultaneously, for example, we wish to maximize profit, minimize the duration of the sales, and distribute the goods to the greatest number of customers. In such cases we have to have several measures of effectiveness; these problems are called multi-criterial.

Let W be a single measure of effectiveness. It would seem that our task is now to find an option x at which W is at a maximum (or, the other way round, at a minimum). However, we should remember that decision making occurs under conditions of uncertainty. There are unknown (random) factors (let us use ξ to denote them), which influence the end result and therefore affect the measure of effectiveness W . There is also always a set of factors known beforehand (let us designate them α). Therefore the measure of effectiveness is dependent on three groups of factors: known factors α , unknown (random) factors ξ , and the selected option x :

$$W = W(\alpha, \xi, x).$$

In the sales example, the α set is goods on sale, the available premises, the season, etc. The ξ factors include the number of customers per day (it varies randomly from day to day), the time customers arrive (random crowding is possible, which leads to long queues), the goods chosen by the customers (the demand for a given commodity varies randomly in time), etc.

Since the ξ , factors are random, the measure of effectiveness W is a random variable. Now, how is it possible to maximize (minimize) a random variable? The answer quite

clearly is that it is naturally impossible. Whichever option x is chosen, W remains random, and it cannot be maximized or minimized. This answer should not discourage the reader. It is true that under conditions of uncertainty we cannot maximize (minimize) the measure of effectiveness with a hundred per cent probability. However, an adequate selection of an option is possible with a reasonably large probability. This is where we should tackle the techniques used in decision making under conditions of stochastic uncertainty.

Substitution of random factors by means. The easiest technique is merely to substitute the random factors ξ by their means. The result is that the problem becomes completely determined and the measure of effectiveness W can be calculated precisely. It can, in particular, be either maximized or minimized. This technique has been widely used to solve problems in physics and technology. Almost every parameter encountered in these fields (e.g., temperature, potential difference, illuminance, pressure) is, strictly speaking, a random variable. As a rule, we neglect the random nature of physical parameters and use their mean values to solve the problems.

The technique is justified if the deviation of a parameter from its mean value is insignificant. However, it is not valid if the random factor significantly affects the outcome. For instance, when organizing the jobs in a motor-car repair shop, we may not neglect the randomness in the way cars fail, or the random nature of the failures themselves, or the random time needed to complete each repair operation. If we are dealing with the noise arising in an electronic device, we cannot neglect the random behaviour of electron flows. In these examples, the ξ factors must indeed be considered as random factors, we shall say they are essentially random.

Mean value optimization. If the ξ factors are essentially random, we can use a technique called *mean-value optimization*. What we do is to use the expected value $E(W)$ as the measure of effectiveness, rather than the random variable W and the expected value is maximized or minimized.

Naturally, this approach does not resolve the uncertainty. The effectiveness of an option x for concrete values of random parameters ξ may be very different from the expected one. However, using mean-value optimization means that we can be sure that after many repeated operations we shall gain overall. It should be borne in mind that mean-value optimization is only admissible when the gains of repeated operations are *totalled*, so that “minuses” in some operations are compensated by the “pluses” in others. Mean-value optimization would be justified should we be trying to increase the profit obtained, for instance, in a sales department. The profit on different days would be totalled, so that random “unlucky” days would be compensated by the “lucky” days,

But here is another example. Suppose we consider the effectiveness of the ambulance service in a large city. Let us select the elapsed time between summoning help and the ambulance arriving as the measure of effectiveness. It is desirable that this parameter be minimized. We cannot apply mean-value optimization because if one patient waits too long for a doctor, he or she is not compensated by the fact that another patient received faster attention.

Stochastic constraints. Let us put forward an additional demand. Suppose we desire that the elapsed time W till the arrival of help after a call for an ambulance be less than some value W_0 . Since W is a random variable, we cannot demand that the inequality $W < W_0$ be always true, we can only demand that it be true for some large probability, for instance, no less than 0.99. In order to take this into account we delete from the X set those options x , for which the requirement is not satisfied. These *constraints* are called *stochastic*. Naturally, the use of stochastic constraints noticeably complicates decision making.

Random Processes with Discrete States

A *random* process can be thought of as the transition of a system from one state to another occurring in a random fashion. We shall consider random processes with *discrete states* in this chapter and so our system will be supposed to have a set of discrete states, either finite or infinite. The random transitions of the system from one state to another are assumed to take place *instantaneously*.

State graphs. Random processes with discrete states can be conveniently considered using a diagram called a *state graph*. The diagram shows the possible states a system may be in and indicates the possible transitions using arrows.

Let us take an example. Suppose a system consists of two machine tools, each of which produces identical products. If a tool fails its repair is started immediately. Thus, our system has four states: S_1 both tools are operating; S_2 the first tool is under repair after a failure while the second is operating; S_3 , the second tool is under repair while the first is operating; S_4 , both tools are being repaired.

The state graph is given in Figure 2.1. The transitions $S_1 \rightarrow S_2$, $S_1 \rightarrow S_3$, $S_2 \rightarrow S_4$ and $S_3 \rightarrow S_4$ occur as a result of failures in the system. The reverse transitions take place upon termination of the repairs. Failures occur at unpredictable moments and the moments when the repairs are terminated are also random. Therefore, the system's transition from state to state is random.

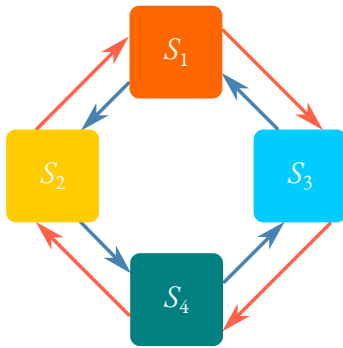


Figure 2.1: A state graph for system with four states.

Note that the figure does not show transitions $S_1 \rightarrow S_4$ and $S_4 \rightarrow S_1$. The former corresponds to the simultaneous failure of both tools and the latter to the simultaneous termination of repair of both tools. We shall assume that the probabilities of these events are zero.

Event arrival. Suppose that we have a situation in which a *stream of uniform events* follow each other at random moments. They may be telephoned orders for taxi, domestic appliances being switched on, the failures in the operation of a device, etc.

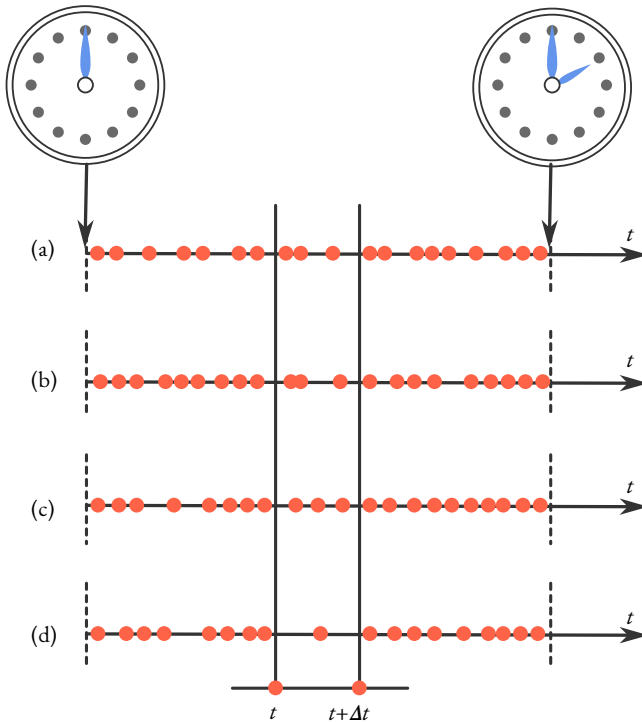


Figure 2.2: A record of taxi orders at a taxi depot.

Suppose the dispatcher at a taxi depot records the time each taxi order is made over an interval of time, for instance, from 12 a.m. to 2 p.m. We can show these moments as points on the time axis, and so the dispatcher might get the pattern illustrated in Figure 2.2 (a). This is the realization of the taxi-call arrivals during that interval of time. Three more such realizations are shown in Figure 2.2 (b), (c), and (d), and they are patterns recorded on different days. The moments when each taxi order is made in each realization are random. At the same time, the taxi-order arrivals possess statistical stability, that is, the total number of events in each interval of time varies only slightly from experiment to experiment (from one arrival realization to another). We can see that the number of events in the arrival realisations presented are 19, 20, 21, and 18.

In the preceding chapter, a random event in an experiment was an outcome which has a definite probability. When we are considering arrivals of events, we must have another meaning for the term “event”. There is no use speaking about the probability of an outcome (event) because each event is uniform, i.e. indistinguishable from the others. For instance, one taxi-order is a single event in a stream and is indistinguishable from another event. Now let us consider other probabilities, for instance, the probabilities that an event will occur during a given interval of time (suppose, from t to $t + \Delta t$, as shown in the figure) exactly once, twice, thrice, etc.

The notion of “event arrival” is applied to random processes in systems with discrete states. It is assumed that the transitions of a system from one state to another occur as a result of the effect of event arrivals. Once an event arrives, the system instantaneously changes state. For the state graph in Figure 2.1 transitions $S_1 \rightarrow S_2$ and $S_3 \rightarrow S_4$ occur due to the arrival of events corresponding to failures in the first tool, while transitions $S_1 \rightarrow S_3$ and $S_2 \rightarrow S_4$ occur due to failures of the second tool. The reverse transitions are caused by the arrival of events corresponding to the “terminations” of repair: transitions $S_2 \rightarrow S_1$ and $S_4 \rightarrow S_3$, are caused by the arrivals of repair terminations of the first tool, and transitions $S_3 \rightarrow S_1$ and $S_4 \rightarrow S_2$ to the arrivals of repair terminations of the second tool.

The system transfers from state S_i to state S_j every time the next event related to the transition arrives. The natural conclusion is that the probability of transition $S_i \rightarrow S_j$ at a definite moment in time t should equal the probability of an event arrival at this moment. There is no sense in speaking of the probability of a transition at a concrete moment t . Like the probability of any concrete value of a continuous random variable, this probability is zero, and this result follows from the continuity of time. It is therefore natural to discuss the probability of a transition (the probability of an event arrival) occurring during the interval of time from t to $t + \Delta t$, rather than its occurrence at time t . Let us designate this probability $P_{ij}(t, \Delta t)$. As Δt tends to zero, we arrive at the

notion of a *transition probability density* at time t , i.e.

$$\lambda_j(t) = \lim_{\Delta t \rightarrow 0} \frac{P_{ij}(t, \Delta t)}{\Delta t}. \quad (2.1)$$

This is also called the *arrival rate of events* causing the transition in question.

In the general case, the arrival rate depends on time. However, it should be remembered that the dependence of the arrival rate on time is not related to the location of “dense” or “rare” arrival realisations. For simplicity’s sake, we shall assume that the transition probability density and therefore the event arrival rate does not depend on time. i.e. we shall consider *steady-state* arrivals.

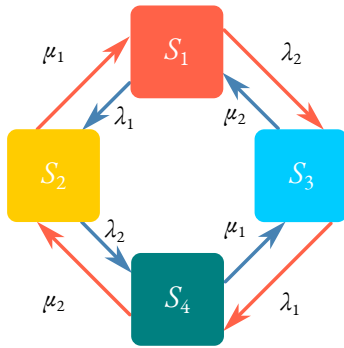


Figure 2.3: A state graph for system with four states with arrival rates.

The Chapman-Kolmogorov equations for steady state. Let us use p_i to denote the probability that a system is in state S_i (since our discussion is only for *steady-state* arrivals, the probabilities p_i are independent of time). Let us consider the system whose state graph is given in Figure 2.1. Suppose λ_1 is the arrival rate for failures of the first tool and λ_2 that for the second tool; let μ_1 be the arrival rate for repair terminations of the first tool and μ_2 that for the second tool. We have labelled the state graph with the appropriate arrival rates, see Figure 2.3.

Suppose there are N identical systems described by the state graph in Figure 2.3. Let $N \gg 1$. The number of systems with state S_i , is Np_i (this statement becomes more accurate the larger N is). Let us consider a concrete state, say, S_1 . Transitions are possible from this state to states S_2 and S_3 with probability $\lambda_1 + \lambda_2$, per unit time. (Under steady state, the probability density is the probability for the finite time interval Δt divided by Δt .) Therefore, the number of departure: from state S_1 , per unit time in the considered set of systems is $Np_1(\lambda_1 + \lambda_2)$. We can discern a general rule here: the number of transitions $S_i \rightarrow S_j$ per unit time is the product of the number of systems with state S_i (the initial state) by the probability of the transition per unit time. We have considered departures from state S_1 . The system arrives at this state from S_2 and S_3 . The number of arrivals at S_1 per unit time is $Np_2\mu_1 + Np_3\mu_1$. Since we are dealing with steady states, the number of departures and arrivals for each particular

state should be balanced. Therefore.

$$Np_1(\lambda_1 + \lambda_2) = Np_2\mu_1 + Np_3\mu_2.$$

By setting up similar balances of arrivals and departures for each of the four states and eliminating the common factor N in the equations, we obtain the following equations for probabilities p_1 , p_2 , p_3 and p_4 :

$$\begin{aligned} \text{for state } S_1 : (\lambda_1 + \lambda_2) p_1 &= \mu_1 p_2 + \mu_2 p_3, \\ \text{for state } S_2 : (\lambda_2 + \mu_1) p_2 &= \lambda_1 p_1 + \mu_2 p_4, \\ \text{for state } S_3 : (\lambda_1 + \mu_2) p_3 &= \lambda_1 p_1 + \mu_1 p_4, \\ \text{for state } S_4 : (\mu_1 + \mu_2) p_4 &= \lambda_2 p_2 + \lambda_1 p_3. \end{aligned}$$

It is easy to see that the fourth equation can be obtained by summing the first three. Instead of this equation, let us use the equation

$$p_1 + p_2 + p_3 + p_4 = 1,$$

which means that the system must be in one of the four states. Therefore, we have the following system of equations:

$$\left. \begin{aligned} S_1 : (\lambda_1 + \lambda_2) p_1 &= \mu_1 p_2 + \mu_2 p_3, \\ S_2 : (\lambda_2 + \mu_1) p_2 &= \lambda_1 p_1 + \mu_2 p_4, \\ S_3 : (\lambda_1 + \mu_2) p_3 &= \lambda_2 p_2 + \mu_1 p_4, \\ p_1 + p_2 + p_3 + p_4 &= 1. \end{aligned} \right\} \quad (2.2)$$

These are the *Chapman-Kolmogomov equations* for the system whose state graph is shown in Figure 2.3.

Which innovation should be chosen? Let us analyze a concrete situation using equations (2.2). The state graph (see Figure 2.3) corresponding to these equations describes a system which, we assumed, consists of two machine tools each producing identical goods. Suppose the second tool is more modern and its output rate is twice that of the first tool. The first tool generates (per unit time) an income of five conventional units, while the second one generates one of ten units. Regrettably, the second tool fails, on the average, twice as frequently as does the first tool: hence $\lambda_1 = 1$ and $\lambda_2 = 2$. The arrival rates for repair termination are assumed to be $\mu_1 = 2$ and $\mu_2 = 3$. Using these

arrival rates for failure and repair termination. let us rewrite (2.2) thus

$$\left. \begin{aligned} 3p_1 &= 2p_2 + 3p_3, \\ 4p_2 &= p_1 + 3p_4, \\ 4p_3 &= 2p_1 + 2p_4, \\ p_1 + p_2 + p_3 + p_4 &= 1. \end{aligned} \right\}$$

This system of equations can be solved to yield $p_1 = 0.4$, $p_2 = 0.2$, $p_3 = 0.27$ and $p_4 = 0.13$. This means that, on the average, both tools operate simultaneously (state S_1 in the figure) 40 per cent of the time, the first tool operates while the second one is being repaired (state S_2) 20 per cent of the time, the second tool operates while the first one is being repaired (state S_3) 27 per cent of the time, and both tools are simultaneously being repaired (state S_4) 13 per cent of the time. It is easy to calculate the income this tool system generates per unit time: $(5 + 10) \times 0.4 + 5 \times 0.2 + 10 \times 0.27 = 9.7$ conventional units.

Suppose an innovation is suggested which would reduce the repair time of either the first or second tool by a factor of two. For technical reasons, we can only apply the innovation to one tool. Which tool should be chosen, the first or the second? Here is a concrete example of a practical situation when, using probability theory, we must justify our decision scientifically

Suppose we choose the first tool. Following the introduction of the innovation, the arrival rate of its repair termination increases by a factor of two, whence $\mu_1 = 4$ (the other rates remain the same, i. e. $\lambda_1 = 1$, $\lambda_2 = 2$ and $\mu_2 = 3$). Now equations (2.2) are

$$\left. \begin{aligned} 3p_1 &= 4p_2 + 3p_3, \\ 6p_2 &= p_1 + 3p_4, \\ 4p_3 &= 2p_1 + 4p_4, \\ p_1 + p_2 + p_3 + p_4 &= 1. \end{aligned} \right\}$$

After solving this system, we find that $p_1 = 0.48$, $p_2 = 0.12$, $p_3 = 0.32$, and $p_4 = 0.08$. These probabilities can be used to calculate the income our system will now generate: $(5 + 10) \times 0.48 + 5 \times 0.12 + 10 \times 0.32 = 11$ conventional units.

If we apply the innovation to the second tool, the rate μ_2 , will be doubled. Now

$\lambda_1 = 1$, $\lambda_2 = 2$, $\mu_1 = 2$ and $\mu_2 = 6$, and equations (2.2) will be

$$\left. \begin{aligned} 3p_1 &= 2p_2 + 6p_3, \\ 4p_2 &= p_1 + 6p_4, \\ 7p_3 &= 2p_1 + 2p_4, \\ p_1 + p_2 + p_3 + p_4 &= 1. \end{aligned} \right\}$$

This system yields: $p_1 = 0.5$, $p_2 = 0.25$, $p_3 = 0.17$, and $p_4 = 0.08$, whence the Income is $(5 + 10) \times 0.5 + 5 \times 0.25 + 10 \times 0.17 = 10.45$ conventional units. Therefore it is clearly more profitable to apply the innovation to the first too.

Queueing Systems

The problem of queueing. Modern society cannot exist without a whole network of *queueing systems*. These include telephone exchanges, shops, polyclinics, restaurants, booking offices, petrol stations, and hairdressers. Despite their diversity, these systems have several things in common and common problems.

When we seek the assistance of a doctor or service from a cafe, restaurant, or barber, we must wait for our turn in a queue, even if we telephone to make an appointment, that is, reserve our place in a queue without actually attending physically. Clearly, we wish to be served straight away and waiting can be frustrating.

It is clear that the source of the problem is the *random nature* of the demands for attention in queueing systems. The arrival of calls at a telephone exchange is random as is the duration of each telephone conversation. This randomness cannot be avoided. However, it can be taken into account and, as a consequence, we can rationally organize a queueing system for all practical purposes. These problems were first investigated in the first quarter of this century. The mathematical problems for simulating random processes in systems with discrete states were formulated and considered, and a new field of investigation in probability theory was started.

Historically, queueing theory originated in research on the overloading of telephone exchanges, a severe problem in the early 20th century. The initial period in the development of the queueing theory can be dated as corresponding to the work of the Danish scientist A. Erlang in 1908-1922. Interest in the problems of queueing rapidly increased. The desire for more rational servicing of large numbers of people led to investigations of queue formation. It soon became evident that the problems dealt with in queueing theory went well beyond the sphere of rendering service and the results are applicable to

a wider range of problems.

Suppose a workman is operating several machine tools. Failures requiring urgent repairs occur at random moments, and the duration of each repair is a random variable. The result is a situation similar to a common queueing system. However, this is a problem of servicing many tools by a worker rather than servicing many people by a queueing system.

The range of practical problems to which queueing theory can be applied is uncommonly wide. We need the theory when we want, say, to organize the efficient operation of a modern sea port, when, for instance, we analyze the servicing rate of a large berth. We apply to queueing theory when we look at the operation of a Geiger-Müller counter. These devices are used in nuclear physics to detect and count ionizing particles. Each particle entering a tube in the counter ionizes gas in the tube, the ionization being roughly independent of the particle's nature and energy, and so a uniform discharge across the tube is generated. But when one discharge is under way, a new particle cannot be registered ("serviced") by the same counter. The moment each particle enters the tube is random, as is the duration of the discharge (the "servicing" time). This is a situation typical for queueing systems.

Basic notions. A queueing system is set up to organize the service of a *stream of requests*. The request may be a new passenger in a booking office, a failure in a machine tool, a ship mooring, or a particle entering a Geiger-Müller counter. The system may have either one or several *servers*. When you go to a large barbershop or hairdresser and want to know the number of barbers or hairdressers, you are in effect asking for the number of servers in the establishment. In other situations, the servers may be the number of cashiers in a booking office, the number of telephones at a post office for making trunk calls, the number of berths in a port, or the number of pumps at a petrol station. If, on the other hand, we wish to see a particular doctor, we are dealing with a single-server queueing system.

When we consider the operation of a queueing system, we must first take into account the number of servers, the number of requests arriving at the system per unit time, and the time needed to service a request. The number of requests arriving at the system, the moments they arrive, and the time needed to service a request are, as a rule, *random* factors. Therefore, queueing theory is a *theory of random processes*.

Random processes of this type (i. e. with *discrete states*) were discussed in the preceding section. A system transfers from state to state when each request arrives at the system and when the requests are serviced. The latter is given by the rate at which requests can be served by a single, continuously occupied server.

Queueing systems. There are two sorts of queueing system: *systems with losses* and *systems with queues*. If a request arrives at a system with losses when all the servers are occupied, the request is “refused” and is then lost to the system. For example, if we want to telephone someone and the number is engaged, then our request is refused and we put down the receiver. When we dial the number again, we are submitting a new request.

The more common types of system are those with queues or systems with waiting. This is why it is called the *theory of queueing*. In such a system, if a request (or customer) arrives when all the servers are occupied, the customer takes a place in a *queue* and waits for a server to become free. There are systems with *infinite queues* (a queueing customer is eventually served and the number of places in the queue is unlimited) and systems with *finite queues*. There are different sorts of restriction, i.e. the number of customers queueing at the same time may be limited (the queue cannot be longer than a certain number of customers and any new customer is refused); the duration of a customer’s stay in the queue may be limited (after a certain length of time queueing, an unserved customer will leave the queue); or the time the system operates for may be restricted (customers may only be served for a certain interval of time).

The service order is also important. Customers are commonly served “first come first served”. However, *priority servicing* is also possible, i.e. a newcomer to a queue is served first irrespective of the queue. A customer with a high priority may arrive at the system and interrupt the servicing of a customer with a lower priority, which may already start, or the higher priority customer may have to wait until the servicing has been completed. The priority is *absolute* in the first case and *relative* in the second. Queueing systems are always *multi-critical*, that is, they have a *set* of measures by which their effectiveness can be estimated. These may be the average number of customers served by the system per unit time, the average number of occupied servers, the average number of customers in the queue, the average time of waiting for servicing, the average percentage of refused customers, and the probability a customer arriving at the system is immediately served. There are other measures of such systems’ effectiveness. It is quite natural that when organizing the operation of a queueing system we should strive to reduce the average number of customers in the queue, and to reduce the time of waiting for servicing. It is also desirable to maximize the probability that a customer arriving at the system is served immediately, to minimize the average percentage of refused customers, and so on.

This eventually means that the productivity of the system must be increased (i.e. the time needed to service each customer be decreased), the system’s operation be rationalized, and the number of servers made as large as possible. However, by raising the number of servers, we cannot avoid decreasing the average number of occupied

servers. This means that the duration of the time for which a server is not occupied will increase, i.e. the server will be idle for some time. The result is that the system's operational efficiency is lowered. Therefore we must in some way *optimize* the system's operation. The number of servers should not be too small (to eliminate long queues and to keep the number of refusals small), but it should also not be too large (so that the number and duration of idle periods for each server is small).

Systems with losses. The simplest type of queueing system is a *single-server system with losses*. Here are some examples: a system with only one telephone line or a particle detector consisting of only one Geiger-Müller counter. The state graph for such a system is shown in Figure 2.4 (a). When the server is unoccupied, the system is in state S_0 , and when the server is occupied, it is in state S_1 . The customer's arrival rate is λ , and the service completion rate is μ . This state graph is very simple. When the system is in state S_0 a customer arriving at the system transfers it to state S_1 , and the servicing starts. Once the servicing is completed, the system returns to state S_0 and is ready to serve a new customer.

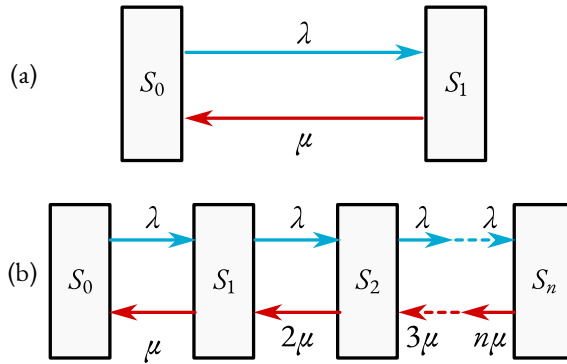


Figure 2.4: State graph of a system with losses.

We shall not go into detail on this type of system and go straight over to a more general case, an *n-server system with losses*. An example is a system consisting of n telephone lines. Erlang, the founder of the queueing theory, considered precisely this system. The corresponding state graph is given in Figure 2.4 (b). The states of the system are designated as follows: S_0 when all servers are unoccupied, S_1 when one server is occupied and the others are unoccupied, S_2 , when two servers are occupied while the others are unoccupied, and so on, and S_n is the state when all n servers are occupied. As in the preceding example, λ is the customer arrival rate, and μ is the service-completion rate.

Suppose the system is in state S_0 . When a customer request arrives, one of the servers becomes occupied, and the system is transferred to state S_1 . If the system is in state S_1 , and a new customer arrives, two servers become occupied, and the system is transferred from S_1 to S_2 . Thus, each customer (with the rate of arrivals λ) transfers the system from one state to the adjacent one *from left to right* (see the state graph in the figure). The arrival of events leading to transitions to adjacent states *from right to left* is somewhat more complicated. If the system is in the state S_1 (only one server is occupied), the next service-completion event will disengage the server and transfer the system to state S_0 . Let me remind you that the service-completion rate is μ . Now suppose the system is in S_1 , i.e. two servers are occupied. The average time of service for each server is the same. Each server is disengaged with the rate it when services are completed. As to the transition of the system from S_2 to S_1 , it is indifferent as to which of the two servers is unoccupied. Therefore, events which transfer the system from S_2 to S_1 arrive at the rate 2μ . As to the transition of the system from S_3 to S_2 , it is indifferent as to which of the three occupied servers is disengaged. Events which transfer the system from S_3 to S_2 arrive at the rate 3μ , and so forth. It is easy to see that the rate of event arrival which transfers the system from S_k to S_{k-1} is $k\mu$.

Let us assume that the system is in a steady state. Applying the rule from the preceding section and using the state graph in Figure 2.4 (b), we can compile the Chapman-Kolmogorov equations for the probabilities $p_0, p_1, p_2, \dots, p_n$, (recall that p_i is the probability that the system is in the state S_i). We obtain the following system of equations:

$$\left. \begin{aligned} \lambda p_0 &= \mu p_1, \\ (\lambda + \mu)p_1 &= \lambda p_0 + 2\mu p_2, \\ (\lambda + 2\mu)p_2 &= \lambda p_1 + 3\mu p_3, \\ \dots &\dots \dots \dots, \\ (\lambda + k\mu)p_k &= \lambda p_{k-1} + (k+1)\mu p_{k+1}, \\ \dots &\dots \dots \dots, \\ [\lambda + (n-1)\mu]p_{n-1} &= \lambda p_{n-2} + n\mu p_n, \\ p_0 + p_1 + p_2 + \dots + p_n &= 1. \end{aligned} \right\} \quad (2.3)$$

This set of equations can be solved easily. Using the first equation, we can express p_1 in terms of p_0 and substitute it into the second equation. Then we can express p_2 in the second equation in terms of p_n and substitute it into the third one, and so forth. At the last but one stage, we express p_n in terms of p_0 . And finally, the results obtained at each

stage can be substituted into the last equation to find the expression for p_0 . Thus

$$p_0 = \left[1 + \frac{\lambda}{\mu} + \frac{(\lambda/\mu)^2}{2!} + \frac{(\lambda/\mu)^3}{3!} + \dots + \frac{(\lambda/\mu)^n}{n!} \right]^{-1}, \quad (2.4)$$

$$p_k = \frac{(\lambda/\mu)^k}{k!} p_0 \quad (k = 1, 2, 3, \dots, n).$$

A customer's request is refused if it arrives when all n servers are engaged, i.e. when the system is in state S_n . The probability that the system is in S_n equals p_n . This is the probability that a customer arriving at the system is refused and the service is not rendered. We can find the probability that a customer arriving at the system will be served,

$$Q = 1 - p_n = 1 - \frac{(\lambda/\mu)^n}{n!} p_0. \quad (2.5)$$

By multiplying Q by λ , we obtain the service-completion rate of the system. Each occupied server serves μ customers per unit time, so we can divide Q by μ and find the average number of occupied servers in the system,

$$E(N) = \frac{\lambda}{\mu} \left(1 - \frac{(\lambda/\mu)^n}{n!} p_0 \right). \quad (2.6)$$

How many servers are required? Let us consider a concrete example. Suppose a telephone exchange receives 1.5 requests per minute on the average, and the service completion rate is 0.5 request per minute (the average service time for one customer is two minutes). Therefore, $\lambda/\mu = 3$. Suppose the exchange has three servers (three telephone lines). Using formulas (2.4)–(2.6) for $\lambda/\mu = 3$ and $n = 3$, we can calculate that the probability of servicing the arriving customers is only 65 per cent. The average number of engaged lines is 1.96, which is 65 per cent of the total number of lines. Thus, 35 per cent of the customers are refused and not served. This is too much.

We may decide on increasing the number of servers. Suppose we add one more, a fourth line. Now the probability of a customer being served increases to 79 per cent (the probability of being turned away decreases to 21 per cent). The average number of engaged lines becomes 2.38, which is 60 per cent of the total number of lines. It would appear that the decision to install a fourth line is reasonable because a relatively small reduction in the percentage of occupied servers (from 65 to 60 per cent) results in a significant rise in the probability to be served, from 65 to 79 per cent. Any further increase in the number of lines may become unprofitable because the effectiveness of the system may fall due to the increasing idleness of the lines. A more detailed analysis would

then be required to allow for the cost of installing each new line. Let me remark that at $n = 5$ we get $Q = 89$ per cent and $E(N)/n = 53$ per cent, while for $n = 6$, $Q = 94$ per cent and $E(N)/n = 47$ per cent.

Single-server systems with finite queues. Suppose the number of queueing customers is restricted, and the queue may only accommodate m customers. If all places in the queue are occupied, a newcomer is turned away. For example, a petrol station with only one pump (only one server) and a parking area for no more than m cars. If all the places at the station are occupied, the next car arriving at the station will not stop and will go on to the next. The state graph for this system is shown in Figure 2.5 (a). Here S_0

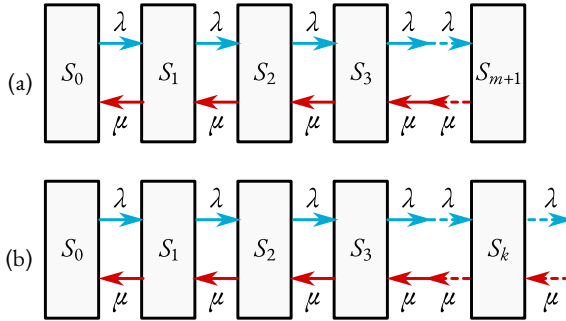


Figure 2.5: State graph for a system (a) with finite queues and (b) with infinite queues.

means the server is unoccupied, S_1 the server is occupied, S_2 the server is occupied and there is one customer in the queue, S_3 the server is occupied and there are two customers in the queue, ..., S_{m+1} means the server is occupied and there are m customers in the queue. As before, λ is the customer arrival rate and μ is the service completion rate. The Chapman-Kolmogorov equations for steady state are

$$\left. \begin{aligned} \lambda p_0 &= \mu p_1, \\ (\lambda + \mu) p_1 &= \lambda p_0 + 2\mu p_2, \\ \dots &\dots \dots \dots, \\ (\lambda + \mu) p_m &= \lambda p_{m-1} + \mu p_{m+1}, \end{aligned} \right\} \quad (2.7)$$

$$p_0 + p_1 + p_2 + \dots + p_m + p_{m+1} = 1.$$

By solving this system and introducing the designation $\rho = \lambda/\mu$ we obtain

$$p_0 = \frac{1}{1 + \rho + \rho^2 + \rho^3 + \dots + \rho^{m+1}} = \frac{1 - \rho}{1 - \rho^{m+2}}, \quad p_k = \rho^k p_0. \quad (2.8)$$

A customer is turned away if the server is engaged and there are m customers in the queue, i.e. when the system is in the state S_{m+1} . Therefore, the probability a customer is turned away is p_{m+1} . The average number of customers in the queue is evidently

$$E(r) = \sum_{k=1}^m k p_{k+1}$$

(p_{k+1} is the probability of k customers being in the queue). The average waiting time in the queue is the ratio $E(r)/\lambda$.

Suppose one car arrives at the petrol station per minute ($\lambda = 1$ customer per minute) and a car is filled, on average, within two minutes ($\mu = 1/2$). Therefore, $\rho = \lambda/\mu = 2$. If the number of places in the queue $m = 3$, it is easy to calculate that the probability of a customer being refused is 51.6 per cent while the average waiting time in the queue is 2.1 min. Suppose that in order to decrease the probability of a customer being refused we double the number of places in the queue. It turns out that at $m = 6$ the probability of refusal is 50.2 per cent, i. e. it is, in fact, the same, but the waiting time in the queue noticeably increases to 5 min. It is clear from (2.8) that if $\rho > 1$, the probability of being refused stabilizes with increasing m and tends, to $(\rho - 1)/\rho$. In order to reduce the probability of being refused significantly, it is necessary (if it is not possible to decrease ρ) to use multi-server systems.

Single-server systems with infinite queues. This sort of queueing system is rather common: for example, a doctor receiving patients, a single public telephone, or a port with only one berth at which a single ship can unload. The state graph for the system is given in Figure 2.5 (b). Here S_0 means that the server is unoccupied, S_1 the server is occupied, S_2 the server is occupied and there is one customer in the queue, S_3 the server is occupied and there are two customers in the queue, and S_k means that the server is occupied and there are $k - 1$ customers in the queue, and so on.

Up till now, we considered graphs with a finite number of states. However, here is a system with an infinite number of discrete states. Is it possible to discuss a steady state for such a system? In fact we can. It is only necessary that the inequality $\rho < 1$ holds true. If so, then the sum $1 + \rho + \rho^2 + \rho^3 + \dots + \rho^{m+1}$ in (2.8) can be substituted by the sum of the decreasing geometric progression $1 + \rho + \rho^2 + \rho^3 + \dots = 1/(1 - \rho)$. The result is

$$p_0 = 1 - \rho \quad \text{and} \quad p_k = \rho^k p_0. \quad (2.9)$$

If $\rho \geq 1$, then the system does not have a steady state, i.e. the queue increases infinitely as $t \rightarrow \infty$.

Method of Statistical Testing

A *statistical testing* involves numerous repetitions of uniform trials. The result of any individual trial is random and is not of much interest. However, a large number of results is very useful. It shows some stability (*statistical stability*) and so the phenomenon being investigated in the trials can be described quantitatively. Let us consider a special method for investigating a random process based on statistical testing. The technique is commonly called the *Monte Carlo method*.

In fact neither the city of Monte Carlo, the capital of the independent principality of Monaco nor its inhabitants nor guests are in any way related to the considered method. Instead, the city is known for its casinos where tourists pay good money playing roulette, and a roulette wheel could be the city's emblem. At the same time, a roulette is a generator of random numbers and this is what is involved when the Monte Carlo method is used.

Two examples indicating the usefulness of statistical testing.

First example. Look at Figure 2.6. It contains a square with side r in which a quarter circle of radius r is inscribed. The ratio of the yellow area to the area of the square is $(\pi r^2)/4r^2 = \pi/4$. This ratio and, therefore, the value of n can be obtained using the following statistical test. Let us place a sheet of paper with the figure on a horizontal surface and let us throw small grains on this paper. We should not aim so that any grain can fall on any part of the paper with equal probability. It is possible, for instance, to blindfold the person throwing the grains. The grains will be distributed over the surface of the paper in a random fashion (Figure 2.6 (b)). Some will land outside the square, but we shall not consider them. We now count the number of grains within the square (and call this number N_1) and count the grains within the yellow area (calling it N_2). Since any grain may land with equal probability on any part of the figure, the ratio N_2/N_1 when the number of trials is large, will approximate the ratio of the yellow area to the area of the square, i.e. the number $\pi/4$. This approximation will become more accurate as the number of trials increases. This example is interesting because a definite number (the number π) can be found following a statistical testing. It can be said that randomness is used here to obtain a deterministic result, an approximation of the real number π .

Second example. Statistical testing is used much more commonly to investigate *random events* and *random processes*. Suppose someone assembles a device consisting of three parts (A , B , and C). The assembler has three boxes containing parts A , B , and C , respectively. Suppose half the parts of each type are larger than the standard and the other half are smaller. The device cannot operate when all three parts are larger than the

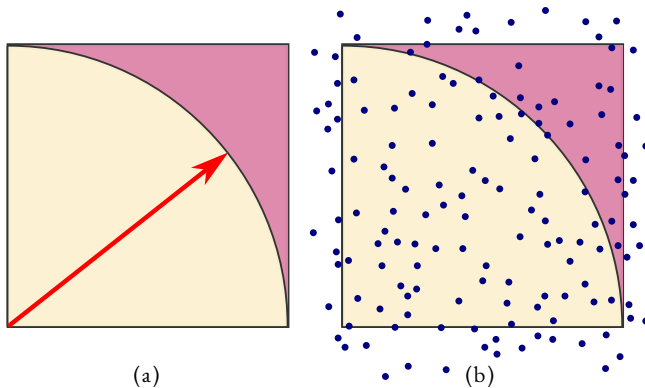


Figure 2.6: Finding out value of π using a random distribution.

norm. The assembler takes the parts from the boxes at random. What is the probability that a normally operating device will be assembled?

Naturally, this example is rather simple and the probability can easily be calculated. The probability of assembling a device that does not work is the probability that all three parts will be larger than the norm, and this equals $1/2 \times 1/2 \times 1/2 = 1/8$. Therefore, the probability that a normally operating device will be assembled is $1 - 1/8 = 0.875$.

Let us forget for a time that we can calculate the probability and instead use statistical testing. We should choose trials such that each one has equally probable outcomes, for instance, tossing a coin. Let us take three coins: A , B , and C . Each coin corresponds to a part used to assemble the device. Heads will mean that the respective part is larger than the norm while tails will mean that it is smaller. Having agreed on this, let us start the statistical testing. Each trial involves tossing all three coins. Suppose after N trials ($N \gg 1$) three heads were recorded in n trials. It is easy to see that the ratio $(N - n)/N$ is the approximation of the probability in question.

Naturally, we could use any other random number generator instead of coins. It would also be possible, for instance, to throw three dice, having agreed to relate three faces of each die with larger than normal parts and three faces with smaller parts.

Let me emphasize that the randomness in these examples was a positive factor rather than a negative one, and was a tool which allowed us to obtain a needed quantity. Here chance works for us rather than against us.

Random number tables come into play. Nobody uses statistical testing in simple

practical situations like the ones described above. It is used when it is difficult or even impossible to calculate the probability in question. Naturally you might ask whether a statistical testing would be too complicated and cumbersome. We threw grains or three coins in the examples. What will be required in complicated situations? Maybe, there will be practically unsurmountable obstacles?

In reality, it is not necessary to stage a statistical experiment with random trials. Instead of real trials (throwing grains, dice, etc.), we need only use *random number tables*. Let me show how this can be done in the above two examples.

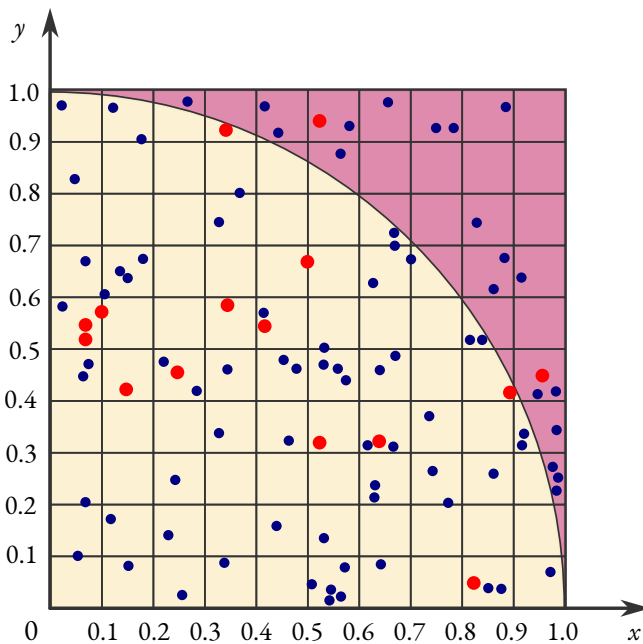


Figure 2.7: Finding out value of π using a random number table.

First example. Let us again discuss the picture in Figure 2.6. We now plot two coordinate axes along the sides of the square and select the scales such that the side of the square equals unity (Figure 2.7). Now instead of throwing grains, we take the random number table in Figure 1.6 and divide each number by 10 000 so that we obtain a set of random numbers between 0 and 1. We take the numbers in the odd lines as x -coordinates and the ones directly below as the y -coordinates of random points. We

plot the points onto the diagram, systematically moving along the random number table (for instance, first down the first column from top to bottom, and then down the second column, and so on). The first fifteen random points are shown in the picture in red, and they have the coordinates as shown in Table 2.1. The figure contains 85 random points in black. From the diagram, it is easy to calculate that using the first fifteen points $N_2/N_1 = 13/15$ and therefore $\pi = 3.47$ while for a hundred points $N_2/N_1 = 78/100$ and therefore $\pi = 3.12$.

(0.0655, 0.5255)
 (0.6314, 0.3157)
 (0.9052, 0.4105)
 (0.1437, 0.4064)
 (0.1037, 0.5718)
 (0.5127, 0.9401)
 (0.4064, 0.5458)
 (0.2461, 0.4320)
 (0.3466, 0.9313)
 (0.5179, 0.3010)
 (0.9599, 0.4242)
 (0.3585, 0.5950)
 (0.8462, 0.0456)
 (0.0672, 0.5163)
 (0.4995, 0.6751)

Table 2.1: Coordinates of fifteen random numbers shown in red in the Figure 2.7.

Second example. Instead of tossing coins, we can use the same random number table (see Figure 1.6). Each number over 5000 can be replaced by a “+” sign and the rest replaced by a “-” sign. The result is a table consisting of a random set of pluses and minuses. We divide these signs into triples as shown in Figure 2.8. Each triple corresponds to a set of three parts. A “+” sign means that a part is larger than the norm while a “-” sign means it is smaller. The approximation of the sought probability is the ratio $(N - n)/N$, where N is the total number of triples and n is the number of triples with three pluses (they are shaded in the figure). It can be seen that $(N - n)/N = 0.9$ in this case, and this is close enough to the accurate value 0.875.

Thus, we have reduced statistical testing to operations on a random number table and used our desk instead of an experimental bench. Rather than performing very many trials, we just look at a random number table.

Computers come into play. Instead of sweating over a random number table, we could program a computer to do the job. We place a random number table in the computer’s memory and program it to search the random numbers and sort them as necessary. In our two examples, we would do the following.

First example. The computer has to check the coordinates of each random point to see whether $x^2 + y^2 < 1$. It counts the number of points for which this is true (the number is N_2) and the number of points for which it is false (this number of points will be the difference $N_1 - N_2$).

Chance complicates our investigation and so randomness is used to investigate it. *Secondly*, this method is *universal* because it is not restricted by any assumption, simplification, or model. There are two basic applications. The first is the investigation of random processes which cannot be dealt with analytically due to their complexity. The second is to verify the correctness and accuracy of an analytical model applied in concrete situations.

The Monte Carlo method was first widely used in operations research, in looking for optimal decisions under conditions of uncertainty, and in treating complicated multi-criterial problems. The method is also successfully used in modern physics to investigate complex processes involving many random events.

A Monte Carlo simulation of a physical process. Let us consider the flow of neutrons through the containment shield of a nuclear reactor. Uranium nuclei split in the core of the reactor and this is accompanied by the creation of high-energy neutrons (of the order of several million electron volts). The reactor is surrounded by a shield to protect the working areas (and therefore, the personnel) from the radiation. The wall is bombarded by an intense flow of neutrons from the reactor core. The neutrons penetrate into the wall and collide with the nuclei of the atoms of the wall. The result is that the neutrons may either be absorbed or scattered. If scattered, they give up some of their energy to the scattering nuclei.

This is a complicated physical process *involving many random events*. The energy and the direction of a neutron when it leaves the reactor core and enters the wall are random, the length of the neutron path before it first collides is random, the nature of collision (absorption or scattering) is random, the energy and the direction of the scattered neutron are random, etc. Let me show in general how the Monte Carlo method is applied to analyze the process. Obviously the computer is first programmed with data on the elementary collisions between neutrons and the wall nuclei (the probabilities of absorption and scattering) the parameters of the neutron flow into the wall, and the properties of the wall. The computer model simulates a neutron with a randomly selected energy and direction (when it leaves the reactor core and enters the wall) in line with appropriate probabilities. Then it simulates (bearing in mind the relevant probabilities) the flight of the neutron until it first collides. Then the first collision is simulated. If the neutron is not absorbed, subsequent events are simulated, i.e. the neutron's flight until its second collision, the collision itself, and so on. The "history" of the neutron is determined from the moment it penetrates the wall until it is either absorbed, scattered back into the reactor core, or scattered into the working area.

The computer simulation is repeated for very many neutrons until a set of possible trajectories of neutrons within the wall is obtained (Figure 2.9). Each trajectory is

the result of one statistical trial simulating the “history” of Chance complicates our investigation and so randomness is used to investigate it. Secondly, this method is universal because it is not restricted by any assumption, simplification, or model. There are two an individual neutron. Given an enormous set of trials the neutron flow through the containment wall as a whole can be analyzed and recommendations for the thickness of the wall and its composition can be made so as to guarantee the safety of the personnel working at the reactor.

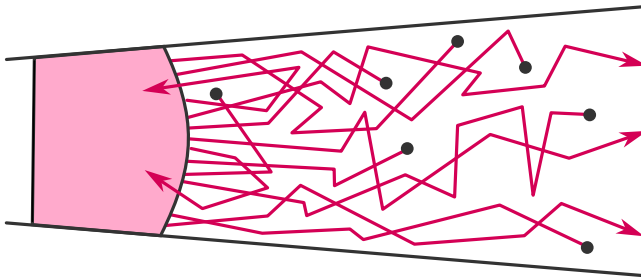


Figure 2.9: A set of all possible trajectories for the neutron.

Modern physics requires the Monte Carlo method on many occasions. Physicists use it to investigate cosmic-ray showers in the Earth’s atmosphere, the behaviour of large flows of electrons in electron discharge devices, and the progress of various chain reactions.

Games and Decision Making

What is the theory of games? Suppose we must make a decision when our objectives are opposed by another party, when our will is in conflict with another will. Such situations are common, and they are called *conflict situations*. They are typical for military actions, games, and every-day life. They often arise in economics and politics.

A hockey player makes a decision that takes into account the current situation and the possible actions of the other players. Every time a chess player makes a decision, he (or she) has to consider the counteraction of the opponent. A military decision should allow for the retaliation of the enemy. In order to decide at what price to sell a product, a salesman must think over the responses of the buyer. In any election campaign, each political party in a capitalist country tries to foresee the actions of the other parties that

are competing for power. In each case, there is a collision of opposing interests, and the decision must be related with *overcoming a conflict*.

Decision making in a conflict situation is hampered by *uncertainty about the behaviour of the opponent*. We know that the opponent will try to act in a way that is least advantageous for us in order to ensure the greatest advantage for himself. However, we do not know to what extent our opponent is able to evaluate the situation and the possible consequences and, in particular, how he evaluates our options and intentions. We cannot predict the actions of the opponent accurately, and the opponent cannot predict our actions. But nonetheless, we both have to make decisions.

Because some way of justifying an *optimal* decision was needed in conflict situations, a new mathematical discipline arose, the *theory of games*. The “game” here is a mathematical model of a conflict situation. Unlike a real conflict, a game has definite rules which clearly indicate the rights and duties of the participants and the possible outcomes of the game (a gain or loss for each participant). Long before the emergence of game theory, simple models of conflicts were used widely. I mean games in the literal sense of the word: chess, checkers or draughts, dominoes, card games, etc. In fact, the name of the theory and the various terms used in it are all derived from these simple models. For instance, the conflicting parties are called players, a realization of a game is a match, the selection of an action by a player (within the rules) is a move.

There are two kinds of move, personal and chance ones. A *personal* move is when the player conscientiously selects an action according to the rules of the game. A *chance* move does not depend on the player’s will: it may be determined by tossing a coin, throwing a die, taking a card from a pack, etc. Games consisting of only chance moves are called *games of chance*, or *games of hazard*. Typical examples are lotteries and bingo. Games with personal moves are called *strategic*. There are strategic games consisting exclusively of personal moves, for instance, chess. There are also strategic games consisting of both personal and chance moves, for instance, certain card games. Let me remark that the uncertainty in games with both personal and chance moves involve both sorts of randomness: the uncertainty of the result of the chance moves and the uncertainty of the opponent’s behaviour in his personal moves.

Game theory is not interested in gambles. It only deals with strategic games. The aim of the game theory is to determine the player’s strategy so as to maximize his chances of winning. The following basic assumption underlies the search for optimal strategies. It is assumed that the opponent is as active and as reasonable as the player, and he or she also takes attempts to succeed.

Naturally, this is not always true. Very often our actions in real conflicts are not as

good as they could be when we assume reasonable behaviour from our adversary; it is often better to guess at the “soft spots” of the opponent and utilize them. Of course, we take a risk when doing so. It is risky to rely too much on the soft spots of the opponent, and game theory does not consider risk. It only detects the most cautious, “safe” versions of behaviour in a given situation. It can be said that game theory gives wise advice. By taking this advice when we make a practical decision, we often take a conscientious risk. E. S. Wentzel writes in *Operations Research*:

“Game theory is primarily valuable in terms of the formulation of the problem, which teaches us never to forget that the opponent also thinks and to take into account his possible tricks and traps. The recommendations following from the game approach are not always concrete or realizable, but it is still useful, while taking a decision, to utilize a game model as one of several possible ones. But the conclusions proceeding from this model should not be regarded as final and indisputable.”

The payoff matrix of a game *Finite two-person zero-sum games* are the best investigated types in game theory. A *two-person* game is a game in which there are exactly two players or conflicting interests. A game is *finite* if both players have a finite number of possible strategies, i.e. a finite number of behaviours. When making a personal move, a player follows a strategy. A *zero-sum* game is a game where the gain by one player equals the loss by the other.

	B_1	B_2	B_3	\dots	B_n
A_1	a_{11}	a_{12}	a_{13}	\dots	a_{1n}
A_2	a_{21}	a_{22}	a_{23}	\dots	a_{2n}
A_3	a_{31}	a_{32}	a_{33}	\dots	a_{3n}
\dots	\dots	\dots	\dots	\dots	\dots
A_m	a_{m1}	a_{m2}	a_{m3}	\dots	a_{mn}

Figure 2.10: Strategies in a finite two-person zero-sum game.

Suppose there is a finite two-person zero-sum game where player A has m strategies

and player B has n strategies (an $m \times n$ game). We use A_1, A_2, \dots, A_m to denote the strategies available to player A and B_1, B_2, \dots, B_n the strategies available to player B . Suppose player A makes a personal move and selects a strategy A_i ($1 \leq i \leq m$), and player B at the same time selects strategy B_j ($1 \leq j \leq n$). We use a_{ij} to denote the gain of player A . Let us identify ourselves with player A and consider each move from his viewpoint. The gain a_{ij} may be either a real gain or a loss (a loss would be a negative gain). The set of gains a_{ij} for different values of i and j can be arranged in matrix form with the rows corresponding to player A strategies and the columns to player B strategies (Figure 2.10). This is called the *payoff matrix* for the game.

Consider the following game. Each player, A and B , writes, simultaneously and independently, one of three numbers 1, 2, or 3. If the sum of the numbers is *even*, player B pays player A the sum, while if the sum is *odd*, A pays it to B . Player A has three strategies: A_1 to write 1, A_2 to write 2, and A_3 to write 3. Player B has the same strategies. The game is a 3×3 one because its payoff matrix contains three rows and three columns. This matrix is given in Figure 2.11(a). Note that a gain by player A of, for instance, -3 is a loss in reality because A pays 3 units to B .

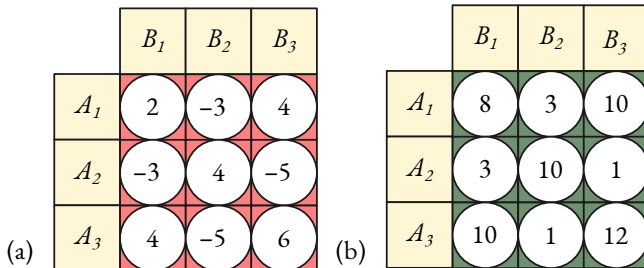


Figure 2.11: A payoff matrix in a 3×3 finite two-person zero-sum game.

Some of the elements are positive and the others are negative in the matrix in Figure 2.11(a). It is possible to make all the elements of the payoff matrix positive by adding some number, say 6, to each element of the matrix. We obtain the matrix in Figure 2.11(b). This matrix is equivalent to the initial one from the viewpoint of analyzing optimal strategies.

The minimax principle Let us analyze the game using the payoff matrix in Figure 2.11(b). Suppose we (player A) pick strategy A_i . Then, depending on the strategy selected by player B , our gain may be either 8 or 3 or 10. Thus, strategy A_1 yields a gain of 3 in the worst case. If we choose either A_2 or A_3 , the worst gain is 1. Let us write down the minimum possible gains for each strategy A_i as an additional column in the

payoff matrix (Figure 2.12). It is clear that we should choose a strategy whose *minimum possible gain is greatest* (as compared with the other strategies). This is strategy A_1 in this case. Three is the largest one out of the minimum gains for each strategy (viz. 3, 1, and 1). This is called the *maximin gain*, or the *maximin*, or just the *maxim*. It is also sometimes called the *lower value of the gain*. Thus, if we select the maximin strategy (strategy A_1 in this case), our gain is guaranteed to be, whatever the behaviour of the opponent, at least the lower value of the game (a gain of 3 in this case). The opponent will reason in a similar way. If he selects strategy B_1 , he will have to give us a gain of 10, which is his worst case.

The same can be said of strategy B_2 . Strategy B_3 yields the worst case for the opponent corresponding to a gain of 12 for us. Numbers 10, 10, and 12 are the maximum values of our gains corresponding to the opponent's strategies B_1 , B_2 , and B_3 , respectively. Let us write these values as a row in the payoff matrix (see Figure 2.12). It is clear that our opponent should select the strategy which *minimizes* our *maximum possible gain*. This is either strategy B_1 or B_2 . Both strategies are minimax ones and both guarantee that our opponent limits our gain to the *minimax*, or, in other words, the *upper value of the game* is 10.

Our maximin strategy and the minimax strategy of the opponent are the most cautious "safe" strategies. The principle of being cautious dictating that the players select such strategies is called the *minimax principle*.

Now let us return to the matrix in Figure 2.12 and try some reasoning. The opponent has two minimax strategies, B_1 and B_2 . Which strategy should he choose? If he knows that we are cautious and have selected the maximin strategy A_1 he would not select strategy B_1 because this would yield a gain of 8. Therefore, it is likely that he would choose strategy B_2 , and our gain would then be 3. But if we perceived our opponent's ideas correctly, shouldn't we take a risk and choose strategy A_2 ? If the opponent then selects strategy B_2 , our strategy A_2 will give us a gain of 10. However, our deviation from the minimax principle may cost us dearly. If the opponent is even cleverer and reasons in a similar way, he would answer our strategy A_2 with strategy B_3 rather than B_2 . And then, instead of a gain of 10, we would only gain 1.

	B_1	B_2	B_3	
A_1	8	3	10	3
A_2	3	10	1	1
A_3	10	1	12	1
	10	10	12	

Figure 2.12: A payoff matrix in a 3×3 finite two-person zero-sum game.

Does this mean that game theory only recommends we adhere to a minimax (maximin) strategy? It depends on whether the payoff matrix has a *saddle point*.

A game with a saddle point. Consider the 3×3 game, whose payoff matrix is given in Figure 2.13. Here both the maximin and minimax gain 4. In other words, the lower and the upper value of the game coincide and both are equal to 4. A gain of 4 is simultaneously the maximum of the minimum gains for strategies A_1 , A_2 , and A_3 and the minimum of the maximum gains for strategies B_1 , B_2 , and B_3 . In geometry, the point on a surface which is at the same time a minimum along one coordinate axis and a maximum along the other is called a saddle point. Point C on the surface in Figure 2.13 is a *saddle point*. It is the maximum along the x -axis and the minimum along the y -axis. It is easy to see that the surface in the vicinity of this point is actually like a saddle. Just as in geometry, element $a_{22} = 4$ of the payoff matrix in question is called the *saddle point of the matrix*, and the game is said to have a saddle point.

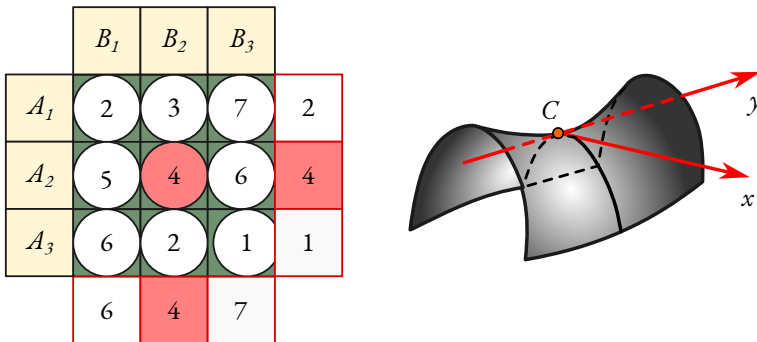


Figure 2.13: A 3×3 game with saddle point.

We need only look through the matrix in Figure 2.13, to see that each player should adhere to his maximin (minimax) strategy. These strategies are optimal in a game with a saddle point. Any deviation from them will be disadvantageous for the player who took the risk.

However, if a game does not have a saddle point (see the matrix in Figure 2.11), neither of strategies A_i or B_j is optimal.

The necessity of a random change of strategy in a game without a saddle point. Suppose that we and our opponent repeatedly play the game whose matrix is given in Figure 2.11. If we choose a definite strategy, for instance, the maximin strategy A_1 , and adhere to it turn after turn, our opponent will see it and select strategy B_2 each

time, so that our gain will not exceed the lower value of the game, i.e. it will equal 3. However, if we suddenly (for the opponent) choose strategy A_2 instead of A_1 , we receive a gain of 10. Having guessed our new strategy (naturally, if we later adhere to it), our opponent will go from strategy B_2 to strategy B_3 right away, thus decreasing our gain to 1. And so forth. We can see here a general rule for games without a saddle point: a player using a *certain strategy* will be worse off than a player who *changes strategy at random*.

However, the random changes in strategies should be done wisely rather than haphazardly. Suppose A_1, A_2, \dots, A_m are the possible strategies of player A (see Figure 2.10). To obtain the greatest benefit, the strategies should be chosen at random but with different (specially calculated) probabilities. Suppose strategy A_1 is used with probability p_1 , strategy A_2 with probability p_2 etc. Player A is now said to have a *mixed strategy* $S_A(p_1, p_2, \dots, p_m)$. Unlike S_A , the A_j strategies are called *pure strategies*. By correctly selecting the probabilities p_j a mixed strategy may be *optimal*. The gain of player A will then be no less than a certain value ν called the *value of the game*. This value is greater than the lower value of the game, but less than the upper one.

Player B should behave in a similar manner. His optimal strategy is also a mixed strategy. Let us designate it $S_B(q_1, q_2, \dots, q_n)$, where q_j are specially selected probabilities with which player B uses strategies B_j . When player B selects an optimal mixed strategy, the gain of player A will be no more than game value ν .

The search for an optimal mixed strategy. Let us use $S_A(p_1, p_2, \dots, p_m)$ to denote an optimal mixed strategy for player A . We must now find probabilities p_1, p_2, \dots, p_m and calculate the game value ν once the payoff matrix of the game is known (see Figure 2.10). Suppose player B selects pure strategy B_1 . Then the average gain of player A will be $a_{11}p_1 + a_{21}p_2 + \dots + a_{m1}p_m$. This gain should be no less than the game value ν , and hence

$$a_{11}p_1 + a_{21}p_2 + \dots + a_{m1}p_m \geq \nu.$$

If player B selects strategy B_2 , the average gain of player A should also be no less than the game value ν , and hence

$$a_{12}p_1 + a_{22}p_2 + \dots + a_{m2}p_m \geq \nu.$$

Whichever strategy player B chooses, the gain of player A should always be no less than the game value ν . Therefore, we can write the following system of n inequalities (recall that n is the number of B 's pure strategies):

$$\left. \begin{aligned} a_{11}p_1 + a_{21}p_2 + \dots + a_{m1}p_m &\geq \nu, \\ a_{12}p_1 + a_{22}p_2 + \dots + a_{m2}p_m &\geq \nu, \\ \dots \quad \dots \quad \dots \quad \dots, \\ a_{1n}p_1 + a_{2n}p_2 + \dots + a_{mn}p_m &\geq \nu. \end{aligned} \right\} \quad (2.10)$$

Recall that

$$p_1 + p_2 + \dots + p_m = 1. \quad (2.11)$$

Introducing designations $x_1 = p_1/\nu$, $x_2 = p_2/\nu$, ..., $x_m = p_m/\nu$ we can rewrite (2.10) and (2.11) as

$$\left. \begin{aligned} a_{11}x_1 + a_{21}x_2 + \dots + x_{m1}p_m &\geq 1, \\ a_{12}x_1 + a_{22}x_2 + \dots + a_{m2}x_m &\geq 1, \\ &\dots \dots \dots \dots, \\ a_{1n}x_1 + a_{2n}x_2 + \dots + a_{mn}x_m &\geq 1. \end{aligned} \right\} \quad (2.12)$$

$$x_1 + x_2 + \dots + x_m = \frac{1}{\nu}. \quad (2.13)$$

It is desirable that the game value ν should be as large as possible, and hence $1/\nu$ should be as low as possible. Therefore, the search for the optimal mixed strategy is thus reduced to the solution of the following mathematical problem: find non-negative values x_1, x_2, \dots, x_m such that they meet inequalities (2.12) and minimize the sum $x_1 + x_2 + \dots + x_m$.

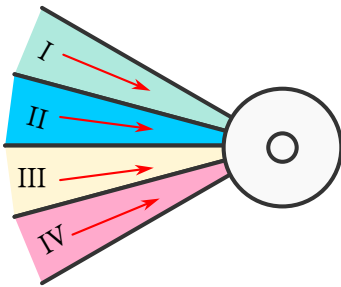


Figure 2.14: Strategies with aeroplanes and anti-aircraft guns.

Airplanes against anti-aircraft guns.

Let us find the optimal mixed strategy for a concrete game. Suppose “player” A wants to attack “player” B . A has two airplanes each carrying a large bomb. B has four anti-aircraft guns defending an important military base. To destroy the base, it is sufficient for at least one airplane to approach it. To approach the base the airplanes may choose one of four air corridors (Figure 2.14, where 0 is the base and I, II, III, and IV are the air corridors). A may send both airplanes along the same corridor or along different corridors. B may place his four anti-aircraft guns to cover the corridors in

different ways. Each gun can only shoot once, but it will hit the airplane if it is in that corridor.

A has two pure strategies: strategy A_1 to send the airplanes along different corridors (no matter which ones), and A_2 , to send both airplanes along the same corridor. B ’s strategies are B_1 to put an anti-aircraft gun into each corridor, B_2 to put two guns into two corridors (leaving the other two corridors unprotected), B_3 to put two guns into

one corridor and one gun into two of the other corridors, B_4 to put three guns into a corridor and one gun into another corridor, and B_5 to put all four guns into one corridor. Strategies B_4 and B_5 are certainly bad because three or four guns in a single corridor are not needed, since A only has two airplanes. Therefore, we need only discuss strategies B_1 , B_2 , and B_3 .

Suppose A chooses strategy A_1 and B chooses strategy B_1 . It is clear that neither airplane will reach base: the A 's gain will be zero ($a_{11} = 0$). Suppose strategies A_1 and B_2 are chosen. Let us assume that the guns are in corridors I and II. If the aircrafts are flying along different corridors, then six variants are equally probable: they fly along corridors I and II, along corridors I and III, along corridors I and IV, along II and III, along II and IV, or along III and IV. In only one of the six cases will neither plane reach the base (when they fly along corridors I and II). Whichever corridor B chooses to place his guns in, airplanes will always have six equally probable variants and only one does not yield a winning move. Therefore, if strategies A_1 and B_2 are chosen, the probable gain for A will be $5/6$ ($a_{12} = 5/6$). Reasoning in the same manner, it is easy to find the rest of the elements of the payoff matrix for this game. The resultant 2×3 matrix is shown in Figure 2.15. Note that the elements of the matrix are *probable* gains; so here even the pure strategies involve chance. The lower value of the game is $1/2$, and the upper one is $3/4$. The maximin strategy is A_2 while the minimax strategy is B_3 . There is no saddle point, and the optimal solution for the game will be a mixed strategy.

	B_1	B_2	B_3
A_1	0	$5/6$	$1/2$
A_2	1	$1/2$	$3/4$

Figure 2.15: Matrix of probable with aeroplanes and anti-aircraft guns.

In order to find the optimal mixed strategy, let us use the payoff matrix and relations (2.12) and (2.13). The relations for this case are

$$x_2 \geq 1, \quad \frac{5}{6}x_1 + \frac{1}{2}x_2 \geq 1, \quad \frac{1}{2}x_1 + \frac{3}{4}x_2 \geq 1, \quad (2.14)$$

$$x_1 + x_2 = \frac{1}{\nu}. \quad (2.15)$$

The solution can be conveniently represented as a diagram. We plot the positive values x_1 and x_2 along the coordinate axes (Figure 2.16). The first inequality in (2.14) corresponds to the area above the straight line CC ; the second inequality is the area above DD ; and the third inequality in (2.14) is the area above EE . All three inequalities are satisfied inside the area shaded red in the figure. The equation $x_1 + x_2 = \text{const}$ defines a family of straight

lines, some of which are shown in figure as dash lines. The straight line FF has the least sum $x_1 + x_2$ of all the lines in the family with at least one point within the red area. Point G indicates the solution corresponding to the *optimal mixed strategy*. The coordinates of this point are $x_1 = 3/5$ and $x_2 = 1$. Hence we find $\nu = 5/8$, $p_1 = 3/8$, and $p_2 = 5/8$. Thus, A 's optimal mixed strategy would be to use strategy A_1 with probability $3/8$ and strategy A_2 with probability $5/8$.

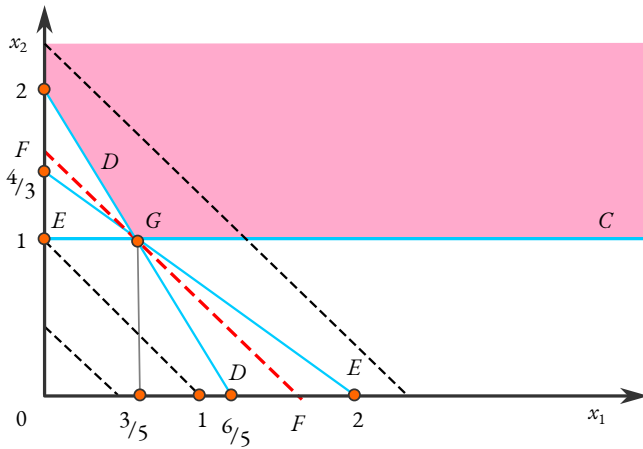


Figure 2.16: Solution for the game of aircrafts and anti-aircraft guns.

How could we use this recommendation in practice? If there is only *one* bombing raid in the “game”. A clearly should select strategy A_2 because $p_2 > p_1$. Suppose now the game has *many* raids (for instance, raids on many bases). If the game is run N times ($N \gg 1$), then A should choose strategy A_1 $3N/8$ times and strategy A_2 $5N/8$ times.

We have so far only discussed the behaviour of A , allowing B to act arbitrarily. If A selects his optimal mixed strategy, his average gain will be between the upper game value of $3/4$ and the game value $\nu = 5/8$. If B behaves unreasonably, the A 's gain may rise to the upper value of the game (or even greater). However, if B in turn adheres to his optimal mixed strategy, the A 's gain will equal the game value ν . The optimal mixed strategy for B precludes his use of strategy B_3 and is to use strategy B_1 with probability $1/4$ and strategy B_2 with probability $3/4$. That strategy B_3 should not be used can be seen from Figure 2.16: the straight line EE corresponding to this strategy does not have any points in the red area. To determine the probabilities with which to apply strategies B_1 and B_2 , we use the game value $\nu = 5/8$, and get $q_1 \times 0 + (1 - q_1) \times 5/6 = 5/8$. It is

clear from this that $q_1 = 1/4$ and $q_2 = 1 - q_1 = 3/4$.

Chapter 3

Control and Self-control

Cybernetics penetrated and continues to penetrate every area of man's work and daily life. This is the science of the optimal control over complex processes and systems.

A.I. Berg

The Problem of Control

Control against disorganization. Although the world around us is full of chance, it nonetheless proves to be organized and ordered in many ways. *The disorganizing effect of chance is countered by the organizing influence of control and self-control.*

Suppose an airplane flies from Moscow to Leningrad. Various random factors affect it during the flight. Therefore, all three space coordinates of the airplane are random functions of time. The flight trajectory is a realization of these random functions. However, these “subtleties” do not bother the passengers; they fasten their belts before takeoff confident that whatever thunderstorms might occur on the way and whichever winds affect the airplane, it will arrive at Leningrad airport. The basis for this confidence lies in the aircraft's control system and the actions of the pilot. We met queueing systems above, and although there is a great deal of chance, they comply with their objectives. This is because the organization of the system and the control of its operation is well-designed.

Controls take on a variety of guises. Suppose we want a set of books to serve public

for a long time. This is impeded by chances both purely physical in nature and those related to the attitudes of some readers. So we control matters: we take care of the binding, regulate the temperature, humidity, and illuminance in the rooms where the books are stored, give the book a library card, and set up the rules governing the use of the books.

No one is safe from disease, and although each disease has a definite cause, the prevalence and lethality of a disease on the scale, say, of a town is governed by chance. When fighting it, we must control matters by improving working and living conditions, taking preventive medical measures, constructing stadiums, swimming pools, sport complexes, ordering pharmacies to supply the necessary drugs, etc.

Thus, there is a *confrontation* of two powerful factors in the world, two basic trends. On the one hand, there is *randomness*, a tendency to disorganization, disorder, and destruction in the long run. On the other hand, there is *control* and self-control, a tendency to organization, order, development, and progress.

Choice as a prerequisite of control. If all the processes and phenomena in the world were strictly predetermined, it would be meaningless even to speak of the possibility of control. *In order to control something, there must be some choice.* How may we make a decision if everything is predetermined in advance? Every phenomenon must have several probable lines of development. One may say that a world built on probability is the only world in which control is possible.

Control acts against chance, even though the possibility of control is brought about by the existence of chance. It is random occurrences that help us avoid predetermination. We can say that randomness “brings to life” its own “grave-digger”, i. e. control. This is a manifestation of the dialectic unity of the necessary and the random in the real world.

Control and feedback. Two different control schemes are shown in Figure 3.1, where S is the controlled system, CU is the control unit, V is the input to the controlled system (the control signal), P are random perturbations affecting the controlled system, and w is the final output from the system. Scheme (b) differs from scheme (a) in having a feedback loop, that is the control unit receives information about the results of control.

What is feedback for? In answering this question, let me remark that the “relationship” between randomness and control is one of active confrontation. Control acts against chance, and chance acts against control. The latter fact requires flexible control, the possibility for adjustment. The control unit must be able continuously to receive data about the results of the control and correct its signals to the system appropriately.

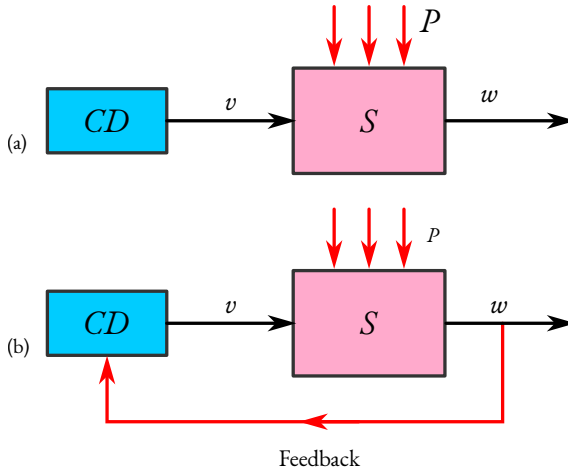


Figure 3.1: Two different control schemes.

In point of fact, any real control system supposes the presence of a feedback loop.

Control without feedback is not only ineffective, it is actually unviable.

Take for example someone driving a motor-car. Imagine for a minute that the feedback suddenly disappeared, that is, the driver stopped attending to the motion of the car. The car would continue to be controlled, but without any feedback. The car is immediately affected by a variety of random events. A small bump or bend in the road, a car moving in the opposite direction all are random and could lead to an accident in only a few seconds.

The control algorithm. Now what should be done and how should the system be controlled? It depends on the situation and the goal being pursued. In fact the answer lies in the algorithm of control.

A control algorithm is a sequence of actions that must be carried out to reach a set of goals.

In the example with the car and a driver, the control algorithm contains rules on how to start the engine, how to brake, how to turn, how to shift gears, and so on. The algorithm also contains the traffic regulations and good driving practice.

In some cases the control algorithm is simple. For instance, in order to use a coffee

machine, only the following two actions need be carried out: put a coin in the slot, and press the appropriate buttons. This is the complete control algorithm for this machine. In other cases, the control algorithm is much more complicated. For instance, it is more difficult to drive a car, while flying a jet is even more complicated. In very complicated cases, the control algorithm cannot even be defined in full. For instance, complete control algorithms for managing a large enterprise or industry simply do not exist.

From the “Black Box” to Cybernetics

Despite the diversity of algorithms, the processes of control can be investigated from general positions, irrespective of the details of the considered system. A typical example is the simulation of a system using the “black box” model.

What is a “black box”? Suppose we consider a controlled system, where V_1, V_2, \dots, V_m are its inputs (control signals), P is a random perturbation, and W_1, W_2, \dots, W_n are its outputs (Figure 3.2). Now let us suppose that we do not know or do not care what is inside the system. We only need investigate the relationships between the inputs (V_1, V_2, \dots, V_m) and the outputs (W_1, W_2, \dots, W_n). It is said in this case that the given system is a “black box”.

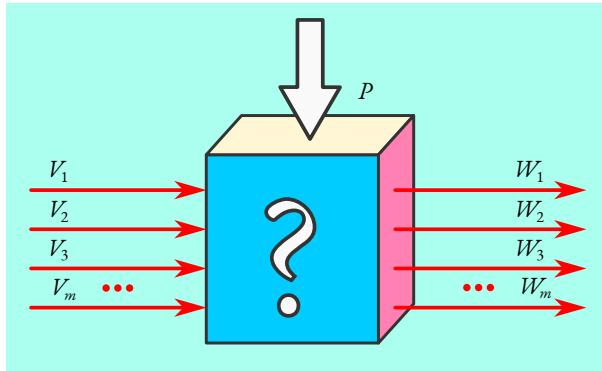


Figure 3.2: A black box is a system whose inputs and outputs are known, but internal structure is not.

Any controlled system is a “black box” if its internal structure is not considered, and only the responses of the outputs to the inputs are investigated.

Man surrounded by black boxes. The advance of science and technology has surrounded mankind by a vast number of controlled systems. As a rule, we are not a bit bothered by this because we quickly get accustomed (sometimes unconsciously) to considering these systems as black boxes. We find out how, what, and where to turn, press, or switch the buttons to obtain the desired effect. If you want to watch a TV show, there is no need to know the structure or workings of a television. We need only press the proper button and select the channel. To make a telephone call, we do not have to be telephone engineers; we just pick up the receiver, wait for the call signal, and dial the telephone number. We use television, telephone, and many other systems and consider them to be black boxes. Naturally, we could learn what is inside the system and how it works if we want to, but in our modern world we often think it's a waste of time to study what we can quite do without in practice. More and more often we prefer to use black boxes and when they fail we call in a professional technician.

We should recognize the validity of the complaints that as modern people we have become less curious, that we do not want to see things in depth because there are too many things to see, and it is not difficult to use them. However, I should not make things appear to be worse than they are. Firstly, there is a system of universal secondary education at least in the developed countries, which ensures each person has a basic minimum knowledge. Secondly, from the viewpoint of the development of society, the knowledge available to the society as a whole is more important than what a single person may know.

Complex systems as black boxes. Modern systems are becoming more and more sophisticated as their functional capacities become more and more diverse. Naturally, the more we need to know about the functions of a system, the further we push our investigation of its inner structure into the background, and in many cases such a total investigation would prove infeasible because of the complexity of the system.

This shift of emphasis leads us to a qualitatively new viewpoint, in which the main aim is to investigate control and self-control as general processes irrespective of the concrete devices comprising the systems. This point of view brings about *cybernetics* as the science of control (self-control) in complex systems.

Curiously this point of view reveals an interesting fact and makes us look at the black-box model in another way. It turns out that we do not need understand every structural subtlety of a complex system, indeed its separation into component parts can obscure essential information. The black-box model becomes *fundamental* as the only acceptable way of analyzing a complex system.

What is cybernetics? The science of cybernetics was founded by the American

scientist Norbert Wiener (1894-1964) and dates from 1948 when he published his famous book *Cybernetics, or Control and Communication in the Animal and the Machine*. Wiener wrote:

“We have decided to call the entire field of control and communication theory, whether in the machine or in the animal, by the name cybernetics, which we form from the Greek κυβερνητής, or steersman.”

It should be noted that the term “cybernetics” was not new. Plato used it meaning the art of controlling ships. The French physicist Ampère classified sciences in the first half of the 19th century and placed a science, which was the study of the methods of government, in section 83. Ampère called this science cybernetics. Today we only use the term “cybernetics” in the sense given to it by Wiener.

Cybernetics is the science of the control and communication in complex systems, be they machines or living organisms.

The Soviet scientist L.A. Rastrigin wrote a book called *This Chancy, Chancy, Chancy World* (Mir Publishers, Moscow, 1984), in which he remarked:

“Until cybernetics made its appearance, control processes in an electric generator were investigated by electrical engineering, control of the motion of a clock pendulum (in effect a swing) was dealt with in mechanics, and control of population dynamics in biology. Norbert Wiener was the first to point to the universal nature of control and to show that the organizing of an object (the lowering of its entropy) could be achieved by means of standard procedures, that is, by applying the methods of cybernetics independently of the physical characteristics of the object.”

L.A. Rastrigin imaginatively calls cybernetics a science which fights randomness, thus emphasizing the idea of control counteracting disorganization and destruction caused by diverse random factors.

Cybernetics and robots. One of the central topics of cybernetics concerns *process automation*, in particular, *self-control in complex systems*. Investigations into this area resulted in the appearance of a discipline called “robotics”. Modern cybernetics literature discusses the possibility of designing automata that can reproduce and teach themselves. Artificial intelligence is also a topic being investigated. The following questions are being studied: Is the machine capable of creativity? Could a machine become cleverer than its designer? Could the machine think?

The more sophisticated types of robots are still in the realms of science fiction, although we often hear discussions about the possibilities of robotics, or rather whether

artificial “men” might be possible. The layman now seems to believe that cybernetics is indeed simply the science of robots, automata, or thinking machines. The true purpose of cybernetics as the science of control is now masked by the fantastic technological promise.

True, cybernetics does include the problems of automation, and thus contributes to scientific and technological progress. The automation of various processes, the design of automatic Lunar explorers, automatic space docking are all achievements of cybernetics. Cybernetics also investigates computer creativity and artificial intelligence. However, this is not so as to evolve an artificial person. When we programme computers to “compose” music or “write” a poem or play chess or give a “talk”, we are attempting to simulate creativity and so find out more about these processes. It could be said that we are investigating the limit of computer abilities, but not that we want to substitute them for human beings in the future: we just want to understand several important topics thus making it possible to go deeper into the control processes occurring in human beings. The reader should remember this and not consider cybernetics to be just the “science of robots”.

We may now start discussing the central notion of cybernetics, i. e. *information*. Let me say right away that cybernetics investigates control and self-control primarily from the viewpoint of information. It investigates the collection, conversion, transmission, storage, and retrieval of information. In a certain sense of the word, cybernetics can be regarded as the “science of information”.

Information

Let me begin with an excerpt from the immortal poem *De Rerum Natura* (On the Nature of Things) by Carus Lucretius (ca. 99-55 B.C.):

“...if things came to being from nothing,
Every kind might be born from all things,
Nought would need a seed.
First men might arise from the sea, and from the land,
The race of scale creatures, and birds burst forth
The sky. Cattle and other herds, and all the tribe
Of wild beasts with no law of birth,
Would haunt tilth and desert ...”

It is interesting that there is here a hint of the conservation of not only matter and energy, but also of something else, which is neither matter nor energy. There is no shortage of

energy and matter in the sea, but people do not appear in the sea. Nor too does the dry land produce fish. Truly, “if things came to being from nothing, ... nought would need a seed”. In the modern terminology of science, we might say that this is a hint of the *conservation of information*. The information needed by plants and animals to live and reproduce cannot appear “from nothing”. It is stored in “seeds” and thus handed down from generation to generation.

The term “information” is now encountered everywhere in science and everyday life. In fact, every activity is related to the *collection, conversion, transmission, storage, and retrieval of information*. We live in a world filled with information, and our very existence is impossible without it. Academician A.I. Berg once said: “Information penetrates every pore of the life of human beings and their societies ... Life is impossible in a vacuum of either mass-energy or information.”

The bit, the unit of information. What is information? What units is it measured in? Let us start with a simple example. A train approaches a station. By remote control, a signalman can switch a train from one track (*A*) to another (*B*). If the switch is up, the train goes along track *A*, and if it is down, the train goes along track *B*. Thus, the signalman, by moving the switch up or down, is sending a control signal containing 1 bit of information. The word “bit” is an abbreviation of “binary digit”.

To see what we mean by “binary digit”, recall how digits are used to write numbers. We commonly use the decimal number system, i.e. a system with ten digits (0, 1, 2, ..., 9). Take a number written in the *decimal* system, say 235. We say “two hundred and thirty five” and, as a rule, do not pause to think that this means the sum of two hundreds, three tens, and five units, i. e. $2 \times 10^2 + 3 \times 10 + 5 \times 10^0$. The same number (235) can also be in the *binary* system, which only has two digits, 0 and 1, as 11101011, which means $1 \times 2^7 + 1 \times 2^6 + 1 \times 2^5 + 0 \text{ times } 2^4 + 1 \times 2^3 + 0 \times 2^2 + 1 \times 2^1 + 1 \text{ times } 2^0$. Since $2^7 = 128$, $2^6 = 64$, $2^5 = 32$, $2^3 = 8$, $2^1 = 2$, and $2^0 = 1$, we have our number $128 + 64 + 32 + 8 + 2 + 1 = 235$. Any number can be written in either the decimal or the binary system. If you don’t follow this explanation try looking at Figure 3.3.

Let us return to the railway example. Remember we have two choices: the switch is either up (track *A*) or down (track *B*). We could write the digit 0 for switch up and digit 1 for switch down. It can be said that the control signal can thus be coded by one of the two binary digits, zero or unity. The signal thus contains one binary digit, or 1 bit of information.

Consider a more interesting example. The railway lines near a station are shown in Figure 3.4.

The railway switches are labelled by the letters *a*, *b*, *c*, *d*, *e*, *f*, and *g*. If a switch

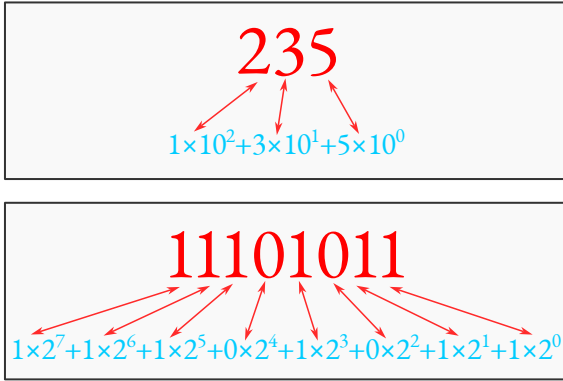


Figure 3.3: Representing the number 235 in decimal and binary systems.

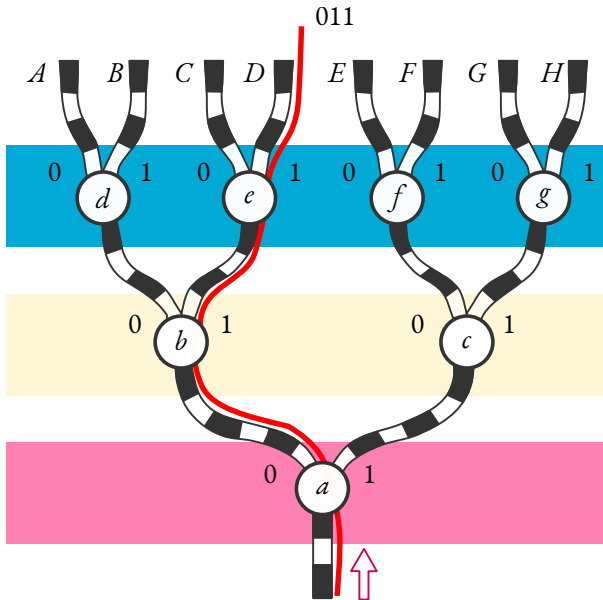


Figure 3.4: Controlling railway lines by using switches.

receives a control signal of 0, it opens the left-hand track, and if it receives a signal of 1, it opens the right-hand track. The signalman has three control switches: the first one

sends a signal (0 or 1) to railway switch a , the second one sends a signal simultaneously to switches b and c , and the third one simultaneously to switches d , e , f , and g . The station has eight tracks: A, B, C, D, E, F, G , and H . To send a train along track A , all three control switches must be turned to the 0 position, i.e. send the three-digit signal 000. To direct a train to track B , it is necessary to send the three-digit signal 001. Each track thus has its own three-digit signal, i.e.

A	B	C	D	E	F	G	H
000	001	010	011	100	101	110	111

We see that to select one of the eight outcomes requires a set of elementary signals, each of which carries 1 bit of information. Therefore, to choose a track in this example requires three bits of information.

Thus, in order to select one option out of two, 1 bit of information is required; in order to select one option out of eight, 3 bits of information are required. In order to select one of N options, I bits of information are required, where

$$I = \log_2 N. \quad (3.1)$$

This is the *Hartley formula*. It was suggested in engineer Ralph Hartley, who was interested information.

The Bar Kohba game. A rebellion against Romans broke in 135 A. D. in the ancient Judea led by one Bar Kohba. As the legend has it, Bar Kohba sent a spy into the camp of Romans, and the spy discovered a great deal before being caught. He was tortured and his tongue was cut out. However, the spy managed to escape, but without his tongue he could not report what he had found out in the enemy's camp. Bar Kohba resolved the problem by asking the spy questions that only required a "yes" or "no" answer (it was only necessary to nod or shake the head). Bar Kohba was able to obtain all the information he wanted from his spy, even though the spy had no tongue.

A similar situation is described in *Le comte de Monte Christo* by Alexandre Dumas père. An old man in the novel had been paralyzed and could neither speak nor move his hands. Nonetheless, his relatives were able to communicate with him asking him questions which required only a "yes" or a "no". If "yes", the old man would close his eyes; if he blinked several times, it was "no".

It turns out that any information can be transmitted in the form of "yes" and "no" answers if the questions are constructed *properly*. This idea underlies the *Bar Kohba game*, which first appeared at the turn of the century in Hungary and then spread

to other countries. A player thinks of something. He may, for instance, make a wish or even think up a sentence. The other player must guess the wish or sentence by asking questions, which must be honestly answered. However, the questions may only require a “yes” or “no” answer. The quantity of information needed for a correct guess can be measured by the number of questions, given that the most rational method of interrogation is used. Each answer can be enciphered by a binary digit, for instance, we could use a one for a “yes” and a zero for a “no”. Then the information needed for a correct guess would be a combination of zeroes and unities.

Let us play a Bar Kohba game with the railway signalman at the station whose tracks are given in Figure 3.4. The signalman thinks of a track along which a train should travel to the station. We want to guess the track. The game would go as follows.

QUESTION: Should switch *a* open the track on the right?

ANSWER: No. (let us cipher this answer by digit 0).

QUESTION: Should switch *b* open the track on the right?

ANSWER: Yes (we cipher: 1).

QUESTION: Should switch *c* open the track on the right?

ANSWER: Yes (we cipher: 1).

Having asked these three questions, we see that the signalman decided on track *D*. The information needed to answer was the chain of answers “no-yes-yes” or, in other words, by the set of binary digits 011. We know that the information capacity of the signalman’s “riddle” was three bits long. Each of the signalman’s three answers contained one bit of information.

Let me cite one more example of the Bar Kohba game. There are 32 pupils in a class. The teacher decides on one of them. How can we find out which one? Let us take the class register, in which the surnames of all the pupils are listed in alphabetical order and enumerated. Let us start asking questions.

QUESTION: Is the pupil among those listed from 17 to 32?

ANSWER: Yes (we cipher: 1).

QUESTION: Is the child among those listed from 25 to 32?

ANSWER: No (0).

QUESTION: Is the child among those listed from 21 to 24?

ANSWER: No (0).

QUESTION: Is the child among those listed either 19 or 20?

ANSWER: Yes (1).

QUESTION: Is it number 20?

ANSWER: No (0).

Consequently, the teacher meant pupil number 19 in the class register. This information required the chain of answers “yes-no-no-yes-no” or, in other words, the set of binary digits 10010. It is clear from Figure 3.5 that the area in which the surname was searched for gradually decreased with each answer. To solve the problem, it only required to ask five questions. According to the Hartley formula, the selection of the option out of 32 requires $\log_2 32 = 5$ bits of information. Therefore, each of the answers in this game contained 1 bit of information.

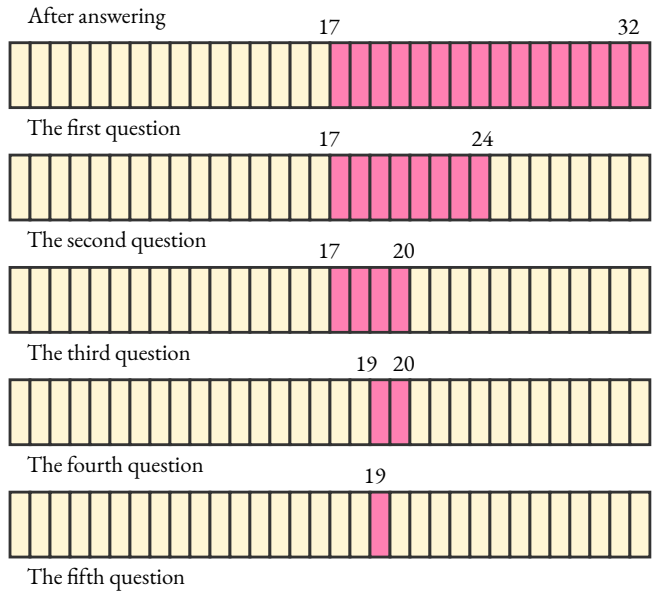


Figure 3.5: Finding out the selected pupil from a group of 32.

Perhaps I have created the impression that each answer in the Bar Kohba game always contains 1 bit of information. It is easy to see that this is not so. Suppose that we established that a surname was listed from 17 to 32 and then ask: Is the surname listed from 9 to 16? It is clear that the answer to this question must be negative. The fact that the answer is obvious means that it does not contain any information at all. Naturally, we might have a situation without “silly” questions.

QUESTION: Is the surname listed from 1 to 8?

ANSWER: No.

QUESTION: Is it listed from 25 to 32?

ANSWER: No.

QUESTION: Is it listed from 9 to 16?

ANSWER: No.

QUESTION: Is it listed from 17 to 24?

ANSWER: Yes.

QUESTION: Is it listed either 23 or 24?

ANSWER: No.

QUESTION: Is it listed either 19 or 20?

ANSWER: Yes.

QUESTION: Is it listed 19?

ANSWER: Yes.

Having chosen this strategy, we extracted the needed information using eight questions rather than five. The quantity of information in the final answer equals 5 bits as before. Therefore, each individual answer in this case contained, on the average, $5/8$ bit of information.

Thus, we see that “yes-no” answers do not always contain 1 bit of information. Running ahead of ourselves, we can note that 1 bit is the *maximum* information that such an answer may contain.

“Just a minute,” you might say, “if this is so, does then a binary digit not always carry one bit of information?”

“Quite true,” I would answer.

“Then how about the definition of a bit of information given above? Can we use the Hartley formula?”

All that has been said about a bit of information (and about the Hartley formula) remains valid, although with a reservation that *every option should be equally probable*. I did not want to discuss this topic too early, but now the time has come to do so.

Information and probability. The Shannon formula. I have emphasized that control is only possible in a world where necessity is dialectically confronted with chance. In order to control something, there must be choice. Any situation we want to control carries with it uncertainty. This uncertainty can be compared with a shortage of information. While we control an object, we introduce information and thus decrease the uncertainty.

For instance, a train may arrive along any of the eight tracks in our example above, so there is uncertainty. By sending a control signal with three bits of information, the signalman eliminates this uncertainty, and the train is directed along one particular track. The teacher could have thought of any of his 32 pupils, so there was uncertainty which surname had been chosen. Having listened to the answers for a number of questions with an overall quantity of information of five bits, we can eliminate this uncertainty and identify the pupil.

Now let us return to the starting point of our reasoning and to the presence of choice. Until now, we assumed that each option was equally probable. The signalman could have chosen any of the eight tracks with equal probability. The teacher could have picked anyone of his 32 pupils. However, we often have to choose between options that are not equally probable, and then it is necessary to pay due attention to the *probability associated with each option*. Suppose the answer to a question may be either “yes” or “no” and both outcomes are equally probable. The answer then will carry precisely 1 bit of information. However, if the “yes” or “no” outcomes have different probabilities, then the answer will contain less than 1 bit of information. And the greater the difference between the probabilities of the two outcomes, the smaller the quantity of information. In the limit of the probability of a “yes” (or a “no”) being unity, the answer will not contain any information at all.

Now, let us look at what happens when different outcomes (different options) have different probabilities. I do not want to cram this book with mathematics, so I shall only discuss the basic results. Suppose ξ is a random discrete variable that may assume the values $x_1, x_2, x_3, \dots, x_N$ with probabilities $p_1, p_2, p_3, \dots, p_N$, respectively. We have N outcomes (N different values of the random variable) which appear with different prob-

abilities. Given an observation of the variable ξ and its value, how much information does this observation carry?

This problem was investigated by the American scientist Claude Shannon in the mid-1940s. He came to the conclusion that we obtain the quantity of information equal (in bits) to

$$I(\xi) = \sum_{i=1}^N p_i \log_2 \frac{1}{p_i}. \quad (3.2)$$

This is a fundamental relation in information theory. It is called the *Shannon formula*.

Suppose that the outcomes are equally probable, and the random variable may take on the values x_i with the same probability p . This probability is clearly $1/N$ and so from (3.2) we obtain

$$I = \frac{1}{N} \sum_{i=1}^N \log_2 N = \frac{1}{N} N \log_2 N = \log_2 N,$$

i.e. the Hartley formula (3.1). Consequently, we see that the Hartley formula is a special case of the Shannon formula when all outcomes are equally probable.

Using the Shannon formula, let us find how much information can be contained in a “yes” or “no” answer. Suppose p is the probability of a “yes”. Then the probability of a “no” answer is $1 - p$. According to (3.2), the information obtained from the answer to a question is

$$I = p \log_2 \frac{1}{p} + (1 - p) \log_2 \frac{1}{1 - p}. \quad (3.3)$$

The graph of I versus p , as defined by (3.3), is given in Figure 3.6.

Maximum information (1 bit) is obtained when $p = 1/2$, i.e. when a “yes” and a “no” are equally probable. Now we can refine our notion of “1 bit of information”.

This is the information contained in a digit that may take on only two values provided both values are equally probable.

It follows that the best strategy in the Bar Kohba game is to ask “yes” or “no” questions, the answers to which are nearly or equally probable. Recall the question: “Is the surname listed from 17 to 32?” Here the answers “yes” and “no” are equally probable because there are 32 pupils and the numbers from 17 to 32 cover half of the pupils. Therefore, the answer to this question gives 1 bit of information. But for the question: “Is the surname listed from 1 to 8?” the range of numbers only covers a quarter of all the numbers and therefore the probability of a “yes” is $1/4$, while that of a “no” is $3/4$. The answer to this question would contain less than 1 bit of information. According to (3.3), in which we substitute $P = 1/4$, each answer contains 0.8 bit of information.

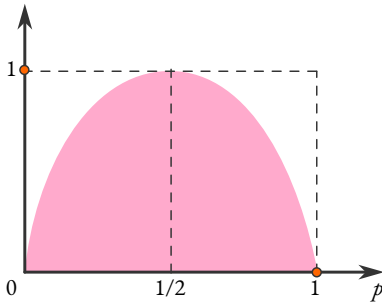


Figure 3.6: Graph of Shannon distribution.

Once again I emphasize that control processes should be regarded in a dialectical unity with the random processes of disorganization. There is a deep relationship between information theory and probability theory. The Shannon formula (3.2) illustrates this point. The probabilistic approach provides a scientific, objective notion of information that is free from a subjective substitution of the quantity of information by its significance or importance.

Information in communication channels with noise. When information is transmitted, some loss is unavoidable. This happens because of the action of random factors, which

are commonly lumped together as noise. A *communication channel* for transmitting information from input set A to output set B is represented in Figure 3.7. The information is affected by noise P as it is transmitted. Suppose that ξ is an input discrete random variable which may assume values $x_1, x_2, x_3, \dots, x_N$, with probabilities $p_1, p_2, p_3, \dots, p_N$, and η is the output variable, which may assume values $y_1, y_2, y_3, \dots, y_M$, with probabilities $q_1, q_2, q_3, \dots, q_M$. Let $P_i(j)$ denote the probability that $\xi = y_j$ is the output variable if $\xi = x_i$ was transmitted. The probability $P_i(j)$ is determined by noise in the communication channel. It has been proved in information theory that the quantity of information about the random variable ξ that can be obtained by observing the random variable η is described by the formula

$$I_\eta(\xi) = \sum_{i=1}^N \sum_{j=1}^M P_i(j) p_i \log_2 \frac{P_i(j)}{q_j}. \quad (3.4)$$

Here the information I is in terms of two types of probability, the probabilities p_i and q_j

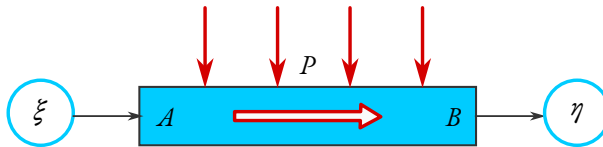


Figure 3.7: A communication channel for transmitting information.

on the one hand and the probability $P_i(j)$ on the other. While the first two probabilities reflect the probabilistic nature of the information at the input of the communication

channel and that “yes” or “no” questions, the answers to which are nearly or equally probable. Recall the question: “Is the surname listed from 17 to 32?” received at the output, probability $P_i(j)$ reflects the random nature of the noise in the channel.

Suppose there is no noise. Then the random variable values at the input and the output of the channel will be the same. Hence

$$N = M, \quad p_i = q_i, \quad \text{and} \quad P_i(j) = \delta_{ij}, \quad (3.5)$$

where $\delta_{ij} = 1$ for $i = j$ and $\delta_{ij} = 0$ for $i \neq j$. Substituting (3.5) into (3.4) and noting that $\lim_{z \rightarrow 0} z \log_2 z = 0$, we get the Shannon formula. This should have been expected because when there is no noise, there is no loss of information in its transmission.

Protection against noise in a communication channel. There are many sorts of communication channel. Information can be transmitted by sound waves propagating in a medium, electric signals running along wires, electromagnetic waves propagating in a medium or in vacuum, etc. Each communication channel is affected by its own sorts of noise. There are general techniques for handling noise that can be applied to any communication channel. First of all, it is desirable to minimize the level of noise and maximize the amount of information in the signals, so that the signal-to-noise ratio is large. The ratio can be increased by coding the transmitted information appropriately, e. g. transmitting it in terms of “symbols” (for instance, impulses of a certain shape) which can be distinctly identified against the background of noise. Coding a signal increases its “noise immunity” or performance in terms of error probability for the transmission.

A special measure against noise is *filtering* (both *smoothing* and *correlation*) the information received at the output of communication channels. If the characteristic noise frequency in a communication channel is substantially greater than the frequency typical for the time change in the signal, we could use a *smoothing filter* at its output to “cut out” the high-frequency oscillations superimposed on the signal as it was transmitted. This is illustrated in Figure 3.8, in which (a) is a diagram of the communication channel with a filter (A is the channel input, B is the channel output, P is noise, and F is a smoothing filter), (b) is the signal at the input, (c) is the signal at the output before filtering, and (d) is the signal after filtering.

Suppose we want to find out whether the output contains a signal of a given shape. If the signal is very different (for instance, by frequency) from the noise background, it will be easily identified. The situation is worse when the signal is “masked” by noise. Correlation filtering is applied in these cases: a device is placed at the output which *multiplies* the output signal by the known signal. If the desired signal is present in the output signal, the multiplication creates a very clear (large) final (correlation) signal; otherwise no correlation signal will appear. This is illustrated in Figure 3.9, in which

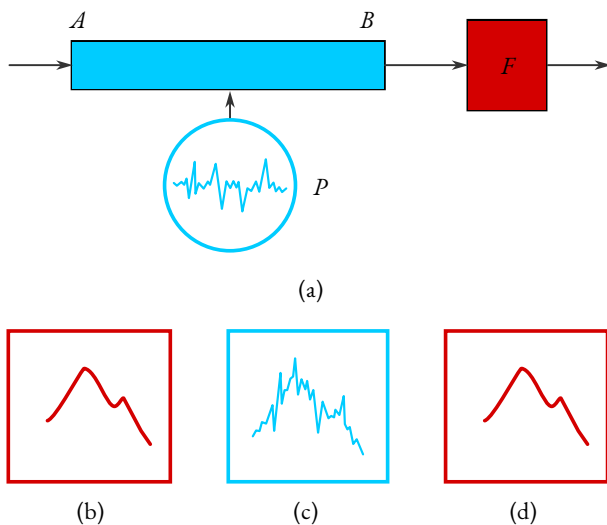


Figure 3.8: A communication channel with a filter.

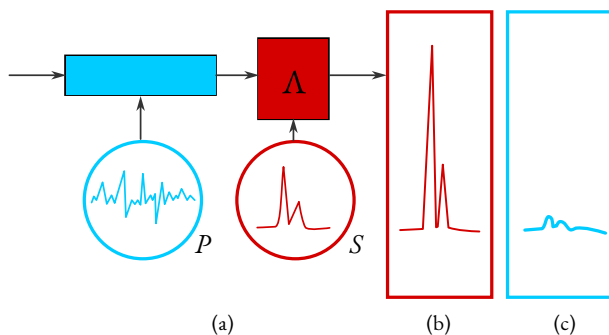


Figure 3.9: A communication channel with a filter and signal multiplier.

(a) is a diagram of the channel (Λ is the signal multiplier, P is noise, and S is the signal shape to be recognized), (b) is the multiplied signal if the recognized signal S is present in the output (the correlation signal), and (c) is the multiplied signal if signal S is absent in the output. Correlation filtering is used, for instance, in radar scanners to recognize the radiation signal emitted by the radar antenna.

Selection of Information from Noise

Where does information come from and some unsatisfactory answers. Any control signal carries certain information. The signal is formed using an algorithm which itself incorporates information, and this algorithm was compiled in turn using information contained in other algorithms. Thus we have a sort of relay race in which information is transmitted from algorithm to algorithm. This idea can be illustrated by a simple example. A teacher educates you, and in turn your teacher had a teacher, who had a teacher, and so on.

This argument leads inevitably to the questions: Whence the “original information”? Whence the first algorithm? An inability (or reluctance) to investigate scientifically the fundamental topic of where information comes from leads to serious misconceptions.

One such misguided hypothesis is that the original information was brought to the Earth by space travellers, who visited us in some long-forgotten past. This hypothesis is materialistic, but it is unsatisfactory because it begs the question of where the aliens got the information. Modern science indicates where the information comes from. The modern scientific answer is that there is no “original information”: the generation of information is a continuous and continuing process.

Chance at the forefront again. The idea of information being handed over like a relay baton in a race is simplistic. I pointed out that any transmission of information is accompanied by loss caused by random factors. However, random factors not only “steal” information, they also generate it.

At first glance, this seems implausible. We witness the continuous creation of information as a result of human creativity. New machines are designed, spacecraft are launched, new books are published, and new drugs become available: these are all a testimony to the explosive generation of information in which everybody participates. So it would seem strange to speak of the fundamental role of chance in generating information.

However, consider the *process of thinking*, how a problem is solved, how an intuition appears, or how a melody or image emerges. If these examples are too philosophical, try and think at least about *associative perception*, that is how we recognize objects and distinguish them. Just try, and you will step into a domain of complicated links, probabilistic relationships, chance guesses, and sudden “revelations”. There are no deterministic algorithms for making discoveries or solving problems. Everything we know about the processes occurring in the brain indicates the *fundamental role* of

random factors. Later I shall illustrate this by the example of *perceptron*, a cybernetic device which can recognize patterns.

Chance and selection. How can chance generate information? How can order appear from disorder? It turns out that the generation of information from noise can be easily observed. You can see this for yourself using the game of scrabble, or rather the small lettered blocks. Put one block with each letter of the alphabet into a bag, mix them, and take one out at random. Write down each randomly taken letter and return the block to the bag. Each time shake the bag. This simple generator of random letters can be used to generate a long chaotic string. If you look closely, you will find some three-letter words, perhaps even words with more letters. Information is being generated from noise.

My son, for example, helped me do an experiment and in a string of 300 random letters found nine three-letter words and two four-letter. This argument leads inevitably to the questions: Whence the “original information”? Whence the first algorithm? An inability (or reluctance) words. The more letters there are in a word, the smaller the probability of generating the word from “letter noise”. The generation of a sentence, let alone a line from a well-known work, is less probable. Nonetheless, the probability of doing is nonzero, and so there is the possibility of any information being generated randomly from noise.

Thus, we can say (although this sounds strange) that *chance generates information by chance*. The greater the information, the smaller the probability of its random generation. That random information can be generated does not solve the basic problem. This randomly generated information must be detected from the enormous flow of meaningless “signals”. In other words, the *information must be selected from the noise*. In the example of taking lettered blocks out, the information is selected from the noise by the person who wrote out the letters and looked through the string.

Selection amplifier Is it possible to use chance conscientiously to generate information? It is, so long as we *amplify the selection*.

You can do a simple experiment to demonstrate the amplification of selection using the random letter generator described above. In order to amplify the selection, we take into account the frequency with which letters appear in each word. Letter frequencies in English are often given when you buy a commercial game of scrabble. To allow for the frequencies, first eliminate the rare letters, e. g. *Z, Q, J, V, X* and add extra blocks with frequent letters, e. g. four blocks with *E* and *T*, three with *A, I, O, L, N, G, R, S*, two with *D, U*, and one of all the rest. I cannot vouch that this selection is optimal, in a similar experiment I found 21 three-letter words, 4 four-letter words and 1 five-letter

word in a succession of 300 random letters.

In order to amplify the selection still greater, we should use *words* rather than *letters*. It is curious that a similar device was suggested in the early 18th century by the English satirist Jonathan Swift in *Gulliver's travels*. When Gulliver visited the Academy in Lagado (the capital of an imaginary kingdom), he met a professor who had an interesting apparatus. Swift wrote:

"He then led me to the frame, about the sides whereof all his pupils stood in ranks. It was twenty feet square, placed in the middle of the room. The super faces were composed of several bits of wood, about the bigness of a die, but some larger than others. They were all linked together by slender wires. These bits of wood were covered on every square with papers pasted on them, and on these papers were written all the words of their language in their several moods, tenses, and declensions, but without any order. The professor then desired me to observe, for he was going to set his engine at work. The pupils at his command took, each of them, hold of an iron handle, there were forty fixed around the edges of the frame, and given then a sudden turn, the whole disposition of the word was entirely changed. He then commanded six and thirty of the lads to read the several lines softly as they appeared on the frame; and where they found three or four words together they might make part of a sentence, they dictated to the four remaining boys who were scribes. This work was repeated three or four times, and at every turn the engine was so contrived, that the words shifted into new places, as the square bits of wood moved upside down."

True, Swift wrote satirically, laughing about such inventions. However, why should we not believe that a talented popular-science writer disguised himself behind mask of a satirist so as not to be laughed at and misunderstood by his contemporaries?

What seemed absurd and laughable in the 18th century has now become the subject of scientific investigation in the mid-20th century. The English scientist W. Ross Ashby suggested a cybernetics device in the early 1950s which could be a *selection amplifier*. Ashby called it an *intelligence amplifier*. A diagram of this amplifier is given in Figure 3.10.

Noise generator 1 supplies "raw material" to the first stage of the amplifier. The noise converter 2 produces various random variants of the subjects to be selected. The selection is performed in unit 3 in compliance with criteria of selection put into this device. In a concrete case, if the result of a selection meets a criterion, control unit 4 opens valve 5 and lets the selected information into the converter of the next stage of the amplifier. One can easily imagine that the first stage of the amplifier, supplied with random letters, selects separate randomly emerging words or separate typical syllables; the second stage of the amplifier selects word combinations; the third stage selects

sentences, the fourth stage selects ideas, etc.

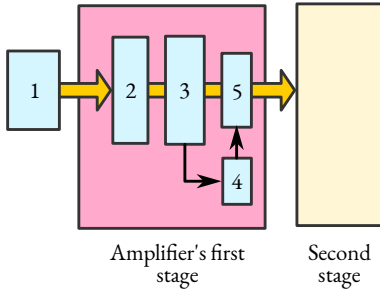


Figure 3.10: A selection amplifier.

was repeated three or four times, and at every turn the system to this new state. How is it to be done? Information is needed firstly on the new state, and, secondly, on how the transition of the system to the new state can be carried out. Since the change in the environment is random in nature, we know neither the new normal state nor how to organize a transition to it. A *random search* may help in such situations. This means that we should randomly change the system's parameters until it randomly matches the new normal state, which can be immediately recognized by monitoring the system's behaviour.

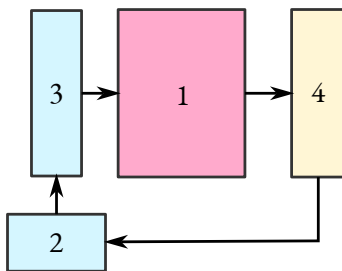


Figure 3.11: A homeostat is a device which possessed the property of self-organization on the basis of random search.

Random search-related self-organization.

The homeostat. Suppose a system is in a state which allows it to carry out certain functions. Let us call this state *normal*. It corresponds to external conditions in which the system operates. Suppose these conditions change all of a sudden, and the result is that the system departs from the normal state. The new conditions correspond to a new normal state. It is desirable to transfer and where they found three or four words together they might make part of a sentence, they dictated to the four remaining boys who were scribes. This work

It can be said that the process of random search generates the information needed to transfer the system to the new normal state. This is nothing else but the *selection of information from noise* about which we have been talking. The selection criterion here is the change in the system's behaviour: once in the new normal state, the system "calms down" and starts functioning normally.

In 1948 Ashby designed a device which possessed the property of self-organization on the basis of random search. He called the device a *homeostat*. A diagram of a homeostat is shown in Figure 3.11.

A homeostat is often compared to a sleep-

ing cat. If the cat is bothered, it wakes up, chooses a new more comfortable position, and goes to sleep again. A homeostat behaves in a similar manner: when it is “woken up”, it carries out random search for new values for its parameters, and when it finds them, it “goes to sleep” again.

System 1 in Figure 3.11 may be either in a stable or unstable state. Without going into detail, let me note that system 1 consists of four electromagnets whose cores can move and control the rheostats which control the voltages across the electromagnets. Therefore, the rotation angle of each electromagnet is dependent on all the other ones. These angles are the parameters of this dynamic system. The magnet cores do not rotate when the system is in a stable state. However, if an external disturbance takes the system out of its stable state, control unit 2 switches on generator 3 of random changes of parameters, and the random search starts. Once system 1 finds a stable state (by chance), the system to this new state. How is it to be done? Information is needed firstly on the new state, and, secondly, on how the transition of the system to the new state can be carried out. Since the chance in the unit 4 having verified the stability sends a signal to control unit 2, which switches off the random parameter generator 3.

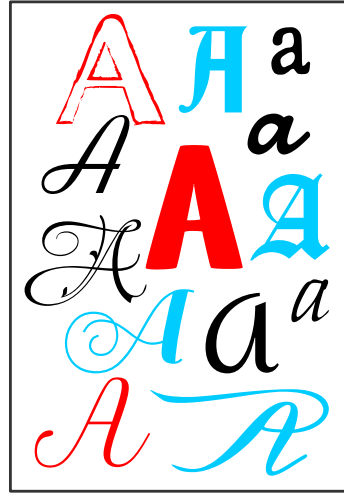


Figure 3.12: A selection of different ways of writing letter A.

On the Way to a Stochastic Model of the Brain

The pattern recognition problem. We do not commonly think about the brain’s ability to *recognize* patterns, although it is amazing. Several characters differing in size, shape, and line breadth are shown in Figure 3.12. Despite this, we immediately recognize the same character, the letter *A*, in every image. It is still more amazing when there is a crowd of variously dressed people with poorly distinguishable faces (because of the distance) and yet we usually manage to distinguish between men and women without error.

The ability to recognize patterns is called *associative perception*, i.e. when certain general, characteristic features are perceived while other more individual aspects recede into the background. Is associative perception possible for a machine? Is it possible to simulate the processes occurring in the brain and relate them to pattern recognition? These questions were answered in the affirmative in 1960 when the American scientist F. Rozenblutt designed a device he called a *perceptron*.

What is a perceptron? A perceptron can be regarded as an oversimplified model of the eye-brain system. The role of the eye, or, more accurately, the retina of the eye, is played by a grid consisting of a large number of *photoelectric cells*, or *receptors*. Each receptor converts the light incident on it into electric signals which are collected by the analysis unit within the perceptron. Before going into detail on the perceptron, let me make two fundamental points. Firstly, the relations between the receptors and the perceptron's internal units which process the information recorded by receptors should not be rigidly defined. If they were so defined, the signals from the images shown in Figure 3.13 (a) and (b) would be "perceived" by the perceptron as different patterns (only five excited receptors shown in red coincide in these images), while the images in Figure 3.13 (a) and (c) would be "perceived", by contrast, to be the same pattern because there are 28 excited receptors in common. In reality, a perceptron should "perceive" the images in Figure 3.13 (a) and (b) as the same pattern while those in Figure 3.13 (a) and (c) as different patterns. Thus, we must accept that the internal relations in a perceptron should be *random*. They have to be *probabilistic* relations.

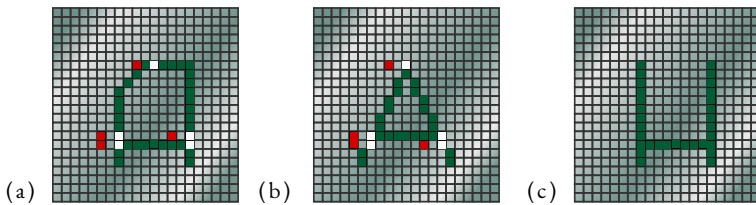


Figure 3.13: Signals and their interpretations by a perceptron.

Secondly, the random nature of these relations suggests the *adjustment* of the perceptron to the patterns being recognized. A perceptron should be presented with different images of the recognized patterns in turn (and several times), and we should *teach* it, the perceptron's parameters being adjusted as needed in the process. A perceptron should take into account its progress at each stage (at each presentation of the image), so a perceptron should have a memory.

Considering both these points, we can define perceptrons to be devices which have

a memory and a random structure of the links between its units. A perceptron can be thought of as a simplified model of the brain, and this model is promising because it is *probabilistic*, or, in other words, *stochastic*. Some scientists believe that stochastic models will be best able to simulate the processes occurring in the brain. Various sorts of perceptron have been designed. Below we shall consider a simple perceptron which can distinguish two patterns.

The arrangement of the simplest perceptron. A diagram of this perceptron is given in Figure 3.14. Here the S_i , are photoelectric cells (receptors), the I_k are phase inverters, which change the sign of the electric voltage, the A_j are associative units (A -units), the λ_j are amplifiers with varying gain factors, Σ is a summator, and R is the receiver. Suppose that the total number of receptors S_i , is N , ($i = 1, 2, 3, \dots, N$). In the first models, N was $20 \times 20 = 400$ receptors. The number of inverters is not fixed in that it can be different in different copies of the same device. The total number of associative units A_j and amplifiers λ_j equals M ($j = 1, 2, \dots, M$). The receptors are wired to the A -units either directly or via the inverters. It is essential that the choice of which receptor is connected to which A -unit and the selection of the potential sign are random. Thus when a circuit is being assembled, the wires connecting the receptors to the A -units are soldered together *randomly*, for instance, in accordance with instructions from a random number generator.

Suppose that an image is projected onto the perceptron's sensor grid. Since the intensity of the light at each point is different, some of the receptors will be excited, generating a logic signal of 1, while others will not, generating an electric signal of 0 at the output of the receptor. If the signal passes through an inverter, a 1 is transformed into a -1 . The system of random links transmits the signals from the receptors to the A -units. Each A -unit algebraically adds up the signals at its input. If the sum is above a threshold, the output of the A -unit goes to logic $+1$, otherwise it goes to logic 0. Let us designate the signals leaving the A -units y_j . Each y_j is either $+1$ or 0. The signal at the output of unit A_j goes to the input of amplifier λ_j , and the amplifier transforms signal v_j to a signal $\kappa_j y_j$. The gain factor κ_j may vary both in absolute value and in sign. The signals from all the amplifiers are summed up in the summator Σ , and hence we get

$$\sum_{j=1}^M \kappa_j y_j.$$

Then it is sent to the input of the R -unit, which checks its sign. If $\sum_j \kappa_j y_j \geq 0$, the R -unit output is $+1$, otherwise the R -unit output is 0.

This perceptron is designed to recognize only two patterns. Irrespective of the concrete images of the patterns, the perceptron will respond to one pattern with an

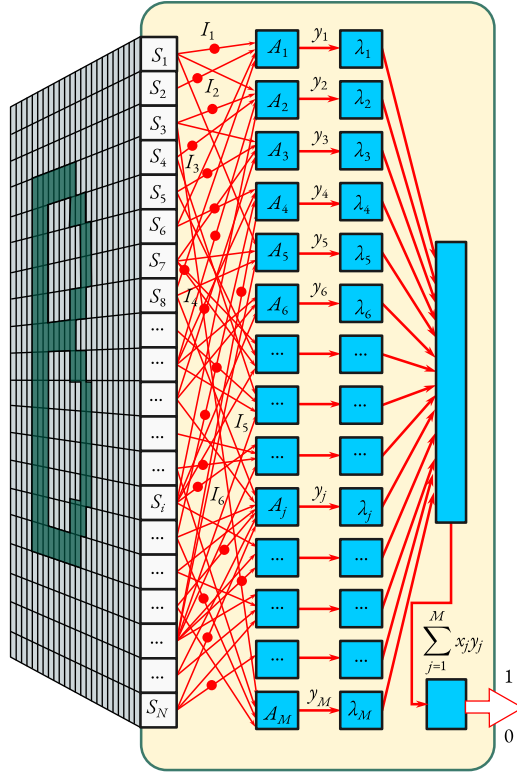


Figure 3.14: Schematic diagram of the simplest perceptron.

output signal of + 1 and with a signal of 0 to the other. The perceptron must *learn* this ability.

Teaching a perceptron. Let us call the two patterns B and C . Suppose pattern B corresponds to an output signal of + 1 and pattern C to an output signal of 0. Suppose $x_1, x_2, x_3, \dots, x_j, \dots, x_M$ are the perceptron's gain factors before it is taught. Let us designate this ordered set $\{x\}$. To teach the perceptron, we present it with an image of pattern B . This will excite a certain set of A -units, i. e. we get a succession of signals $y_1, y_2, y_3, \dots, y_j, \dots, y_M$, or, in short, $\{y\}$. Now suppose the sum $\sum_j x_j y_j$ is non-negative, so the perceptron's output signal is + 1. If so, then everything is true, and we can present the perceptron with a second image of pattern B . The second image will excite a new set

of A -units, i.e. a new succession of signals $\{y'\}$. The set of gain factors $\{x\}$ remains yet the same, but the sum $\sum_j x'_j y'_j$ may be negative, and then the signal at the perceptron's output will be 0. This is not good, and therefore the perceptron is "discouraged": the gain factors of the excited A -units are incremented by, say, unity, so that a new set of gain factors $\{x'\}$ ensures that the sum $\sum_j x'_j y'_j$ is non-negative. Now the perceptron responds correctly to the second image of pattern B . But what about the first image? The set of gain factors has been changed, so that the sign of the sum $\sum_j x'_j y_j$ may be changed. We present the perceptron with the first image of pattern B again and identify the sign of the sum $\sum_j x'_j y_j$ by the output signal.

If the sum is non-negative, we are satisfied because the set of gain factors $\{x'\}$ has caused the perceptron to respond correctly to both the first and the second images of pattern B . Now we can present the perceptron with a third image of pattern B . If the sum is negative, the gain factors of the excited A -units should be incremented by unity again (set $\{x'\}$ is replaced by set $\{x''\}$), and so on.

Gradually, by varying the set of gain factors step by step, we will find a set of factors such that the perceptron will produce a signal of +1 for any presented image of pattern B . However, our job is not yet over. It is quite possible that after many increments of the various gain factors, the perceptron will produce a +1 signal for both pattern B and pattern C images. However, the perceptron should produce a +1 signal for all pattern B images and a 0 signal for all pattern C images. This means that while the perceptron is being taught, we should alternate between both patterns and when presenting an image of pattern C , we should (if need be) decrement rather than increment the gain factors of the excited A -units to get $\sum xy$ below zero.

Ultimately, we will find a set of gain factors $\{x^0\}$ such that the perceptron will always recognize patterns B and C . Suppose $\{y(n)\}$ is the set of excited A -units corresponding to an n th image of pattern B and $\{Y(m)\}$ is a set corresponding to an m th image of pattern C . The M gain factors $\{x^0\}$ should be such that

$$\sum_{j=1}^M x_j^0 y_j(n) \geq 0 \text{ for all } n \text{ and } \sum_{j=1}^M x_j^0 y_j(m) \leq 0 \text{ for all } m$$

The teaching (and the learning) is over when such gain factors have been found.

Conclusion. In concluding this chapter, I went to emphasize the main point, i.e. the deep, intrinsic relationship between *information theory* and *probability theory*. The very notion of information is underlain by probability. And this is natural because control processes and random processes are always *dialectically united*. Randomness

both “steals” information and generates it, because the most complicated information devices are fundamentally based on random internal structures.

We see that the phrase “the world is built on probability”, which is the title of this book, has a deeper meaning. Mankind lives and acts in a world filled with information. This information arises by nature through probability and, moreover, is created in probabilistic processes. Thus, a *world filled with information* is naturally a *world built on probability*.

Part II

Fundamentality of the Probability Laws

Chapter 4

Probability in Classical Physics

Probability theory is used in physics, and its first application of fundamental importance for our understanding of the laws of nature can be found in the general statistical theory of heat founded by Boltzmann and Gibbs ... The most elegant and important advantage of this theory is the understanding of thermodynamical “irreversibility” as a picture of transition to more probable states.

W. Pauli

Thermodynamics and Its Puzzles

All bodies consist of molecules in chaotic thermal motion. This fundamental point can be disregarded when considering the basic problems of *thermodynamics*, the branch of physics which seeks to derive, from a few basic postulates, relationships between the properties of matter, especially those which are affected by changes in temperature, and a description of the conversion of energy from one form to another.

Thermodynamics is a branch of physics in which the energy transfers between macroscopic bodies and their environment are investigated from the most general positions (without using molecular concepts). Thermodynamic considerations are underlain by a description of the states of the bodies using thermodynamic *variables or the thermodynamic functions of state* or state parameters, and the use of several basic principles called the *laws of thermodynamics*. You already know about such thermodynamic variables as

temperature and *pressure*.

Thermodynamic equilibrium. Let us perform a simple experiment. Take a vessel with hot water into a room and put a thermometer into the water. By recording the readings of the thermometer over time, we will see that the temperature of the water gradually decreases until finally equals the air temperature in the room, after which the temperature will remain constant. This means that the water in the vessel has reached a *thermodynamic (heat) equilibrium* with the environment. If a system is in a thermodynamic equilibrium, its thermodynamic functions of state (temperature and pressure) remain constant until disturbed. Another feature of a thermodynamic equilibrium is that the temperature is constant at all points of the system.

If a system does not exchange energy with bodies around it, it is a *closed system*. When we talk about a thermodynamic equilibrium of a closed system, we mean an equilibrium between its various parts, each of which can be regarded as a macroscopic body.

Suppose we heat a body unevenly and then put it in a vessel which does not conduct heat. It can be said that we first disturb the thermodynamic equilibrium in the body and then leave it. The temperature of the hotter regions will decrease, and that of cooler ones will increase, and finally the temperature will become the same throughout the body: they will reach a thermodynamic equilibrium with each other. *An unperturbed macro-system will always reach a state of thermodynamic equilibrium and remain there* until some external action brings it out of this state. If this action stops, the system will again reach a thermodynamic equilibrium.

And here is the first puzzle of thermodynamics. Why does a system brought out of thermal equilibrium and left to itself return to an equilibrium state, while systems in a thermal equilibrium and left to themselves do not leave it? Why is it not necessary to spend energy to maintain thermal equilibrium, while energy is needed to maintain a system in a thermodynamic equilibrium? By the way, this is a far from futile question. The weather outside may be below freezing, e.g. -1°C , while it's warm in the room, 25°C . The walls of houses conduct heat fairly well, and therefore, there is a non-equilibrium "room-outside" system. To maintain this thermodynamic non-equilibrium state, it is necessary to spend energy continuously to heat.

The first law of thermodynamics. A system may exchange energy with its environment in many ways, or, as is said, along many channels. For simplicity's sake, let us limit ourselves to a consideration of two channels, namely, the transfer of energy by *heat conduction* and the transfer of energy by *performing work*. The *first law of thermodynamics* is simply the law of the conservation of energy involving the possible

energy transfer between a body and its environment via different channels, i.e.

$$\Delta U = A + Q, \quad (4.1)$$

where $\Delta U = U_2 - U_1$ is the change in the internal energy of the body (U_1 and U_2 being the internal energies of the initial and final states of the body, respectively), A is the work performed by external forces with respect to the body, and Q is the amount of heat transferred to or from the body by conduction. Note that unlike internal energy, which is a *function of state* of the body (it varies when the body transfers from one state to another), neither work nor heat are functions of state. It is equally absurd to say that a body in a state has so much heat or so much work. The heat Q and work A in formula (4.1) are the changes in the body's energy carried out through different channels. Let us consider a simple macro-system, an *ideal gas* (m is the mass of the gas). The internal energy of an ideal gas is proportional to the absolute temperature T of the gas and does not depend on the volume V it occupies. Let us change the gas volume using a piston. By pushing a close-fitting piston down a cylinder and thus compressing the gas in the cylinder, we perform some work A . When the gas expands, it performs work A' to move the piston back: $A' = -A$. This work is related to the change in the gas volume. It is numerically equal to the area under the pressure-volume curve, which describes the process, from $V = V_1$ to $V = V_2$, where V_1 and V_2 are the initial and final volumes of the gas.

Let us consider, from the viewpoint of the first law of thermodynamics, two types of gas expansion, *isothermal* and *adiabatic*. The former process occurs at constant gas temperature while the latter occurs when there is no heat exchange between the gas and the environment. The change in the gas volume should be carried out very slowly (compared to the rate at which thermal equilibrium is reached within the gas), and so the gas can be regarded at any moment in time as being in thermodynamic equilibrium. In other words, we assume that the gas passes from one thermodynamic equilibrium state to another, as it were, via a succession of intermediate equilibrium states.

If the expansion is *isothermal*, the gas's temperature remains constant, and therefore, $\Delta U = 0$ ($U_1 = U_2$). Noting this, we obtain from (4.1):

$$-A = Q \quad \text{or} \quad A' = Q. \quad (4.2)$$

The expanding gas performs as much work as it receives heat from the environment during its expansion.

When the expansion is *adiabatic*, there is no heat exchange with the environment ($Q = 0$). Therefore,

$$\Delta U = A \quad \text{or} \quad A' = -\Delta U. \quad (4.3)$$

The expanding gas performs work owing to a decrease in its internal energy, and the gas's temperature therefore falls.

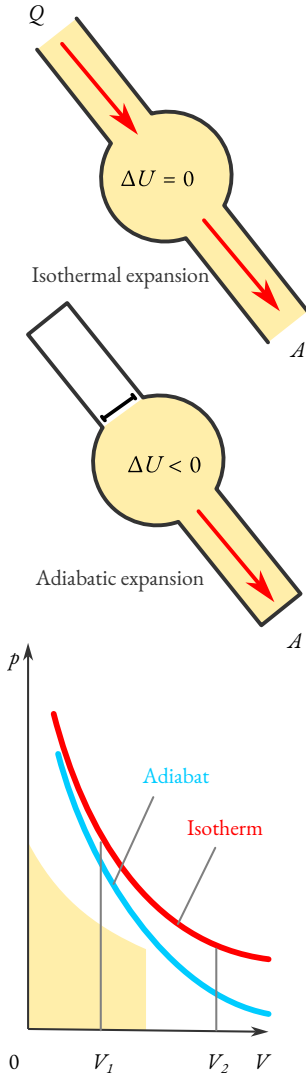


Figure 4.1: Isothermal and adiabatic processes.

Both of these processes are conventionally shown in Figure 4.1. The processes are also represented on $p - V$ diagrams (where p is the gas pressure). The work A' performed by the gas in an isothermal expansion from volume $V = V_1$ to $V = V_2$ equals numerically the yellow area under the plot of $p(V)$ in the figure:

$$A' = \int_{V_1}^{V_2} p(V) dV. \quad (4.4)$$

Using an equation of state for an ideal gas (the Mendelev-Clapeyron equation), we get

$$p = \frac{mRT}{MV}, \quad (4.5)$$

where M is the molar mass of the gas and R is the universal gas constant. Substituting (4.5) into (4.4) and given that the temperature of the gas is constant, we obtain

$$A = \frac{mRT}{MV} \int_{V_1}^{V_2} \frac{1}{V} dV = \frac{mRT}{MV} \ln \frac{V_2}{V_1}, \quad (4.6)$$

(the symbol \ln designates a logarithm to base $e = 2.71828 \dots$).

The Carnot cycle. In 1824, a 28-year-old engineer called Sadi, Carnot published a book in Paris entitled *Refléxions sur la puissance motrice du feu et le machine propre à développer cette puissance* (Reflections on the Driving Force of Fire and Machines Capable of Developing This Force). Unfortunately, his ideas as presented in the book were only appreciated many years later, and long after he had died.

Carnot was investigating the work obtained from heat engines. He showed that a heat machine not only needs a hot body, it also requires a second body with a lower temperature. The first body is conventionally called the heat source, and the second is called the heat sink. Besides the heat source and heat sink, there must be a working substance (a liquid, steam, or gas), which transmits the heat from the heat source to the heat sink and performs work in the process. Carnot considered a closed cycle consisting of two isotherms and two adiabats. Later this cycle was called the *Carnot cycle*. It is shown in Figure 4.2 for an ideal gas.

Suppose T_1 is the temperature of the heat source and T_2 is that of the heat sink. Moving from point 1 to point 2 (the isotherm for T_1), the gas receives a heat Q_1 from the heat source and expands, thus spending energy to perform work A . From point 2 to point 3 (along an adiabat), the gas performs work A ; and its temperature falls to T_2 . From point 3 to point 4 (the isotherm for T_2) the gas gives a heat Q_2 to the heat sink, and this heat equals the work A_2 performed to compress the gas. From point 4 to point 1 (another adiabat), the work A_4 is expended to compress the gas, and this goes to increasing the internal energy of the gas, so its temperature rises to T_1 . The result is that the working substance returns to its initial state 1.

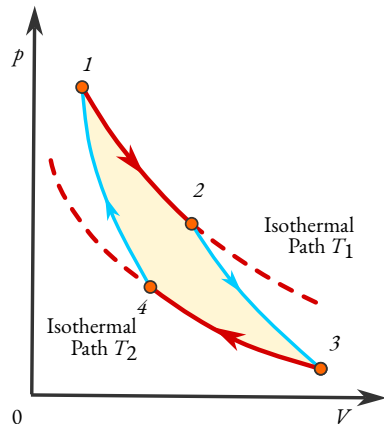


Figure 4.2: Carnot cycle for an ideal gas.

Suppose that a heat engine operates following the Carnot cycle. The gas receives a heat Q_1 from the heat source and gives a heat Q_2 to the heat sink. In compliance with (4.2), we can write $Q_1 = A'$ and $|Q_2| = A_2$. Note here that $Q > 0$ when heat is given to the gas, and that $Q < 0$ when the heat is taken from the gas. It is clear from Figure 4.2 that the area under isotherm 3-4 is smaller than that under isotherm 1-2, and therefore, $A_2 < A'_1$. Consequently, $|Q_2| < Q_1$ i.e. the gas gives the heat sink less heat than it receives from the heat source. At the same time, the internal energy of the gas, when the cycle is completed, remains the same. Therefore, the difference $Q_1 - |Q_2|$ equals the work performed by the heat engine during its cycle. Hence the efficiency of the heat engine is

$$\eta = \frac{(Q_1 - |Q_2|)}{Q_1}. \quad (4.7)$$

Carnot showed that

$$\frac{Q_1}{T_1} = \frac{(|Q_2|)}{T_2}. \quad (4.8)$$

This allows us to rewrite (4.7) in the form

$$\eta = \frac{(T_2 - T_1)}{T_1}. \quad (4.9)$$

The efficiency of a heat engine, as defined by (4.7) and (4.9), is the best possible efficiency. The efficiency of real heat engines is always less because of unavoidable irreversible processes.

Reversible and irreversible processes. The notions of reversible and irreversible processes are essential for thermodynamics. A process is said to be *reversible* if the system (the working substance) is in thermal equilibrium all the time, continuously passing from one equilibrium state to another. This process is completely controlled, while it lasts, by the changes in its parameters, for instance, the temperature or volume. If the parameters are changed in the reverse direction, the process will also go backwards. Reversible processes are also called *equilibrium processes*.

Boyle's (Mariotte's) and Gay-Lussac's (Charles') laws define reversible processes in an ideal gas. The expressions (4.7) and (4.9) we have just obtained are related to a reversible Carnot cycle, which is also called the ideal Carnot cycle. Each part of the cycle and the whole cycle can be reversed if desired.

An *irreversible* process is a process that cannot be controlled. It proceeds independently, or, in other words, spontaneously. The result is that we cannot reverse such a process. It was noted above that once a system is moved from its thermodynamic equilibrium, it tends spontaneously to another thermodynamic equilibrium state. Processes related to transition of a system from a non-equilibrium state to an equilibrium one are irreversible. They are also called *non-equilibrium* processes.

Here are some examples of irreversible processes: conduction of heat from a hotter body to a cooler one, mixing of two or more gases in the same vessel, expansion of a gas in vacuum. All of these processes occur spontaneously, without any external control. Heat does not spontaneously transfer from a cooler body to a hotter one. The components of a gas mixture do not spontaneously separate. A gas cannot spontaneously compress. I wish to emphasize: *every irreversible process is characterized by a definite direction*. It develops in a certain direction and does not develop in the opposite one. Which direction a process can develop along and which it cannot are problems related to the second law of thermodynamics.

The second law of thermodynamics. One of the first formulations of the *second law of thermodynamics* was given by the English physicist William Thompson (Lord Kelvin):

“It is *not* possible that, at the end of a cycle of changes, heat has been extracted from a reservoir and an equal amount of work has been produced without producing some other effects.”

This means that it is impossible to design a machine to carry out work by reducing the internal energy of a medium, sea water, for instance. Kelvin called such a machine a *perpetuum mobile of the second kind*. While some *perpetua mobile* violate the law of the conservation of energy (*perpetua mobile of the first kind*), those of the second kind do not contradict the first law of thermodynamics; they are instead forbidden by the second law.

In 1850, the German physicist Rudolf Clausius formulated the second law of thermodynamics as follows:

“The transfer of heat from a cooler body to a hotter one cannot proceed without compensation.”

It is useful to demonstrate the equivalence of the formulations given by Kelvin and Clausius. If we could, despite Kelvin’s formulation, “extract” heat from a medium and, using a cyclic process, turn it into work, then, using friction, transform this work into heat at a higher temperature, we would contradict Clausius’s formulation because it would involve the conduction of heat from a cooler body to a hotter one within a closed cycle without any external force performing work.

On the other hand, suppose that, despite Clausius’s formulation, we succeed in getting some quantity of heat Q to conduct itself from a cooler body (at a temperature T_2) to a hotter one (T_1), and subsequently, allow this heat to go naturally from the hotter body to the cooler at the same time performing some work A' while the rest of the heat $Q_1 = Q - A'$ returns to the cooler body. This process is shown in Figure 4.3 (a). It is clear that this process corresponds to direct transformation of heat $Q - Q_1$ into work A (Figure 4.3 (b)), which evidently contradicts Kelvin’s formulation.

Entropy. As he was studying Carnot’s investigations, Clausius discovered that relationship (4.8) is similar to a conservation law. The value of Q_1/T_1 “taken” by the working substance from the heat source equals the $|Q_2|/T_1$ “conducted” to the heat sink. Clausius postulated a variable S , which like the internal energy is a state function of the body. If the working substance (an ideal gas in this case) receives heat Q at temperature T , then S is incremented by

$$\Delta S = Q/T. \quad (4.10)$$

Clausius called S *entropy*.

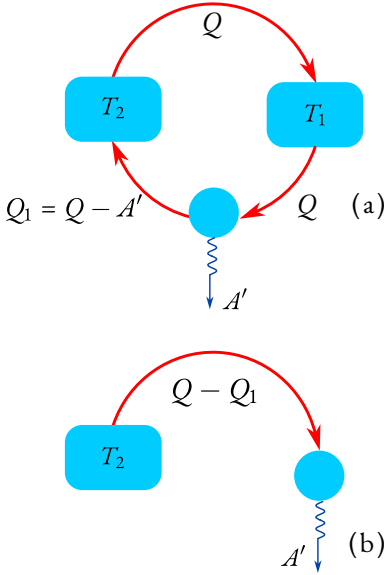


Figure 4.3: Work done in transfer of heat from bodies at different temperatures.

We shall later need to deal with a property of entropy called its additivity. This means that the entropy of a system is the sum of the entropies of the system's parts. Mass, volume, and internal energy are also additive. However, neither temperature nor pressure are additive.

The second law of thermodynamics as the law of increasing entropy in irreversible processes within closed systems. Using the notion of entropy, we can formulate the second law of thermodynamics as follows:

Any irreversible process in a closed system proceeds so that the system's entropy increases.

From point 1 to point 2 of the Carnot cycle (see Figure 4.2), a heat Q_1 is conducted from the heat source to the working substance at a temperature T_1 , and the entropy of the working substance increases by $\Delta S_1 = Q_1/T_1$. From point 2 to point 3 and from point 4 to point 1, there is no conduction of heat, and therefore, the entropy of the working substance does not vary. From point 3 to point 4, a heat Q_2 is conducted from the working substance to the heat sink at temperature T_2 , and the entropy of the body is decreased by $|\Delta S_2| = |Q_2|/T_2$ ($\Delta S_2 < 0$). According to (4.8) and (4.10),

$$\Delta S_1 + \Delta S_2 = 0. \quad (4.11)$$

Consequently, when an ideal (reversible) Carnot cycle comes to an end, the working substance's entropy returns to its initial value.

Note that entropy can be defined as the state function of a body (system) whose value remains constant during an adiabatic process. Similarly, temperature can be regarded as the state function of a system whose value remains constant during an isothermal process.

Consider the following irreversible process by way of an example. Suppose a closed

system consists of two subsystems 1 and 2 which are at temperatures T_1 and T_2 , respectively. Suppose that an infinitesimal amount of heat ΔQ is conducted from subsystem 1 to subsystem 2, so that the temperatures of the subsystems almost remain the same. The entropy of subsystem 1 reduces by $\Delta Q/T_1$, ($S_1 = -\Delta Q/T_1$) while the entropy of subsystem 2 increases by $\Delta S_2 = \Delta Q/T_2$. The entropy of the whole system is the sum of its subsystems' entropies, and therefore, the change in the system's entropy will be

$$\Delta S = \Delta S_1 + \Delta S_2 = \Delta Q \left(\frac{1}{T_2} - \frac{1}{T_1} \right). \quad (4.12)$$

Heat conduction from subsystem 1 to subsystem 2 is irreversible if $T_1 > T_2$. Using this inequality, we can conclude from (4.12) that $\Delta S > 0$. Thus, we see that the process of heat conduction from a heated body to a cooler one is accompanied by an increase in the entropy of the system consisting of the two.

A gain in entropy during irreversible processes is only a necessary law for closed systems. If a system is open, a reduction in its entropy is possible. Thus, if some external body does work with respect to the system, heat can be transferred from a heat sink to a heat source. It is essential that if the system includes a heat source, a heat sink, a working substance, and all the bodies that perform work (i.e. if we consider a closed system again), then, the total entropy of this system will increase.

I shall now formulate the basic conclusions concerning the change in the system's entropy.

The first conclusion. If a system is closed, its entropy does not decrease over time:

$$\Delta S \geq 0. \quad (4.13)$$

The system's entropy does not vary if the processes within it are reversible. If the processes are irreversible, the system's entropy increases. The gain in entropy can be regarded as a measure of the irreversibility of the processes occurring in it.

The second conclusion. Generally, nothing can be said about the change in entropy in an open system. It can either remain constant or increase or even decrease.

The puzzles of thermodynamics. These puzzles focus on the second law of thermodynamics. Since it gives a definite direction to the processes in nature, it introduces a fundamental irreversibility. How can this irreversibility be explained by physics? Why can heat be transferred from a hotter body to a cooler one while it cannot be spontaneously conducted in the opposite direction? Why does any gas expand in vacuum but does not compress spontaneously? Why, when in the same vessel, do two or more gases mix, but not spontaneously separate? A hammer strikes an anvil. The temperature of the anvil rises a bit. But however strongly we might heat the anvil with the

hammer resting on it, the reverse will not happen: the hammer will not jump off the anvil. Why? Very many similar “whys” can be asked. Thermodynamics does not answer these questions in principle. The answer must be sought in the kinetic theory of matter. We should now look into the picture of chaotically moving molecules.

Molecules in a Gas and Probability

A dialogue with the author. Imagine that we are talking with a physicist of the 1860s. We do not need a “time machine”. We shall just believe that my partner adheres to the views typical of physicists in the mid-19th century, the same physicists, many of whom later, in the 1870s, could not understand or accept the ideas of the Austrian physicist Ludwig Boltzmann (1844-1906). Anyway, let us imagine that it is 1861.

AUTHOR: “Let us consider a gas to be an ensemble of very many chaotically moving molecules.”

PARTNER: “Good. I’m aware of the recent investigations of James Clerk Maxwell, who calculated the velocity distribution of molecules in a gas.”

AUTHOR: “I would like to discuss some thing more fundamental than the distribution established by Maxwell. The point is that there is a qualitative difference between considering thermodynamic equilibria and considering the motion of molecules. In the first we have *dynamic* laws with strictly determined dependences, and in the second we have the *probabilistic* laws that govern processes in large ensembles of molecules.”

PARTNER: “But the movements of molecules are governed by Newton’s laws of classical mechanics rather than by probabilistic laws. Suppose we assign coordinates and velocities to all the molecules in a gas at a certain moment. Suppose that we can follow all the collisions of the molecules with each other and with the walls of the vessel. It is clear that in this case we will be able to predict where a molecule will be at some other moment and what velocity it will have.”

AUTHOR: “Why aren’t you bothered by the fact that you’re very much like the superbeing of which Laplace wrote?”

PARTNER: “I have a concrete problem in mechanics. True, the number of bodies is extremely great.”

AUTHOR: "There are about 10^{19} molecules in a cubic centimetre of gas under normal conditions. You have a problem in which some 10^{20} bodies have to be accounted for."

PARTNER: "Naturally, it would be exceptionally difficult. But the difficulty is purely technical and not fundamental. So long as our calculational abilities are limited, we shall have to resort to probabilities, the probability of a molecule arriving in a volume, its probability of having a velocity in a certain range, etc."

AUTHOR: "Thus, you believe that the use of probabilities is only related to our practical inability to perform a very cumbersome calculation, but that in principle an ensemble of molecules behaves according to Newton's laws as applied to individual molecules."

PARTNER: "Precisely. This is why I do not see the qualitative difference you mentioned."

AUTHOR: "I have at least three hefty arguments to support my position that the probabilistic description of large ensembles of molecules is necessary in principle, that chance is present in the very nature of these ensembles rather than simply being related, as you seem to believe, with our inadequate knowledge and inability to perform cumbersome calculations."

PARTNER: "I'd like to know of these arguments."

AUTHOR: "I'll start with the first. Suppose there is, as you postulate, a rigid system of strictly determined links (as given by Newton's laws) between the molecules in a gas. Now imagine that some of these molecules suddenly escape from this system (e.g. they escape from the vessel through a slit). Clearly the disappearance of these molecules will bring about the disappearance of all that is predetermined by their presence, I mean their later collisions with other molecules, which, in its turn, will change the behaviour of the other molecules. All this will affect the whole system of rigid relationships and, as a consequence, the behaviour of the ensemble as a whole. However, we know that from the viewpoint of gas as a whole you can suddenly withdraw a large number of molecules without any noticeable effect (for instance, 10^{12} molecules or more). The properties of the gas and its behaviour do not change in the least. Does this not indicate that the dynamic laws governing the behaviour of individual molecules do not actually interfere with the behaviour of the gas as a whole?"

PARTNER: “Still, it is hard to believe that molecules obey some laws while the ensemble of the same molecules obeys quite different laws.”

AUTHOR: “But this is exactly so. And my second argument will emphasize this fundamental point. I’ll give you some simple examples. A stone is thrown from point *A* at some angle to the horizontal (Figure 4.4 (a)). Imagine that we can change the direction of the stone’s velocity to the opposite at point *B* of its trajectory. It is clear that the stone should return to point *A* and have the same velocity (in absolute value) it had when it was thrown. The flying stone, as it were, ‘remembers’ its history.”

PARTNER: “This is natural because each state of the thrown stone is determined by its preceding one and, in its turn, determines the subsequent one.”

AUTHOR: “Another example: a ball hits a wall elastically and bounces off (Figure 4.4 (b)). If you change the direction of the ball’s velocity to the opposite one at point *B*, the situation will recur in the reverse order: the ball will hit the wall and return to point *A*.

“I cited these examples in order to illustrate an essential idea: the movements determined by the laws of classical mechanics have a kind of ‘memory’ of the past. This is why these movements can be reversed.”

“Another thing is the behaviour of gas. Imagine the following situation. There is a beam of molecules whose velocities are parallel. After entering a vessel, the molecules collide many times with each other and the walls. The result is that the molecules reach a state of thermodynamic equilibrium, and they lose all ‘memory’ of their past. It can be said that any gas in a state of thermal equilibrium, as it were, ‘forgets’ its prehistory and does not ‘remember’ how it arrived at the equilibrium state. Therefore, it is absurd to think of reversing the situation: the molecules could not recollect into a beam and depart from, the vessel in one definite direction. Many examples of such forgetfulness can be cited.”

“Suppose there is some gas on one side of a partition in a vessel and another gas is on the other side. If you take away the partition, the molecules of both gases will mix. Evidently, we should not expect this picture to reverse: the molecules will not move back into their own halves of the vessel. We might say that the mixture of two gases does not remember its prehistory.”

PARTNER: “Do you want to say that the equilibrium state of a gas is not pre-

determined by the preceding states of the gas?”

AUTHOR: “When we use the word predetermined, we mean strictly unambiguous predetermination. There is no such predetermination here. A gas may arrive in an equilibrium state from different initial states. No information may be obtained about the initial states by studying the gas in thermal equilibrium. This means that the gas forgets its prehistory.”

PARTNER: “Yes, this is true.”

AUTHOR: “And when does this loss of memory occur? It occurs when *chance* comes into play. You throw a die, and, say, a four turns face up. You throw again and a two appears. The appearance of the two is not related to the appearance of the four before it. You throw the die many times and obtain a set of digits. This set possesses stability (for instance, the four occurs approximately in one-sixth of all trials). This stability does not have any prehistory, it is not related to the occurrence of any other digit in the previous trials.”

“The same happens in a gas. The loss of prehistory indicates that we must deal with *statistical* laws, laws in which chance plays a fundamental role.”

PARTNER: “It seemed to me before that everything was clear. Newton developed his mechanics. Then the temperature and pressure of gas appeared. Using the notion of molecules, we reduced these physical variables to mechanical ones by relating temperature to the energy of molecules and the pressure of the gas to the impulses transferred to the wall by the molecules striking it. Therefore, the laws of mechanics were and continue to be fundamental laws. Are you suggesting we put probabilistic laws on the same level as the laws of mechanics?”

AUTHOR: “I believe that you are aware of the fact that some thermodynamic variables do not have analogues in classical mechanics. And here is my third argument. Entropy does not have a mechanical analogue. The very existence of a variable such as entropy is sufficient to disprove the thesis of the total fundamentality of the laws of classical mechanics.”

PARTNER: “I would not like to discuss entropy at all ...”

Let us finish with this dialogue because it has become a bit too long. We agreed that it referred to 1861. Therefore, I could not use arguments that were unknown at the time. But here I can cite two more arguments in favour of my position. Firstly, note

that entropy is explicitly expressed in terms of probability, and that namely this makes it possible to explain every puzzle of thermodynamics. We shall discuss this in detail in the next sections. Secondly, it follows from quantum physics that the assumption (made by my partner) that he can assign coordinates and velocities to all the molecules simultaneously proves to be inconsistent. This cannot be done due to fundamental considerations, which we shall talk about in detail in Chapter 5.

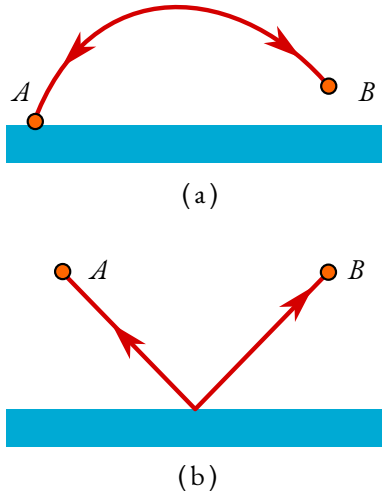


Figure 4.4: Bounce of a stone and an elastic ball.

And now let us discuss molecules moving in a gas.

Movements of gas molecules in thermodynamic equilibrium. Suppose a gas of mass m is in thermal equilibrium. The gas occupies volume V and has temperature T and pressure p .

Each gas molecule moves with a velocity which is constant in magnitude and direction until the molecule collides with either another molecule or the wall. On the whole, the picture of molecular movements is chaotic: the molecules move in different directions with different velocities, there are chaotic collisions leading to changes in the direction of movement and the absolute value of the velocities of molecules. Let us take an imaginary “photograph” of the molecules’ positions at a single moment in time. It might look like the one in (Figure 4.5), where for simplicity’s sake only two rather than three dimensions are consid-

ered (the “photograph” is flat). It is clear that the points (molecules) fill the volume of the vessel uniformly (the vessel in the figure is the square). Suppose N is the total number of molecules in the vessel; $N = N_A m / M$, where N_A is Avogadro’s number. At any site within the vessel and at any moment in time, the number of molecules per unit volume is the same (on average), N/V . Molecules may be found with equal probability at any point within the vessel.

Let us use $G(x, y, z) \Delta x \Delta y \Delta z$ to denote the probability of finding a molecule within a volume $\Delta V = \Delta x \Delta y \Delta z$ in the vicinity of a point with coordinates (x, y, z) . To be more accurate, this is the probability that the x -coordinate of the molecule will take a value from x to $x + \Delta x$, its y -coordinate from y to $y + \Delta y$, and its z -coordinate

from z to $z + \Delta z$. At small Δx , Δy , and Δz , the function $G(x, y, z)$ will be the density of the probability of finding a molecule at point (x, y, z) . The probability density in this case does not depend on the coordinates, hence $G = \text{const}$. Since the probability of finding a molecule somewhere within the vessel is unity, we have

$$\int_V G dV = 1, \quad \text{or} \quad G \int_V dV = GV = 1.$$

Consequently, $G = 1/V$.

Wherever a unit volume is taken within the vessel, the probability of finding a molecule within the unit volume is $1/V$, i.e. the ratio of the unit volume to the volume of the vessel. Generalizing this conclusion, we can state that the probability of finding a molecule within volume V_0 is V_0/V .

Now let us discuss the velocities of the gas molecules. It is clear from the start that the velocities cannot all be equally probable: there should be few molecules with very high and very small velocities. When considering the velocities of molecules, it is convenient to use the concept of a *velocity space*, i.e. the molecular velocities are projected onto the coordinate axes v_x, v_y, v_z . For simplicity's sake, Figure 4.6 shows only two axes: the v_x -axis and the v_y -axis (a two-dimensional velocity space). The figure shows a molecular velocity distribution in a gas for some moment in time. Each point in the figure relates to a molecule. The abscissa of the point is the x -projection of the molecule's velocity and the ordinate is its y -projection.

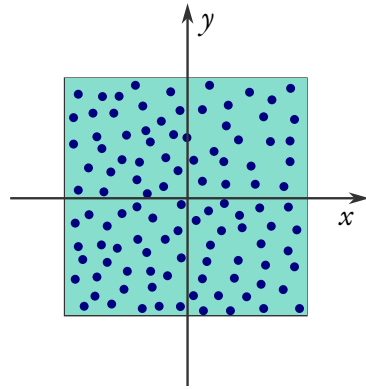


Figure 4.5: A snapshot of the molecules in motion.

It is interesting to compare Figure 4.5 and Figure 4.6. The points in Figure 4.5 are within a certain area and the distribution is uniform. The scatter of points in Figure 4.6 is unlimited in principle. These points clearly focus around the origin. This means that although the projection of a molecule velocity may be as large as you wish, the projections of the velocities in the neighbourhood of zero are the most probable. The scattering in Figure 4.6 is rotationally symmetric for any angle about the origin. This means that all directions of movement are equally probable: a molecule may be found moving in any direction with equal probability.

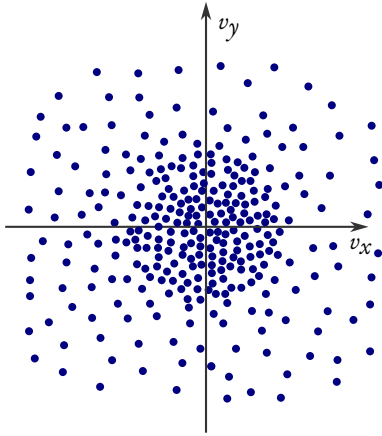


Figure 4.6: A snapshot of molecular velocity distribution.

In order to have a correct picture of the molecular movements in a gas, we should use both figures. It is still better, instead of each figure, to consider a sequence of snapshots taken at regular intervals in time.

We should then see that the points in Figure 4.5 move in different directions: the trajectories change during collisions. The points in Figure 4.6 do not move; however, some suddenly disappear and some appear. Each time a pair of points disappears another pair of new points appears: this is the result of collision between two molecules.

Maxwell's distribution law. Suppose $F(v_x)\Delta v_x$ is the probability that a certain molecule (at a certain moment in time) has an x -velocity component from v_x to $v_x + \Delta v_x$ the other two velocity components taking any arbitrary value. At small Δv_x the function $F(v_x)$ is the density of the probability of finding a molecule with velocity component v_x .

The English physicist James Clerk Maxwell (1831-1879) showed that the probability density $F(v_x)$ corresponds to *Gauss's law*:

$$F(v_x) = A \exp(-\alpha v_x^2), \quad (4.14)$$

where α is a parameter ($\alpha > 0$) and the constant A is determined from

$$\int_{-\infty}^{\infty} F(v_x) dv_x = 1, \quad (4.15)$$

which is a reflection of the fact that the probability of a molecule having an x -component in its velocity is unity. Substituting (4.14) into (4.15), we obtain

$$A \int_{-\infty}^{\infty} \exp(-\alpha v_x^2) dv_x = 1.$$

The integral in this expression is known in mathematics as Poisson's integral and, evaluates to $\sqrt{\pi/\alpha}$. Consequently, $A = \sqrt{\pi/\alpha}$. Thus, we can rewrite (4.14) as

$$F(v_x) = \sqrt{\pi/\alpha} \exp(-\alpha v_x^2). \quad (4.16)$$

Similar functions can be derived for the probability densities for the y - and z -components of a molecule's velocity. The function $F(v_x)$ is plotted in Figure 4.7. Suppose $f(v_x, v_y, v_z)$ is the density of the probability of finding a molecule with velocity components v_x, v_y , and v_z . Using the theorem of probability multiplication, we can write:

$$f(v_x, v_y, v_z) \Delta v_x \Delta v_y \Delta v_z = [F(v_x \Delta v_x)][F(v_y \Delta v_y)][F(v_z \Delta v_z)].$$

Whence

$$f(v_x, v_y, v_z) = \left(\frac{\alpha}{\pi}\right)^{3/2} \exp(-\alpha(v_x^2 + v_y^2 + v_z^2)) = \left(\frac{\alpha}{\pi}\right)^{3/2} \exp(-\alpha v^2). \quad (4.17)$$

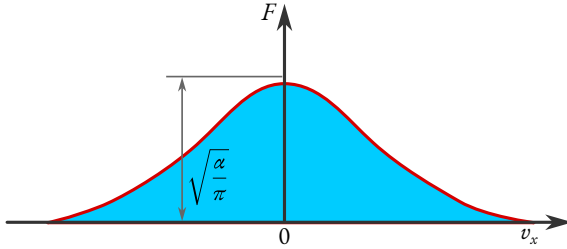


Figure 4.7: The Gauss velocity distribution.

We see that the probability density depends on the squares of the velocity components, viz. $v_x^2 + v_y^2 + v_z^2 = v^2$. This we might have expected because, as it was already noted, each velocity direction is equally probable, and so the probability density may only depend on the absolute value of a molecule's velocity.

Thus, the probability of finding a molecule with velocity components taking the values $v_x - v_x + \Delta v_x$, $v_y - v_y + \Delta v_y$, $v_z - v_z + \Delta v_z$, is:

$$\Delta w_v = \left(\frac{\alpha}{\pi}\right)^{3/2} \exp(-\alpha v^2) \Delta v_x \Delta v_y \Delta v_z, \quad (4.18)$$

where $v^2 = v_x^2 + v_y^2 + v_z^2$.

Let us take one more step: since each velocity direction is equally probable, let us look at the probability of finding a molecule with an absolute velocity from v to $v + \Delta v$, irrespective of its direction. If we consider a velocity space (Figure 4.8), then Δw_v (see (4.18)) is the probability of finding a molecule in the "volume" Δv , shown in (Figure 4.8 (a)) (the word "volume" is enclosed in quotation marks to remind us that we are dealing with a velocity space rather than with a normal space). Now we want

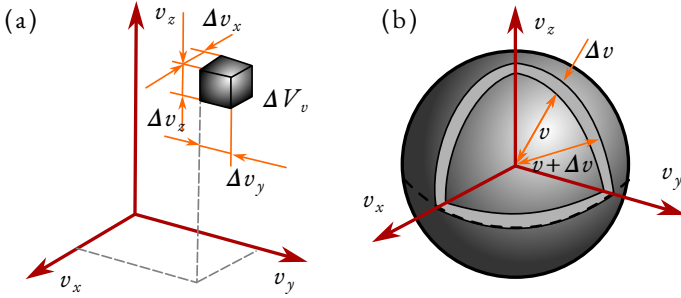


Figure 4.8: The velocity space.

to consider the probability of finding a molecule within the spherical layer shown in Figure 4.8 (b) and confined between spheres with radii v and $v + \Delta v$. The “volume” of this layer is the surface area of a sphere of radius v multiplied by the thickness of the layer Δv , i.e. $4\pi v^2 \Delta v$. Therefore, the probability we want has the form:

$$\Delta w_v = \left(\frac{\alpha}{\pi}\right)^{3/2} \exp(-\alpha v^2) 4\pi v^2 \Delta v. \quad (4.19)$$

This formula expresses the distribution of molecules in an ideal gas by the absolute value of their velocities, i.e. the *Maxwellian distribution*. The probability density $g(v) = \Delta w_v / \Delta v$ is shown in Figure 4.9. It vanishes both when v tends to zero and when it tends to infinity. The “volume” of the spherical layer shown in Figure 4.8 (b) vanishes when v tends to zero and the factor $\exp(-\alpha v^2)$ in the distribution law vanishes when v tends to infinity.

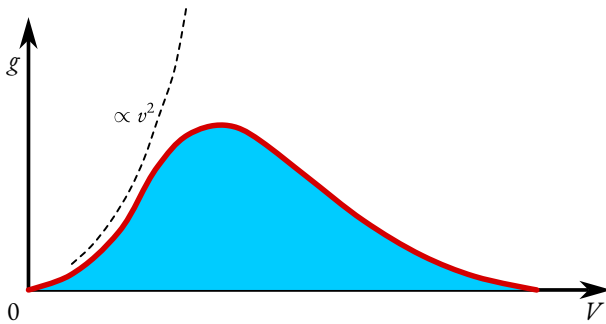


Figure 4.9: The Maxwellian velocity distribution.

Chance and necessity in the pattern of moving molecules. Suppose we could record the position and velocity of every molecule in a volume of gas at some moment in time. Imagine now that we divide the volume into numerous identical cells, and look at our instantaneous “photograph” from cell to cell. It will turn out that the number of molecules varies from cell to cell in a random fashion. Let us only pay attention to those molecules whose velocities are within the range from v to $v + \Delta v$. The number of such molecules varies randomly from cell to cell. Let us divide the solid angle for all space at a point, i.e. 4π steradians, into many identical elementary solid angles. The number of molecules whose velocities lie within an elementary solid angle varies randomly from one such an angle to another.

We could look at the situation in another way, that is, we could focus our attention on some cell or an elementary solid angle and take snapshots at different moments in time. The number of molecules (in a cell or a solid angle) at different times will also randomly change.

To emphasize the *randomness* in the picture of moving molecules, the term “chaotic” is applied: chaotic collisions between molecules, chaotically directed molecule velocities, or generally, the chaotic thermal movement of molecules. However, there is some *order* in this “chaos” or, in other words, *necessity* or what we have repeatedly called *statistical stability*.

The statistical stability shows itself in the existence of definite probabilities: the probability of a molecule being in a volume ΔV (the probability is $\Delta V/V$), the probability of a molecule moving within a solid angle $\Delta\Omega$ (the probability is $\Delta\Omega/4\pi$), and the probability of a molecule having an absolute value of velocity from v to $v + \Delta v$ (the probability is defined by (4.19)).

The number of molecules per unit volume each possessing an absolute value of velocity from v to $v + \Delta v$ is, to a great degree of accuracy,

$$\Delta n = \frac{N}{V} \Delta w_v = 4\pi \frac{N}{V} \left(\frac{\alpha}{\pi} \right)^{3/2} \exp(-\alpha v^2) 4\pi v^2 \Delta v. \quad (4.20)$$

Collisions between molecules push some molecules out of this range of velocity values; however, other collisions bring new molecules into it. So order is maintained: the number of molecules in a given interval of velocity values remains practically constant and is defined by (4.20). Let me emphasize that chance and necessity, as always, are dialectically united here. Collisions among a great number of molecules give the picture of the moving molecules its randomness. But at the same time the collisions maintain the thermodynamic equilibrium in the gas, which is characterized by definite probabilities, and in turn reveals statistical stability.

Pressure and Temperature of an Ideal Gas

Pressure as the result of molecular bombardment. The walls of a vessel containing a gas are continuously struck by gas molecules. This molecular bombardment results in the pressure exerted by a gas on a wall. Let us take an x -axis at right angles to the wall. It is clear from Figure 4.10 (a) that the x -component of a molecule's momentum in an elastic collision with the wall changes by $2m_0v_x$ where m_0 is the mass of the molecule. This means that when it strikes the wall, the molecule gives it an impulse of $2m_0v_x$. Let us first look at those gas molecules whose x -components of velocity lie between v_x and $v_x + \Delta v_x$ (note that $v_x > 0$, otherwise the molecule will be moving away from the wall rather than towards it); the other components of the molecule's velocity are not important. The number of collisions between the molecules in question and an area s of the wall per unit time equals the number of molecules in a volume equal to $s v_x$ Figure 4.10 (b). (The reader should not be confused by the fact that the product $s v_x$ does not have the dimensions of volume. In reality, we deal herewith the product $s(\text{cm}^2) \times v_x(\text{cm s}^{-1}) \times 1(\text{s})$.) Regarding (4.16)), this number of collisions is

$$\Delta R = \frac{N}{V} s v_x F(v_x) \Delta v_x = \frac{N}{V} s v_x \sqrt{\frac{\alpha}{\pi}} \exp(-\alpha v_x^2) \Delta v_x.$$

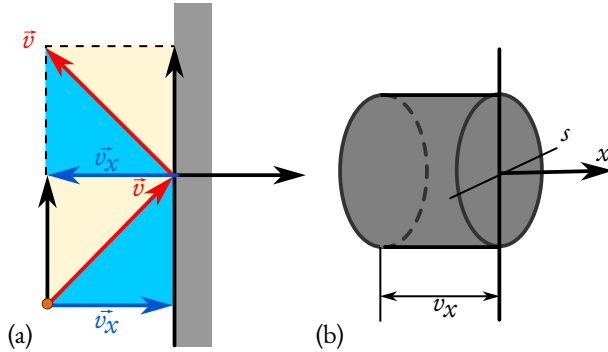


Figure 4.10: The collisions of molecules with walls of the container.

The wall receives an impulse of $2m_0v_x$ at each collision. The force acting on an area s of the wall per unit time is the impulse transferred to the area. Dividing the force by the area s , we can find the pressure exerted by the gas on the wall caused by the molecules whose x -velocity components take values from v_x to $v_x + \Delta v_x$:

$$\Delta p = 2m_0v_x \Delta R \frac{1}{s} = 2m_0 \frac{N}{V} \sqrt{\frac{\alpha}{\pi}} \exp(-\alpha v_x^2) v_x^2 \Delta v_x. \quad (4.21)$$

The only thing left is to sum up, or, more accurately, to integrate (4.21) over all non-negative values of velocity v_x :

$$p = 2m_0 \frac{N}{V} \sqrt{\frac{\alpha}{\pi}} \int_{-\infty}^{\infty} \exp(-\alpha v_x^2) v_x^2 dv_x. \quad (4.22)$$

The following is a standard identity:

$$\int_0^{\infty} \exp(-\alpha v_x^2) v_x^2 dv_x = \frac{1}{4} \sqrt{\frac{\pi}{\alpha^3}}.$$

Therefore,

$$p = m_0 N / 2\alpha V. \quad (4.23)$$

Maxwellian distribution finally becomes clear. We have long tried the reader's patience with the mysterious parameter α . It is clear from (4.23) that $\alpha = m_0 N / 2pV$. Since the gas is in a thermal equilibrium, we can use the Mendelev-Clapeyron equation $pV = mRT/M$. Inasmuch as $R = N_A k$ (N_A is Avogadro's number and k is Boltzmann's constant and equal to $1.38 \times 10^{-23} \text{ J C}^{-1}$), and moreover $N_A m / M = N$, we can rewrite the Mendelev-Clapeyron equation in the form

$$pV = NkT. \quad (4.24)$$

Now we obtain from (4.23) and (4.24)

$$\alpha = \frac{m_0}{2kT}. \quad (4.25)$$

Consequently, (4.19) becomes

$$\Delta w_v = g(v) \Delta v = 4\pi \left(\frac{m_0}{2\pi kT} \right)^{3/2} \exp \left(-\frac{m_0 v^2}{2kT} \right) v^2 \Delta v. \quad (4.26)$$

Temperature as a measure of mean molecular energy. The mean value of the squared velocity of molecules in an ideal gas can be found using (1.17) and (4.26):

$$E(v^2) = \int_0^{\infty} v^2 g(v) dv = 4\pi i \left(\frac{m_0}{2\pi kT} \right)^{3/2} \int_0^{\infty} \exp \left(-\frac{m_0 v^2}{2kT} \right) v^4 dv. \quad (4.27)$$

Another standard integral is

$$\int_0^{\infty} \exp(-\alpha v^2) v^4 dv = \frac{3}{8} \sqrt{\frac{\pi}{\alpha^5}}.$$

whence we obtain from (4.27):

$$E(v^2) = \frac{3}{2\alpha} = \frac{3kT}{m_0}. \quad (4.28)$$

If we apply the model of an ideal gas, we can neglect the energy of the collisions between the molecules as compared with their kinetic energy, i.e. we can present the energy of a molecule as $\varepsilon = m_0 v^2/2$. From (4.28) we find the following expression for the mean energy of a molecule in an ideal gas:

$$E(\varepsilon) = \frac{m_0}{2} E(v^2) = \frac{3}{2} kT. \quad (4.29)$$

Therefore, we see that the temperature can be considered as a *measure of the mean energy of a molecule*.

It follows from (4.29) that the *internal energy* of an ideal gas in equilibrium and containing N molecules and possessing temperature T is

$$U = \frac{3}{2} NkT. \quad (4.30)$$

Molecular kinetics has allowed us to explain why the internal energy of an ideal gas is proportional to its absolute temperature and does not depend on the volume occupied by the gas. We have used this fact while considering some problems of thermodynamics.

Fluctuations

Fluctuations of micro-variables and macro-variables. Let us call the variables governing a particular molecule *micro-variables* and those governing a macroscopic body, for instance, a gas as a whole, *macro-variables*. The velocity v and energy ε of a molecule, are micro-variables; while the internal energy of a gas U , temperature T , and pressure p are macro-variables.

Let us imagine that we are following the energy of a molecule in a gas. The energy varies randomly from collision to collision. Knowing the function $\varepsilon(\tau)$ for a long enough time interval τ , we can find the mean value of the molecule's energy:

$$E(\varepsilon) = \frac{1}{\tau} \int_0^\tau \varepsilon(t) dt. \quad (4.31)$$

Recall that we approached the notion of mean energy in another manner in the section Pressure and Temperature of an Ideal Gas. Instead of following the energy of a molecule

during a time interval, we recorded the instantaneous energies of all the molecules and divided the sum by the number of molecules; this is the idea behind equation (4.27). It can be said that here we regarded *averaging over the collective (ensemble) of molecules*. Now (4.31) corresponds to *averaging over time*. Both lead to the same result.

However, let us return to the energy of a molecule in a gas. In the course of time, the energy $\varepsilon(t)$ varies randomly, or rather it fluctuates around a mean value $E(\varepsilon)$. In order to select a measure for the deviation of energy from the mean value, we choose the variance

$$\text{var } \varepsilon = E(\varepsilon^2) - (E(\varepsilon))^2. \quad (4.32)$$

The variance $\text{var } \varepsilon$ is called the *quadratic fluctuation* of energy ε . Once we know the distribution of molecules by velocities, we can calculate $E(\varepsilon^2)$ thus:

$$E(\varepsilon^2) = \int_0^\infty \left(\frac{m_0 v^2}{2} \right)^2 g(v) dv. \quad (4.33)$$

By substituting here the probability density $g(v)$ from (4.26), we can find (the mathematical calculations are omitted for simplicity's sake):

$$E(\varepsilon^2) = \frac{15(kT)^2}{4}. \quad (4.34)$$

From (4.29) we obtain

$$\text{var } \varepsilon = E(\varepsilon^2) - (E(\varepsilon))^2 = \frac{3}{2}(kT)^2. \quad (4.35)$$

The ratio of the square root of the quadratic fluctuation to the mean value of a variable is called its *relative fluctuation*. The relative fluctuation of the energy is approximately unity:

$$\xi = \frac{\sqrt{\text{var } \varepsilon}}{E(\varepsilon)} = \sqrt{\frac{2}{3}}. \quad (4.36)$$

The amplitude of a micro-variable's fluctuation proves to be of the same order as its mean value.

Now let us consider the fluctuation of a macro-variable, for instance, the internal energy of the gas consisting of N monoatomic molecules. Suppose $U(t)$ is the instantaneous value of the gas internal energy at time t :

$$U(t) = \sum_{i=1}^N \varepsilon_i(t). \quad (4.37)$$

The values of $U(t)$ fluctuate around mean value $E(U)$. The fluctuations of the gas internal energy can be related to the chaotic elementary exchanges of energy between the gas molecules and the vessel wall. Since the mean of a sum is the sum of the means, we have

$$E(U) = \sum_{i=1}^N E(\varepsilon) = NE(\varepsilon). \quad (4.38)$$

We have made use of the fact that the mean energy is the same for any molecule.

Let us first write the variance $\text{var } U$ in the form

$$\text{var } U = E(U^2) - (E(U))^2 = E((U(t) - E(U))^2).$$

We shall use δU to denote the difference $U(t) - E(U)$,

$$\text{var } U = E(\delta U)^2. \quad (4.39)$$

Using (4.37) and (4.38), we can find:

$$\begin{aligned} \delta U &= U(t) - E(U) = \sum_{i=1}^N \varepsilon_i(t) - NE(\varepsilon) \\ &= \sum_{i=1}^N (\varepsilon_i(t) - E(\varepsilon)) = \sum_{i=1}^N \delta \varepsilon_i. \end{aligned}$$

Therefore,

$$\text{var } U = E\left(\sum_{i=1}^N \delta \varepsilon_i\right)^2. \quad (4.40)$$

Thus we have to square the sum of N terms and then average each of the resultant terms. Squaring a sum of N terms yields N terms of the form $(\delta \varepsilon_i)^2$ ($i = 1, 2, \dots, N$), which after averaging yield $NE(\delta \varepsilon)^2$. In addition, squaring a sum of N terms generates a number of what are usually called cross-terms, i. e. terms of the form $2\delta \varepsilon_i \delta \varepsilon_j$ where $i \neq j$. Each of these terms will vanish after averaging. Indeed, $E(\delta \varepsilon_i \delta \varepsilon_j) = E(\delta \varepsilon_i) E(\delta \varepsilon_j)$. As to the averaged terms $E(\delta \varepsilon_i)$ and $E(\delta \varepsilon_j)$, they vanish too because a variable is equally likely to deviate from its mean on either side. Thus,

$$\text{var } U = NE(\delta \varepsilon)^2 = N \text{var } \varepsilon. \quad (4.41)$$

Using (4.35) we can obtain the following expression for the quadratic fluctuation of the gas internal energy:

$$\text{var } U = \frac{3}{2}N(kT)^2. \quad (4.42)$$

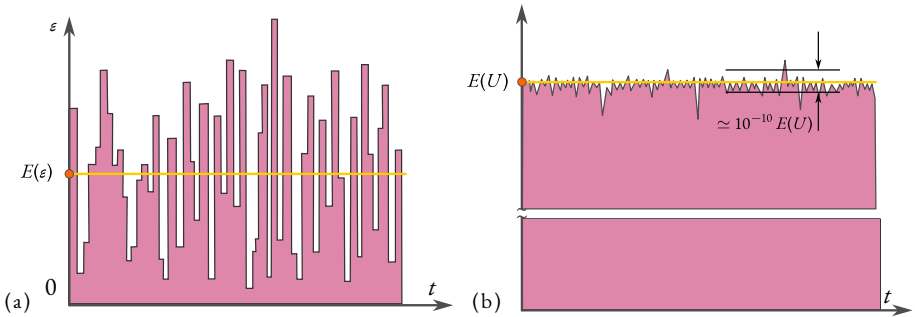


Figure 4.11: A comparison of the fluctuations of the micro-variable ε and macro-variable U .

The relative fluctuation of the internal energy is

$$\xi = \frac{\sqrt{\text{var } U}}{E(U)} = \sqrt{\frac{2}{3}} \frac{1}{\sqrt{N}}. \quad (4.43)$$

We can see, therefore, that the relative fluctuation of the internal energy of a gas of N molecules is proportional to $1/\sqrt{N}$, i.e. it is very small (recall that a cubic centimetre of a gas contains about 10^{19} molecules at normal pressure). In fact, $\xi \propto 1/\sqrt{N}$ for all macro-variables, which allows us to neglect their fluctuations for all practical purposes, and to regard the mean values of macro-variables as the true values. The fluctuations of the micro-variable ε and macro-variable U are compared in Figure 4.11.

Thus, the total internal energy U is not a fixed value for an equilibrium state of a macroscopic body. It varies slightly in time, going through small fluctuations around its mean value. Temperature, pressure, and entropy fluctuate around their mean values too.

Brownian movement. Having seen (4.43), a reader may conclude that under ordinary conditions, i.e. when we deal with macroscopic bodies and the macro-variables characterizing them, fluctuations do not show themselves. However, we can actually observe fluctuations by eye. Consider the *Brownian movement* as an example.

In 1827, the English biologist Robert Brown (1773-1858) used a microscope to study small particles (plant pollen) suspended in water. He discovered that they were in constant chaotic motion. He was sure that this movement was due to the particles themselves rather than a result of flows in the liquid or its evaporation.

A correct explanation of Brownian movement was given in 1905 by Albert Einstein

(1879-1955). He showed that the cause of the Brownian movement is the chaotic bombardment of the small suspended particles by the molecules of the surrounding liquid.

Imagine a small disc with a diameter of, for instance, 10^{-4} cm suspended in a liquid. The number of collisions between the liquid molecules and one side of the disc per unit time equals, on average, the number of collisions on the other side. But this is only on the average. In reality, the number of collisions on one side of the disc during a small interval of time may be noticeably greater than the number of collisions on the other side. The result is that the disc receives an overall unbalanced impulse and so moves in the appropriate direction. We can say that the disc moves because of the *fluctuations in the pressure* exerted by the liquid molecules on the two sides of the disc.

Einstein considered a concrete physical model with a ball as a Brownian particle. He showed that the mean square of the displacement of such a particle during an observational period τ is defined by the following formula

$$E(l^2) = \frac{\tau}{8\pi\eta r} kT, \quad (4.44)$$

where r is the ball's radius, η is the viscosity coefficient of the liquid, T is its temperature.

Why the sky is blue. The colour of the sky is due to the diffusion of sunlight through the Earth's atmosphere. Let us imagine the atmosphere to be separated into a great number of small cubic cells each with an edge a wavelength of light long (about 0.5×10^{-4} cm). The chaotic motion of the air molecules results in that the number of molecules within the cell varies randomly from cell to cell. It will also vary randomly within a cell if we observe it at different instants in time. Sunlight diffuses through these *fluctuations of air density*.

The intensity ΔI of light diffused through a volume of air ΔV at distance R from the observer is defined by the relationship

$$\Delta I = a \frac{\Delta V}{R^2} \frac{1}{\lambda^4} kT, \quad (4.45)$$

where λ is the light wavelength, T is the air temperature, and a is a factor we shall not deal with here. It is clear from (4.45) that the shorter the wavelength the more light diffuses ($\Delta I \propto \lambda^4$). Therefore, the spectrum of the light which diffuses through the Earth's atmosphere proves to have a peak at the *shortwave end*, which explains why the sky is *blue*.

The Nyquist formula. It follows from Ohm's law that if there is no electromotive force in an electric circuit, there is no current in it. However, this is not quite true. The

point is that fluctuations related to the thermal movement of electrons in a conductor result in fluctuating currents, and hence a fluctuating electromotive force. In 1927, the American physicist and engineer Harry Nyquist (1889-1976) showed that if there is a conductor with resistance R and temperature T , a *voltage fluctuation* δV appears at the ends of the resistor, the mean square of the fluctuation being

$$E(\delta V)^2 = 4RkT\Delta\nu, \quad (4.46)$$

where $\Delta\nu$ is the range of frequencies within which the voltage fluctuations are measured.

Fluctuating electrical variables play an essential role in modern technology. They are, in principle, an unavoidable source of noise in communication channels and define the sensitivity limits of measuring instruments. Besides fluctuations caused by the thermal motion of electrons in conductors, let me mention another essential type of fluctuation, the fluctuation in a number of electrons leaving the heated cathode of an electron tube.

Fluctuations and temperature. I would like to draw the reader's attention to expressions (4.35) and (4.35). It is clear that a quadratic fluctuation is related to the absolute temperature: $\sqrt{\text{var}} \propto T$. The same result can be derived from formulas (4.44)-(4.46). The relation between the quadratic fluctuation of a physical variable and temperature has a deep meaning. The greater the temperature of a body the more a physical parameter will fluctuate.

We noted above that the temperature of a body can be regarded as a measure of the average energy of the body's particles. Recall that this is only valid if the body is in thermal equilibrium. If an ensemble of particles is very far from equilibrium (suppose we are discussing a cosmic shower or the beam of particles from an accelerator), then the average energy of the particles cannot be measured by temperature. A more general approach to the notion of a body's temperature is its relation with the fluctuations of its physical parameters rather than the average energy of its particles. Temperature can be regarded as a measure of fluctuation. By measuring the fluctuations, we can measure the absolute temperature of the body in principle. The fluctuations in the electrical variables suit this purpose best.

The relationship between temperature and fluctuations indicates, in particular, that the notion of temperature, strictly speaking, has no analogue in Newtonian mechanics. Temperature involves probabilistic processes and is a measure of the variance of random variables.

Entropy and Probability

From the formula of the work done by a gas during an isothermal expansion to Boltzmann's formula. Suppose an ideal gas with mass m and temperature T expands isothermally from volume V_1 to volume V_2 . According to (4.6), the work performed by the gas during the expansion is $(mRT/M) \ln (V_2/V_1)$. During an isothermal expansion, the work is done due to a quantity of heat Q drawn by the gas from the environment. Therefore,

$$Q = \frac{mRT}{M} \ln \left(\frac{V_2}{V_1} \right). \quad (4.47)$$

Using (4.24) for the equation of state of an ideal gas, we can transform (4.47) into

$$Q = NkT \ln \left(\frac{V_2}{V_1} \right), \quad (4.48)$$

where N is the number of molecules in the gas. Taking into account (4.10), we can conclude that the increment of entropy in the gas is

$$\Delta S = Nk \ln \left(\frac{V_2}{V_1} \right). \quad (4.49)$$

The isothermal expansion of a gas is a *reversible* process. The increase of entropy in a reversible process should not surprise the reader: we consider the entropy of a gas, and the gas here is an open system (it performs work on a piston or draws heat from an external body). The same increase in entropy is observed in an *irreversible* process of gas expansion from V_2 to V_1 when the gas is a closed system. This irreversible process can be carried out as follows. Suppose that a thermally insulated vessel of volume V_0 has a partition, and first all the gas is on one side of the partition and occupies volume V_1 . Then the partition is removed and the gas expands into vacuum. The expansion is considered to start when the partition is removed and to end when the gas occupies volume V_2 . The increment in the gas's entropy during this process is also defined by formula (4.49).

Using the example of gas expansion into a vacuum, we can explain the increase in entropy on the basis of *probabilities*. The probability that a gas molecule occurs in volume V_1 is evidently equal to V_1/V_0 . The probability that another molecule will occur in volume V_1 simultaneously with the first one is $(V_1/V_0)^2$. The probability that all N molecules will gather in volume V_1 is $(V_1/V_0)^N$. Let us use w_1 to denote the probability that all molecules are in volume V_1 and w_2 to denote the probability that all molecules will occur in volume V_2 . The first probability is $(V_1/V_0)^N$ while the second one is $(V_2/V_0)^N$.

Therefore,

$$\frac{w_2}{w_1} = \left(\frac{V_2}{V_1} \right)^N. \quad (4.50)$$

We can therefore obtain from (4.49):

$$\Delta S = Nk \ln \left(\frac{V_2}{V_1} \right) = k \ln \left(\frac{V_2}{V_1} \right)^N = k \ln \left(\frac{w_2}{w_1} \right). \quad (4.51)$$

Thus, using rather simple reasoning, we have arrived at an essential result, namely Boltzmann's formula.

Boltzmann's formula. In 1872, Ludwig Boltzmann (1844-1906) published a formula in which the entropy of a system in a certain state is proportional to the *logarithm of the probability of the state*. The proportionality factor in this formula was refined later and was called *Boltzmann's constant*. Boltzmann's equation is now given as

$$S = k \ln w. \quad (4.52)$$

Formula (4.51) is obtained from (4.52) if we assume that $S_1 = k \ln w_1$, $S_2 = k \ln w_2$, and $\Delta S = S_2 - S_1$.

Suppose a system consists of two subsystems, one of which is in state 1 with entropy S_1 and probability w_1 and the other is in state 2 with entropy S_2 and probability w_2 . Let S and w be the entropy and the probability of the entire system's state, respectively. Entropy is additive, and therefore

$$S = S_1 + S_2. \quad (4.53a)$$

This state is realized when the first subsystem is in state 1 and the second subsystem is in state 2 at the same time. According to the theorem of probability multiplication,

$$w = w_1 \cdot w_2. \quad (4.53b)$$

It is clear that (4.53a) and (4.53b) are in agreement with Boltzmann's formula:

$$S = k \ln(w_1 w_2) = k \ln w_1 + k \ln w_2 = S_1 + S_2.$$

Macro-states and micro-states. Now what is the "probability of the system's state"? Consider a simple system consisting of four particles, each of which may be in either of two states with equal probability. We can imagine a vessel divided into two equal parts (left and right) and only four molecules inside the vessel. Each of the

molecules may be found in the left or right half with equal probability. This system has five possible macro-states: 1, there are no molecules in the left half; 2, there is one molecule in the left half; 3, there are two molecules in the left half; 4, there are three molecules in the left half; and 5, there are four molecules in the left half. These macro-states may be realized by *different numbers of equally probable ways*, or, in other words, different macro states correspond to different numbers of micro-states. This is clear from Figure 4.12, where different colours are used to mark the different molecules. We can see that macro-states 1 and 5 may only occur in one way each. Each therefore corresponds to one micro-state. Macro-states 2 and 4 correspond to four micro-states. Macro-state 3 corresponds to six equally probable micro-states. There can be 16 equally probable micro-states in all. The probability of a macro-state is *proportional to the number of corresponding micro-states*, and this is the probability involved in Boltzmann's formula. The number of micro-states corresponding to a given macro-state is called its *statistical weight*.

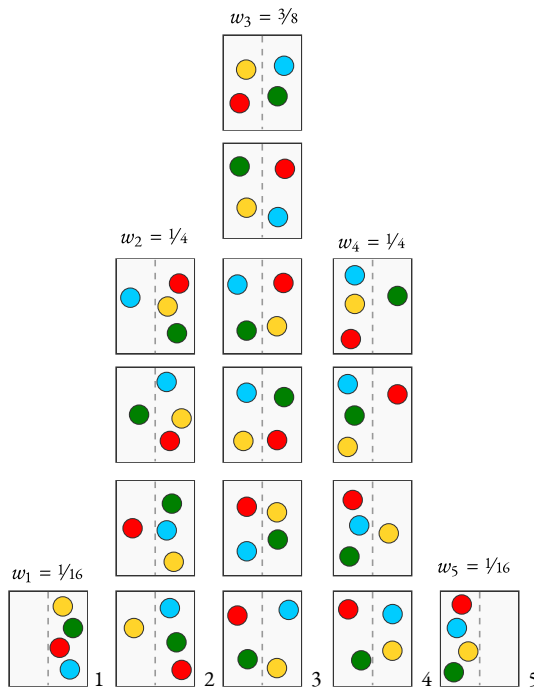


Figure 4.12: Entropy as a measure of disorder in the system.

Suppose that there are N molecules rather than four in the vessel divided into two equal halves. Now there are $N + 1$ macro-states, which can be conveniently designated by the numbers $0, 1, 2, 3, \dots, N$, according to the number of molecules present, say, in the left half. The statistical weight of the n^{th} macro-state equals the number of combinations of N things taken n at a time:

$$\binom{N}{n} = \frac{N!}{(N-n)!n!}. \quad (4.54)$$

This is the number of micro-states corresponding to the n^{th} macro-state.

The total number of micro-states is defined by the sum

$$\sum_{n=0}^N \binom{N}{n}$$

The probability of the n^{th} macro-state is

$$w_n = \binom{N}{n} / \sum_{n=0}^N \binom{N}{n} \quad (4.55)$$

An example using Boltzmann's formula. Suppose a gas consisting of N molecules expands into vacuum. Its volume doubles. Find the increase in the gas's entropy.

The *initial* state of the gas is the macro state with $n = 0$ (all molecules are in the right half of the vessel), and the final state is the macro-state with $n = N/2$ (the molecules are uniformly distributed between both halves of the vessel, which means the volume of the gas has doubled). Here we assume that N is an even number (this reservation is not essential for large N). In agreement with (4.54) and (4.55), we can write:

$$\frac{w_{N/2}}{w_0} = \binom{N}{N/2} / \binom{N}{0} = \binom{N}{N/2} = \frac{N!}{(N/2)!(N/2)!}. \quad (4.56)$$

According to Boltzmann's formula, the increase in the gas's entropy is

$$\Delta S = k \ln \frac{w_{N/2}}{w_0} = k \ln \frac{N!}{(N/2)!(N/2)!}, \quad (4.57)$$

Since N is a very large number, we can use the approximation

$$\ln(N!) = N \ln N, \quad (4.58)$$

hence (4.57) takes the form

$$\Delta S = kN \ln 2. \quad (4.59)$$

The same result follows from (4.49) if we assume $V_2/V_1 = 2$.

Entropy as a measure of disorder in a system. Let us return to Figure 4.12. Macro-states 1 and 5 clearly show the structure of the system, its separation into two halves. There are molecules in one half and no molecules in the other. On the contrary, macro-state 3 does not have this structure at all because the molecules are evenly distributed in both halves. The presence of a definite structure is related to the *order* in a system while the absence of structure is related to *disorder*. The greater the degree of order in a macro-state, the smaller its statistical weight (i.e. the number of corresponding micro-states is smaller). Disordered macro-states with no inner structure have large statistical weight. They can be realized in many ways, in other words, by many micro-states.

All this allows us to regard entropy as a *measure of disorder in a system*. If the disorder in a given macro-state is large, its statistical weight is large, and therefore, its entropy is large.

A statistical explanation of the second law of thermodynamics. Boltzmann's formula makes it possible to explain the increase in entropy during irreversible processes in a closed system as postulated by the second law of thermodynamics. The *increase in entropy* means the transition of the system from a *less probable* state to a *more probable* one. The example of gas expanding into vacuum illustrates this. While the gas expands, the system moves from a less probable to a more probable macro-state.

Any process in a closed system proceeds in a direction such that the system's entropy does not decrease. This means that transitions to more probable states or, at least, transitions between equally probable states correspond to real processes.

When a probabilistic approach is used entropy becomes a measure of the disorder in a system. The law requiring the increase of the entropy in a closed system is, therefore, a law which demands that the *degree of disorder* in these systems increases. In other words, a transition from a less probable to a more probable state corresponds to an order-disorder transition. For instance, when a hammer strikes an anvil, the ordered component of the hammer's molecular movement related to its overall downward movement is transformed into the disordered thermal molecular movement of the anvil and the hammer.

The *quantity* of energy in a closed system does not vary in time. However, the *quality* of the energy varies. In particular, its capacity to perform usable work decreases. The increase of entropy in a closed system is, in its essence, a gradual destruction of the

system. Any closed system is unavoidably disordered and degraded as time passes. The isolation of a system subjects it to the power of destructive chance, which always sends the system into disorder. As the French scientist Leon Brillouin once said, “the second law of thermodynamics means death due to isolation”.

Maintaining or, moreover, increasing the order in a system requires that the system be *controlled*, for which it is necessary, first of all, that the system should *not be isolated* or closed. Naturally, when the system loses its “protecting envelope”, it is open to external disorganizing factors. However, it also becomes available to control factors. The action of the latter can decrease the system’s entropy. Of course, this does not contradict the second law of thermodynamics: the decrease of entropy is local in nature, only the entropy of the given system decreases. This decrease is more than compensated by an increase in the entropy in other systems, in particular, those that control the given system.

Fluctuations and the second law of thermodynamics. The probabilistic approach both explained the second law of thermodynamics and showed that the demands of this law are not absolute. The direction in which a process must proceed is dictated by the second law, but it is not strictly predetermined. It is only the *most probable* direction. In principle, violations of the second law of thermodynamics are possible. However, we do not observe them because their *probability is low*.

A gas expands into vacuum spontaneously. This is the most probable direction of the process. However, there is another possible situation, viz. the velocities of the molecules in the gas suddenly point in directions such that the gas spontaneously compresses. This situation has an exceptionally low probability because of the enormous number of molecules in any macro-volume of gas. The spontaneous compression of the gas should be regarded as a fluctuation of its density. If the number of molecules in the gas is large, then, as is known, the characteristic value of the relative fluctuation is small (recall that it is proportional to $1/\sqrt{N}$), and therefore, it is very improbable that a fluctuation on the scale of the macrocosm would be observed.

Suppose a phenomenon requires the participation of a relatively small number of molecules. Then it is not difficult to observe various kinds of fluctuations that violate the second law of thermodynamics. In the preceding section, we discussed density fluctuations in air inside volumes whose linear dimensions are comparable to the light wavelengths. These fluctuations appear as spontaneous compressions and rarefactions in the air, bringing about the blue colour of the sky.

It is most probable for a Brownian particle to collide with the same number of liquid molecules on both sides per unit time. However, because of the small dimensions of

the Brownian particle, fluctuations of pressure due to unbalanced number of collisions from different directions are quite probable such that the particle will randomly move. A moving Brownian particle demonstrates the spontaneous transformation of heat taken from a liquid into the kinetic energy of the particle's motion.

Therefore, we see that the probabilistic explanations of entropy and the second law of thermodynamics help comprehend more deeply the nature of processes in macro-systems. The probabilistic approach explains the puzzles thermodynamics could not solve and, moreover, indicates that the second law of thermodynamics itself has the *probabilistic nature* because it is only valid on the average, and various fluctuations violate this law of thermodynamics. We come to an essential conclusion:

probabilistic laws rather than strictly deterministic ones underlie the second law of thermodynamics.

Entropy and Information

The relation between entropy and information. It was shown in Chapter 3 that the notion of information is underlain by probability. Now we have seen that probability is the basis of entropy. The unity of the nature of *information* and *entropy* proves to be essential. An increase in the entropy of a system corresponds to its transition from a less ordered state to a more ordered one. This transition is accompanied by a decrease in the information contained in the structure of the system. Disorder and uncertainty can be regarded as a lack of information. In turn, information is nothing else but a decrease in uncertainty.

According to the second law of thermodynamics, the entropy of a closed system increases in time. This process corresponds to the loss of information due to random factors, as was considered in Chapter 3. Fluctuations in physical parameters cause random violations of the second law of thermodynamics. Random decreases of entropy are observed. These processes correspond to the generation of information from noise which we discussed above. By influencing the system in a certain way, we can decrease its entropy (by increasing the entropy of another system). This is the process of control, which demands definite information. All this speaks in favour of relation between information and entropy. The Hungarian physicist Leo Szilard (1898-1964) was first to indicate this relation, doing so in 1929.

Thus, *entropy is a measure of disorder* and uncertainty in a system, and *information*

is a measure of order and structural certainty. An increase in information corresponds to a decrease in entropy and, vice versa, a decrease in information corresponds to an increase in entropy.

Boltzmann's formula and Hartley's formula. We came across Hartley's formula in Chapter 3 (see (3.1)). According to this formula, the information required to indicate which of N_1 equally probable outcomes is wanted is $I = \log_2 N_1$. Suppose N_1 is the number of railroad tracks at a station. The signalman has to send a signal indicating the track along which the train is to approach the station. Sending the signal, the signalman selects from N_1 equally probable outcomes. This signal contains $I_1 = \log_2 N_1$ bits of information. Now suppose that some of the tracks must be repaired, so that the signalman must select from N_2 outcomes ($N_2 < N_1$). Now his signal contains information $I_2 = \log_2 N_2$. The difference

$$\Delta I = I_1 - I_2 = \log_2 \left(\frac{N_1}{N_2} \right), \quad (4.60)$$

is information about the repair of the tracks. In other words, this is the information required to decrease the number of equally probable outcome from N_1 to N_2 .

Let us compare the existence of N equally probable outcomes with the presence of N equally probable micro-states, i.e. with the statistical weight N of a certain macro-state. According to Boltzmann's formula, a decrease in the statistical weight of a macro-state from N_1 to N_2 means that the system's entropy is incremented by

$$\Delta S = -k \ln \left(\frac{N_1}{N_2} \right). \quad (4.61)$$

I used a minus sign here because the entropy decreases (the increment is negative) as the statistical weight decreases. In compliance with (4.60), to realize this negative entropy requires an increment in the information of $\Delta N = I_1 - I_2 = \log_2 (N_1/N_2)$. Comparing (4.60) with (4.61) and given that

$$\log_2 \left(\frac{N_1}{N_2} \right) = \frac{\ln (N_1/N_2)}{\ln 2}.$$

Therefore, an increment in the information ΔI corresponds to a decrease in the system's entropy of $\Delta I k / \ln 2$.

Norbert Wiener called information to be negative entropy. Louis Brillouin suggested using the term "negentropy" rather than "negative entropy".

Maxwell's demon and its exorcism. In 1871, Maxwell formulated the following paradox. Suppose a vessel with a gas is separated into two halves (A and B) by a partition

with a trapdoor over a microscopic hole in it. And suppose, Maxwell continued, a “being” (Maxwell called it a “demon”) controls the trapdoor causing it to close and open the hole so as to let the fastest molecules from the A half of the vessel enter the B half and to let the slowest molecules from the B half into the A half. Thus, the demon would increase the temperature in the B half and decrease it in the A half without doing any work, which evidently contradicts the second law of thermodynamics. When looking

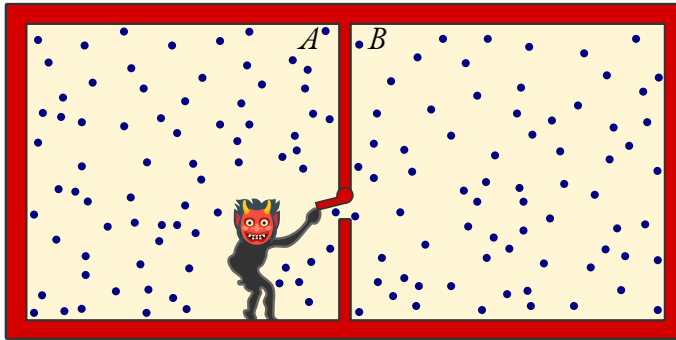


Figure 4.13: Maxwell’s demon controls the flow of molecules from one section to another.

at the illustration of Maxwell’s demon (Figure 4.13), the reader clearly should not think of an evil force. The point of contention is a device that opens and closes a hole in the way the demon described above would act.

Three types of device could be suggested in principle. The first type would be a device controlled by the gas molecules present in the vessel. Imagine there is a one-way trapdoor which responds to the energy of the molecules striking it: fast molecules open the door and slow ones do not. So that it open when struck by an individual molecule, the door would have to be exceedingly light. However, such a door, if it could be produced, would be unable to carry out the functions of the demon. The door would in fact be affected both by the fluctuations due to the motion of the gas molecules and by the fluctuations related to the thermal motion of the molecules of the material making up the door. The door would therefore operate chaotically and would not sort molecules by speed.

The second type of demon would be a device controlled from the outside. Suppose we could monitor the molecules arriving at the hole in the partition. The monitoring device would signal at the right moment and the trapdoor would open or close. If we ignore the technical problems, we might have to admit that this way of sorting the

molecules is possible in principle. However, it will not be a substitute for Maxwell's demon because the latter should work in a *closed* system.

This is essential because it is a decrease in the entropy of a closed system that violates the second law of thermodynamics. But our system is open, the "demon" obtaining information from the outside. The reception of information must be regarded as an inflow of negative entropy (negentropy) into the system, which is equivalent to a decrease in the system's entropy.

There is one more type of the demon, an *intelligent* demon. However, such a demon would not be what we are looking for because, as Einstein said, an intelligent mechanism cannot act in an equilibrium medium. In other words, life and intelligence are impossible in a closed system, that is in a state of equilibrium.

Entropy and life. A living organism is a very ordered system with low entropy. The existence of living organisms suggests a continuous maintenance of the system's entropy at a low level, a continuous reaction to disordering factors, and, in particular, the factors causing diseases. It may seem that an organism does not obey the demands of the second law of thermodynamics.

Naturally, this is not so. We should take into account that any organism is an *open* system in an essentially *non-equilibrium* state. This system actively interacts with its environment, continuously drawing negentropy from it. For instance, it is well-known that food has lower entropy than waste.

Man does not just live. He works, creates, and therefore, actively decreases entropy. All this is only possible because man obtains negentropy (information) from the environment. It is supplied to him via two different channels. The first one is related to the process of learning. The second channel is related to physiological processes of metabolism occurring in the "man-environment" system.

Chapter 5

Probability in the Microcosm

To date quantum theory led us to a deeper comprehension: it established a closer relation between statistics and the fundamentals of physics. This is an event in the history of human thought, whose significance is beyond science itself.

M. Born

Quantum mechanics allowed us to postulate the existence of primary probabilities in the laws of nature.

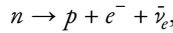
W. Pauli

Spontaneous Micro-processes

Classical physics proceeded from that randomness only reveals itself in large collections, for instance, in ensembles of molecules, in appreciable volumes of gas. However, classical physics did not see randomness in the behaviour of individual molecules. The investigations resulting in the appearance and development of *quantum physics* showed that this viewpoint was invalid. It turned out that randomness is seen both in ensembles of molecules and in the behaviour of individual molecules. This is demonstrated by *spontaneous* microprocesses.

Neutron decay. A typical example of a spontaneous microprocess is the decay of a free neutron. Usually, neutrons are in a bound state. Together with protons,

they are the “bricks” from which atomic nuclei are built. However, neutrons can also be observed outside nuclei, in the free state. For instance, free neutrons appear when uranium nuclei split. It turns out that a free neutron can *randomly, without any external influence*, transform into three particles: a proton, an electron, and an antineutrino (more accurately, an electron antineutrino). This transformation is called neutron decay, and it is commonly written down as:



where n is a neutron, p is a proton, e^- an electron, and $\bar{\nu}_e$ is an antineutrino. Note that the term “decay” is not entirely suitable here because it conveys the idea that a neutron consists of a proton, electron, and antineutrino. In reality, all three particles are born at the moment the neutron annihilates, and it is no use looking for them “inside” the neutron.

The very fact of spontaneous neutron decay is *random*, but there is also a dialectic necessity here as well. In order to reveal it, we should consider a large number of neutrons. Suppose there are N_0 neutrons in a volume at moment $t = 0$, where $N_0 \gg 1$. Let us measure the number of neutrons in the volume at different moments t , the result being a function $N(t)$ whose plot has a certain shape (Figure 5.1). The resultant function is

$$N(t) = N_0 \exp(-at), \quad (5.1)$$

where a is a constant and is commonly given as $1/\tau$, measurements show that $\tau = 10^3$ s.

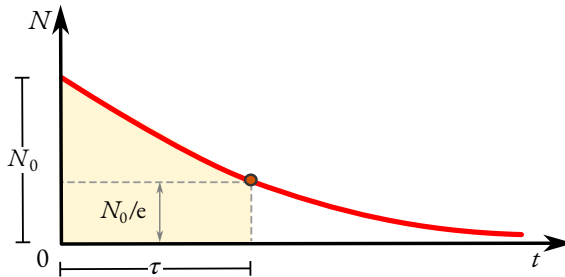


Figure 5.1: The number of neutrons decaying as a function of time.

The value τ is called the neutron’s *lifetime*. It is called this conventionally not because it is the true lifetime of a neutron, but because it is the time needed for the number of intact (un-decayed) neutrons to decrease e times. Whence from (5.1) we have

$$\frac{N(t)}{N_0} = \exp\left(-\frac{t}{\tau}\right) = \frac{1}{e}.$$

The true lifetime of a neutron may vary considerably from τ in both directions. It is in principle impossible to predict when a neutron will decay. We can only consider the *probability* that a neutron will live a while until it decays. When the number of neutrons is large, the ratio $N(t)/N_0$ is the probability that a neutron will survive for a time t . It follows from (5.1) that this probability is $\exp(-t/\tau)$.

I would like to draw your attention to an interesting detail. When we discuss the probability that a neutron will survive for a time t , we do not suppose that this interval is measured from the moment of the neutron's birth. It is not essential how long a neutron has lived by $t = 0$. The probability that it will survive a further time t is always equal to $\exp(-t/\tau)$. It can be said that neutrons "do not get old". This means that there is no meaning in looking for the cause of the neutron's decay within its "internal mechanism".

It is interesting that (5.1), which expresses a certain necessity, is nothing but a direct consequence of the fact that the decays occur independently and randomly. Since the decay is a random process, the decrease in the number of neutrons (in other words, the number of decays) ΔN during an interval of time from t to $t + \Delta t$ is proportional to the number of neutrons $N(t)$ at that instant and the lapse of time Δt , i.e. $\Delta N = -a N(t) \Delta t$. Let us rewrite this equality as $\Delta N / \Delta t = -a N(t)$. In the limit as $\Delta t \rightarrow 0$, we obtain a differential equation known as the *equation of exponential decay*:

$$\frac{dN}{dt} = -a N(t). \quad (5.2)$$

The function (5.1) is the solution of this equation given the initial condition $N(0) = N_0$.

In conclusion, let me remark that if a neutron is not free but is bound with protons and other neutrons in an atomic nucleus, it loses its ability to decay. However, it regains this ability in some cases. The phenomenon of beta decay is then observed (we shall discuss it below).

The instability of elementary particles. The neutron is not at all the only elementary particle that turns spontaneously into other particles. Most elementary particles possess this property, which might be called *instability*. There are only several particles that are stable: the photon, the neutrino, the electron, and the proton.

The instabilities of different particles teach us additional things of randomness. For instance, let us take the particle called the sigma-plus-hyperon Σ^+ . It has a positive electric charge equal in its absolute value to the charge of electron, and has a mass 2328 times that of an electron. Like the neutron, this particle decays spontaneously. Its lifetime (this term is understood in the same way as it was for a neutron) is 0.8×10^{-10} s.

Unlike the neutron, the hyperon may decay in two ways:

$$\text{either } \Sigma^+ \rightarrow p + \pi^0 \quad \text{or} \quad \Sigma^+ \rightarrow n + \pi^+.$$

(π^0 and π^+ are neutral and positively charged pions, respectively). Approximately 50 per cent of the hyperons decay in one way, and the others decay in the other way. We cannot unambiguously predict either when the hyperon decays or how.

The instability of atomic nuclei (radioactivity). Each element may have several types of atomic nuclei. They contain the same number of protons (the atomic number determining the position of the element in the periodic table), but the number of neutrons in them differs; these different nuclei are called *isotopes*. Most isotopes of an element are *unstable*. The unstable isotopes of an element transform spontaneously into isotopes of other elements simultaneously emitting particles. This phenomenon is called *radioactivity*. It was first discovered by the French physicist Antoine Henry Becquerel (1852-1908) in 1896. The term “radioactivity” was introduced by Pierre Curie (1859-1906) and Marie Skłodowska-Curie (1867-1934) who investigated the phenomenon and won the Nobel Prize for physics (with A.H. Becquerel) in 1903.

Investigations showed that the lifetime of unstable isotopes is essentially different for different isotopes and follow different decay routes (different types of radioactivity). The lifetime of an isotope may be measured in milliseconds, or it may be years or centuries. There are isotopes with lifetimes of over 10^8 years. The study of long-lived unstable isotopes in nature have allowed scientists to determine the age of rocks.

Let us discuss different types of radioactivity. Let us use Z to denote the number of protons in a nucleus (the atomic number of an element) and use A to denote the sum of the number of protons and neutrons in the nucleus (the mass number). One type of radioactivity is called *alpha decay*. During the process, the initial nucleus (Z, A) decays into an alpha-particle (a helium nucleus, which consists of two protons and two neutrons) and a nucleus with two less protons ($Z - 2$) and a mass number four units smaller ($A - 4$):

$$X(Z, A) \rightarrow \alpha(2, 4) + Y(Z - 2, A - 4).$$

Another type of radioactivity is *beta decay*. During this process, one of the neutrons in the initial atomic nucleus turns into a proton, an electron, and an antineutrino, like a free neutron does. The proton stays within the new nucleus while the electron and the antineutrino escape. The scheme of beta decay can be presented as:

$$X(Z, A) \rightarrow Y(Z + 1, A) + e^- + \bar{\nu}_e.$$

Proton radioactivity is also possible:

$$X(Z, A) \rightarrow p + Y(Z - 1, A - 1).$$

Let me draw your attention to the *spontaneous fission* of atomic nuclei. The initial nucleus disintegrates spontaneously into two “fragments” (two new nuclei), approximately equal in mass, and several free neutrinos are formed in the process.

A chain of consecutive spontaneous transformations is shown in Figure 5.2. The neptunium isotope ^{237}Np ($Z = 93$, $A = 237$) finally turns into the stable isotope of bismuth ^{209}Bi ($Z = 83$, $A = 209$). The chain consists of “links” corresponding to alpha decays (the blue arrows in the figure) and beta decays (the red arrows). The lifetime (in the probabilistic sense) is indicated by each arrow. These chains are called *radioactive families* (or series).

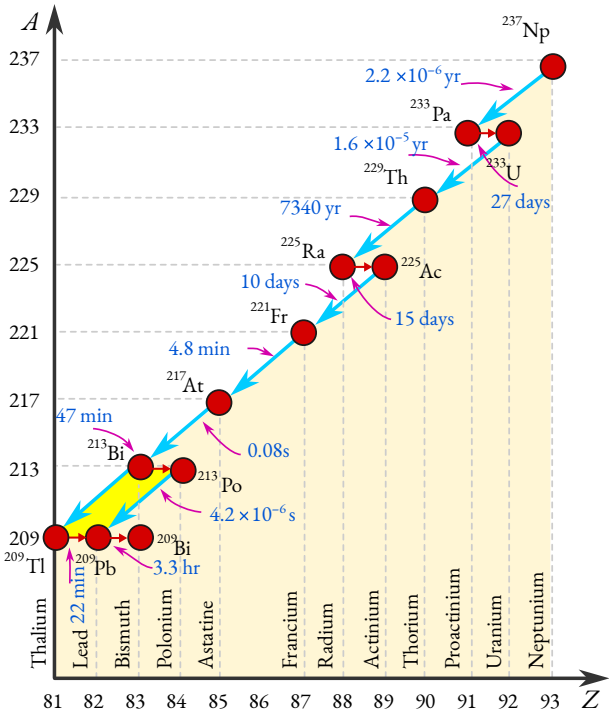


Figure 5.2: The number of neutrons decaying as a function of time.

Induced and spontaneous transitions in an atom. The reader will know that the energy of an atom can only have a set of discrete values that are specific to each atom. These allowed states of the atom are called *energy levels*. When we excite atoms by irradiating them, they jump from low energy levels to higher ones. The excited atoms

return to the lower levels by emitting light. These jumps are called *quantum transitions*.

A quantum transition may be either *induced* (stimulated) or *spontaneous*. Transitions due to the excitation of an atom are always induced. The reverse transitions may be both induced and spontaneous.

For simplicity's sake, let us only consider two atomic energy levels: energies E_1 and E_2 (Figure 5.3). The transition $E_1 \rightarrow E_2$ is an induced and occurs when an atom absorbs a photon with energy $\varepsilon_{12} = E_2 - E_1$. The atom may return to level E_1 either spontaneously or by being induced to. A photon with energy ε_{12} is emitted in the process. The spontaneous transition $E_2 \rightarrow E_1$ is a random event. The induced transition $E_2 \rightarrow E_1$ is caused when a photon passes near the atom. The energy of the photon should be equal to ε_{12} . The figure shows each of these three processes:

- (a) the absorption of a photon with energy ε_{12} by the atom (atom transition $E_1 \rightarrow E_2$),
- (b) the spontaneous emission of a photon with energy ε_{12} by the atom (atom transition $E_2 \rightarrow E_1$), and
- (c) the induced emission of a photon possessing energy ε_{12} by the atom while it interacts with the stimulating primary photon also possessing energy ε_{12} (atom transition $E_2 \rightarrow E_1$).

It should be noted that the photon emitted during an induced transition, as it were, copies every property of the primary photon that caused the atom transition. For instance, it moves in the same direction as the primary photon.

How does a laser generate radiation? Many books on science for the general reader cover lasers and explain the induced emission of photons as being due to simultaneous emission by a large number of specially selected atoms or molecules (they are called *active centres*). The photons resulting from induced radiation move in the same direction, thus forming laser radiation (laser is the abbreviation for light amplification by stimulated emission of radiation).

The explanation of how a laser generates is commonly given as follows. First, the active centres are excited, for instance, by an intense flash of light. It is necessary that the number of active centres in the higher energy level should be greater than those in the lower one. Then photons begin to appear with an energy equal to the difference between the energies of the higher and lower levels of the active centres, and the active centres radiate by induced emission more often than the reverse process (the process of photon absorption) occurs. This is easy to see if we take into account that each primary photon can cause with equal probability the transition of an active centre both upwards (the process of light absorption) and downwards (induced emission). Therefore, everything

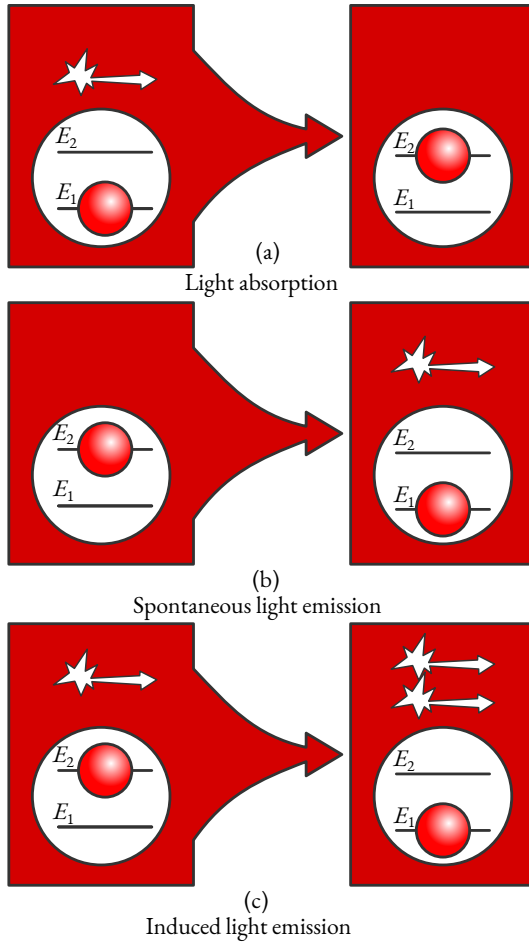


Figure 5.3: Induced and spontaneous emissions in a two level quantum system.

depends on whether the number of active centres is greater in the higher or in the lower level. If there are more centres in the higher level, more downward transitions will occur i.e. induced emission prevails. The result is an intense beam of laser photons.

Everything is correct in this explanation. However, most writers ignore the appearance of the primary photons which induce emission of the new photons and trigger

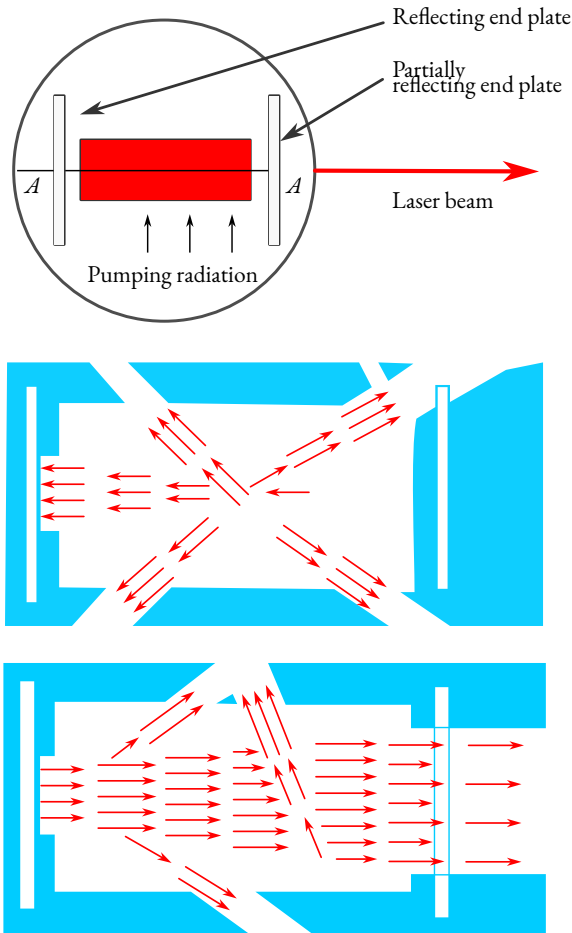


Figure 5.4: A schematic explanation of how does the laser emission works.

the process of laser generation. The photons appear due to the *spontaneous* transition of active centres from the higher level to the lower one. Because they are so important for lasers, we should not forget the *primacy* (and fundamentality) of the spontaneous emission processes. We could stop discussing lasers at this point. However, a reader might want to ask some questions.

READER: “You said that the induced photon copies every property of the primary photon, in particular, its direction of motion.”

AUTHOR: “Quite right.”

READER: “But spontaneous transitions yield photons moving in random directions. Therefore, the induced photons should also move in random directions. A photon that has appeared spontaneously, passing by a number of excited active centres, will induce an avalanche of photons in the direction it is moving in. The second spontaneous photon will cause an avalanche of induced photons in another direction, and so on. Now how come a laser beam has a single direction?”

AUTHOR: “You have made an essential point. Suppose AA is the beam direction (Figure 5.4). The active medium of a laser is formed into a cylinder with its long axis in the AA direction. Two mirrors (end plates) are placed at right angles to AA , one mirror being partially silvered: it lets the emission out. Some photons will be randomly born in the AA direction (or close enough to it), and then will pass the active substance along a relatively long path, which is increased because it might be reflected many times from the mirrors at both ends. By interacting with induced active centres, these photons, sooner or later, will cause a powerful flux of induced photons to appear, and these form the laser beam. Photons randomly born in other directions and their associated induced photons will only travel a short distance along the active substance and will very soon be ‘out of play’. This can be seen clearly in the figure.”

“Let me note that the mirrors which set the direction of the laser beam constitute the *resonator* of the laser.”

READER: “So the laser radiation appears from noise (spontaneous radiation) owing to the *selectivity* of amplification, i.e. because the amplification occurs mainly in a certain direction.”

AUTHOR: “Exactly. Once again we encounter the *selection of information from noise*. The ordered (coherent) laser radiation is, as it were, selected from noise by the mirrors (end plates) of the resonator. The *amplification* of selection occurs owing to induced emission: when the secondary photon copies the properties of the primary one.”

From Uncertainty Relations the Wave Function

As we discussed spontaneous micro-processes, we found that the random in the microcosm reveals itself even in the behaviour of an individual body. This brings us close to a discussion of the *primacy* and *fundamentality* of the notion of probability in quantum mechanics. We shall start with the *uncertainty principle* suggested in 1927 by the German physicist Werner Heisenberg (1901-1976).

Uncertainty relations. A micro-body moving according to the laws of quantum mechanics does not have, strictly speaking, a trajectory of motion. This is because a micro-body does not have both a momentum and a set of coordinates *simultaneously*. Suppose a micro-body has a certain x -component of its momentum. It turns out that the x -coordinate of the micro-body in this state does not have any certain value. The other extreme case corresponds to the state of a micro-body in which, vice versa, its x -coordinate has a certain value while the x -component of its momentum does not. There are an infinite number of intermediate cases when both the x -coordinate of the body and the x -component of its momentum are not certain, although they take values within certain intervals.

Suppose Δx is the interval within which the x -coordinate values lie; let us call Δx the *uncertainty* of the x -coordinate. Let us consider the uncertainty of the x -component of the momentum Δp_x in a similar way. Heisenberg showed that the uncertainties Δx and Δp_x are related as:

$$\Delta x \Delta p_x \approx \hbar, \quad (5.3)$$

where $\hbar = 1.05 \times 10^{-34}$ J s is Planck's constant. Similar relations can be written down for other components of the coordinates and the momentum of the microbody: $\Delta y \Delta p_y \approx \hbar$ and $\Delta z \Delta p_z \approx \hbar$.

These are Heisenberg's famous *uncertainty relations*. We shall limit ourselves to a discussion of the *coordinate-momentum* uncertainty relations. However, there are similar relations for some other pairs of variables, for instance, for *energy* and *time*, and *angle* and the *moment of momentum*. Heisenberg wrote that we cannot interpret the processes on the atomic scale as we might a large-scale process. However, if we use common notions, their applicability is limited by the uncertainty relations.

When we discuss the uncertainty relations, we shall only use (5.3). Do not, however, think that this relation outlaws accurate measurements of the momentum or coordinates of a micro-body. It only states that a micro-body cannot simultaneously have both accurately defined coordinates and an accurately defined momentum. For instance, if we try to measure the x -coordinate of a micro-body more accurately (in

other words, to decrease Δx), we will cause its momentum's x -component to become more uncertain. In the limit when the x -coordinate of the micro-body has a certain value (the micro-body is accurately localized), the uncertainty of the x -component of its momentum becomes very large. And vice versa, establishing the x -component of the micro-body's momentum more accurately unavoidably causes its x -coordinate to become more uncertain.

Let us consider a plane in which the x -coordinate of a body is plotted along one axis (the x -axis) and its momentum's x -component is plotted along the other axis (the p_x -axis) (Figure 5.5). If the body obeyed the laws of classical mechanics, its any state would be a point in the plane. However, the state of a micro-body corresponds to a rectangle with area \hbar . Other types of state are also possible. They correspond to rectangles of various shapes. Some of them are presented in the figure.

Uncertainty relations and the wave properties of a micro-body.

In 1924, the French physicist Louis de Broglie (b. 1892) hypothesized that a microbody possesses the properties of both a *particle* and a *wave*. Its particle characteristics (energy ε and momentum p), de Broglie postulated, are related to its wave characteristics (frequency ω and wavelength λ) thus:

$$\varepsilon = \hbar \omega \quad \text{and} \quad p = 2\pi \hbar / \lambda. \quad (5.4)$$

This hypothesis seemed absurd to many physicists. They could not understand what a particle's wavelength might be.

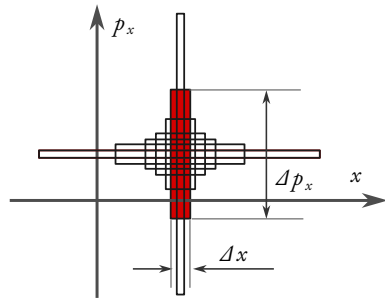


Figure 5.5: States of a micro-body.

In 1927, a striking result was obtained during experiments in which an electron beam was sent through thin metal plates. After leaving the plate the electrons spread out in a *diffraction* pattern (Figure 5.6). *Electron diffraction* by a crystalline lattice became an experimental fact, and yet diffraction and interference are wave properties. Therefore, the experiments on electron diffraction were unanimously accepted as proof of the wave properties of the electron. The nature of the electron waves remained as puzzling as before, but nobody doubted their existence.

We shall return to the waves below. Let us use de Broglie's hypothesis to explain the uncertainty relations. Suppose that a strictly parallel electron beam with a momentum p passes through a plate with a very narrow slit whose width in the x -direction is d (the x -axis is at right angles to the beam) (Figure 5.7).

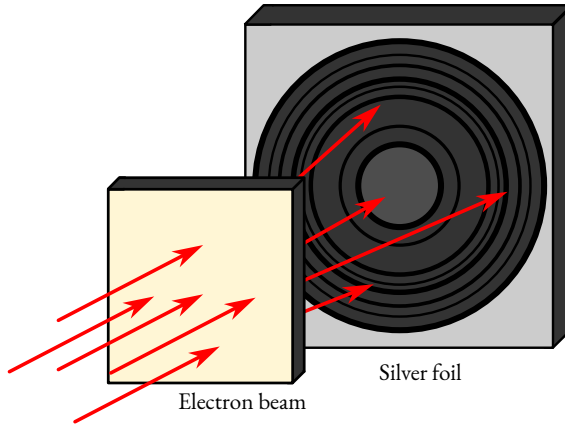


Figure 5.6: Electron diffraction experiment proved the wave properties of the electrons.

The electrons are diffracted when they pass through the slit. According to classical wave theory, the angle through which the electrons are diffracted to the first diffraction maximum is $\theta \approx \lambda/d$. If we use λ as the wave characteristic of the electron and use the second relation in (5.4), we can write θ as $\theta \approx \hbar/pd$. However, what does the angle θ mean in terms of particles? In fact what happens is that when the electron passes through the slit, it acquires a momentum Δp_x in the x -direction. Clearly, $\Delta p_x \approx p\theta$. Since $\theta \approx \hbar/pd$, we obtain $\Delta p_x d \approx \hbar$. If d is thought of as the uncertainty Δx of the x -coordinate while the electron passes through the slit, we obtain the uncertainty relation (5.3).

The wave function. Suppose a micro-body is in a state such that the x -component of its momentum has a value p_0 . We know that the value of the x -coordinate of the micro-body in this state is very uncertain. In other words, the micro-body may be found at any place on the x -axis.

Does this mean that we can say nothing about the x -coordinate of the micro-body? No, it does not. It turns out that we can establish the probability that the micro-body's x -coordinate takes a value from x to $x + \Delta x$. This probability can be written as $|\Psi_{p_0}(x)|^2 \Delta x$.

We see that the probability density needed to find the micro-body at a point x is the square of the function $\Psi_{p_0}(x)$. This function is commonly called the *wave function*. The reader should not understand the term “wave” literally. The point is that in the

1930s the researchers looking at the microcosm got so carried away by wave concepts (due to the experiments on electron diffraction) that they spoke of “wave mechanics” rather than “quantum mechanics”.

Thus, the state of a micro-body such that the x -component of its momentum possesses a value p_0 and given that the x -coordinate does not have any certain value is described by the wave function $\Psi_{p_0}(x)$ whose squared absolute value is the probability density of the micro-body to be found at point x . I want to emphasize that the results of measuring a micro-body’s coordinate in state $\Psi_{p_0}(x)$ prove to be *random* each time. A value of the coordinate is realized with the probability density $|\Psi_{p_0}(x)|^2$.

I have only selected one state of the micro-body without dealing with, for instance, the states where both the momentum and coordinate are uncertain. Besides, I limited the discussion to the coordinate and momentum without dealing with other variables, for instance, energy or the moment of momentum. I believe that this is sufficient to see the main point: any state of a micro-body is described by a function defining the probability (or the probability density) of some characteristics of the micro-body. Thereby it is clear that quantum mechanics of even one micro-body is a *probabilistic* theory.

The electron in the atom. The electrons in atoms may occur in different states. A change in the electron’s state may, for instance, be related to the atom’s transition from one energy level to another. Let us put down possible states of an electron in an atom by means of wave functions $\Psi_j(x, y, z)$, where j is a set of some numbers characterizing a state and (x, y, z) are coordinates of the electron. Given what we said above, we can conclude that $|\Psi_j(x, y, z)|^2$ is the density of probability that we can find an electron in state j at point (x, y, z) . Now imagine an “object” whose density is proportional to $|\Psi_j(x, y, z)|^2$ at various points of space. We can imagine a cloud with the density varying from point to point. The density inside the cloud is the greatest. While the point approaches the surface of the cloud, the density falls to zero, and thus the cloud has some *shape* (although without a distinct bounding surface).

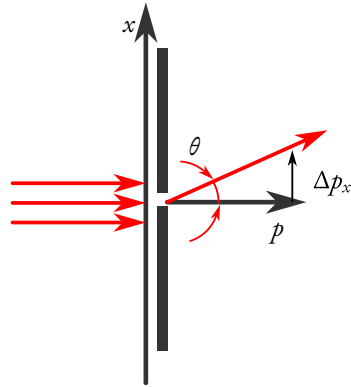


Figure 5.7: Electron diffraction experiment as understood by the uncertainty of the momentum and direction.

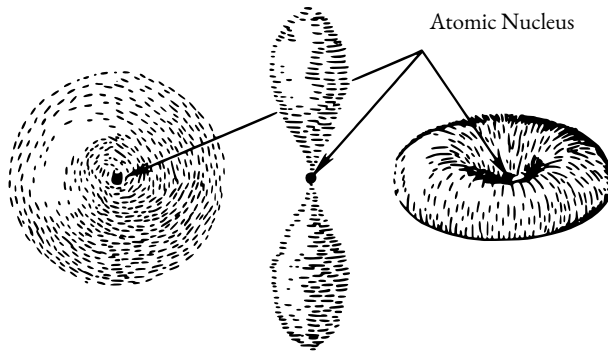


Figure 5.8: Electron clouds in an atom.

This “cloud” is the probabilistic “image” of an electron in an atom. Several “electron clouds” are shown in Figure 5.8 for the electron’s several states in an atom.

Interference and Summing Probability Amplitudes

After reading this section, we shall see that the probabilities in the microcosm obey the laws we have not dealt with above. It is noteworthy that these laws allow us to make a rather unexpected conclusion, namely that interference and diffraction are possible in principle even in the absence of waves. They may be an effect of *specific rules for the summation of probabilities*.

The puzzling behaviour of a microbody in an interferometer. Without discussing the technical details, let us consider an experiment in which particles pass through an interferometer containing two close slits and then are detected on a special screen (Figure 5.9).

Let us consider the x -coordinate of the particles. In order to deal with a *probability* rather than a *probability density*, suppose that the x -axis on the screen is separated into small identical intervals, so that when we speak of the probability that a particle arrives at a point x we mean the probability of arriving at the appropriate part of the axis around point x .

Suppose slit A is closed while slit B is open. After a large enough number of particles have been detected on the screen, we obtain a distribution defined by the function $w_B(x)$ (Figure 5.9 (a)). This function is the probability that a particle passing through slit B

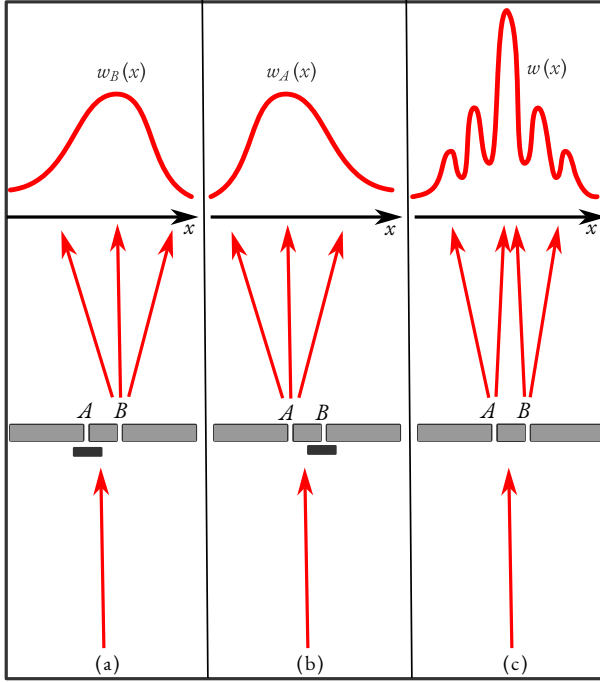


Figure 5.9: The two slit experiment.

(when slit A is closed) will arrive at point x . Given our remarks in the preceding section, we have

$$w_B(x) = |\Psi_B(x)|^2, \quad (5.5)$$

where $\Psi_B(x)$ is the wave function for the particle passing through slit B . I should remark that recently the term “wave function” is being more often substituted by a better term, “probability amplitude” (or “probability density amplitude”). Therefore, the probabilistic nature of the particle’s state is emphasized in this way. We shall now use the term *probability amplitude* and not wave function. Thus, $\Psi_B(x)$ is the probability amplitude that a particle will arrive at point x after passing through slit B (when slit A is closed).

Now suppose that slit B is closed while slit A is open. If this is the case, the screen (Fig. 5.9b) will show the distribution $w_A(x)$:

$$w_A(x) = |\Psi_A(x)|^2, \quad (5.6)$$

where $\Psi_A(x)$ is the probability amplitude of a particle arriving at point x after passing through slit A (when slit B is closed).

And finally, let us open both slits. It would be natural to believe that if it passes through one of the slits, a particle “does not feel” the other slit. It can be said that it is “indifferent” as to whether the other slit is open or closed. And in this case the distribution on the screen should be the sum of distributions (5.5) and (5.6), which, by the way, corresponds to the rule of *probability summation*:

$$w_{AB}(x) = w_A(x) + w_B(x) = |\Psi_A(x)|^2 + |\Psi_B(x)|^2. \quad (5.7)$$

In reality, the screen yields a typical *interference* distribution (Figure 5.9 (c)). rather than distribution (5.7). It turns out that when it passes through one slit the particle somehow “feels” the other slit. Or, perhaps more incomprehensible, the particle somehow manages to pass through both slits at the same time. How does it actually pass the interferometer?

“Spying” destroys the interference pattern. Let us try and “spy” on how the particle behaves when both slits are open. The “spying” would seem to be possible in principle. For instance, we might place a source of light near each slit and detect the photons reflected by the particles near each slit. Such experiments have in fact been carried out. They showed that the particle passes through only one slit, and at the same time it turned out that the distribution on the screen was described by (5.7). This means that “spying” helps establish the details of the particle’s passing through the interferometer, but the interference distribution is destroyed.

We have thus a curious situation. If the light is turned off (no “spying”), there is interference, but the mechanism by which the particle passes through the interferometer cannot be uncovered. If the light is on, the mechanism can be ascertained, but the interference distribution is destroyed.

When should we sum up probabilities and when probability amplitudes? Let me start to explain the amazing results described above. A particle has two options (two alternatives): to pass through either slit A or slit B . If the light is off, these alternatives are *indistinguishable*. They become *distinguishable* if the light is on, and therefore, “spying” or, in terms of science, observation is possible.

One of the basic conclusions of quantum mechanics is that

if alternatives are distinguishable, the respective probabilities are to be summed up; but if the alternatives are indistinguishable, probability amplitudes rather than probabilities are summed up.

Therefore, when the light is on, the probabilities should be summed up, but when the light is off, the probability amplitudes should be summed up. In the former case, we arrive at distribution (5.7), and in the latter case, we obtain the distribution

$$w_x(x) = |\Psi_A(x) + \Psi_B(x)|^2. \quad (5.8)$$

This is an interference distribution. It can be shown that

$$|\Psi_A + \Psi_B|^2 = |\Psi_A|^2 + |\Psi_B|^2 + \left[\frac{\Psi_A}{\Psi_B} |\Psi_B|^2 + \frac{\Psi_B}{\Psi_A} |\Psi_A|^2 \right]. \quad (5.9)$$

The expression in the square brackets is “responsible” for the interference nature of the distribution $w(x)$. In classical physics, the problem of distinguishable (indistinguishable) events does not exist since classical events are always distinguishable. In the microcosm, the situation is qualitatively different. Here we encounter the possibility of complete *indistinguishability* of some random events. This possibility arises because of the fundamental *identity* of all particles of the same type. An electron is like any other to a far greater extent than the proverbial two peas in a pod. Naturally, electrons may be in different states, which allows us to distinguish between them. However, any electron (as a physical particle) is indistinguishable from any other electron. Here we are dealing with absolute identity. In the last analysis, it allows for indistinguishable alternatives.

We see that interference should not be limited to wave concepts. The interference in microphenomena is not necessarily related to waves, it may be a consequence of probabilistic laws, or more accurately, a consequence of the fact that we should sum up probability amplitudes rather than probabilities for indistinguishable events.

Quantum-mechanical superposition. Consider

$$\Psi_A(x) + \Psi_B(x) = \Psi(x). \quad (5.10)$$

The function $\Psi(x)$ in quantum mechanics is on an equal footing with functions $\Psi_A(x)$ and $\Psi_B(x)$ and like them it defines a state, or rather the probability amplitude for a random event. In this case, $\Psi_A(x)$ is the amplitude of the probability that a particle arrives at point x after passing through the interferometer with two open slits. This amplitude is said to be the *superposition* of the amplitudes $\Psi_A(x)$ and $\Psi_B(x)$.

It is impossible to imagine such a superposition in a demonstrative way. Otherwise, we should quite seriously have to believe that the particle passes simultaneously through both slits (A and B). Any attempt to reveal the details of this event *destroys the superposition*. It is destroyed each time either in favour of Ψ_A (the particle passed through slit A) or in favour of Ψ_B (the particle passed through slit B). Here we encounter one more

manifestation of the random. We have noted above that the arrival of the particle at a point on the screen is a random event, and probabilities (5.7) and (5.8) characterize these random events. It turns out that the “selection” of a slit by a particle is also random. The particle passes through slit A with a probability proportional to $|\Psi_A|^2$ and passes through slit B with a probability proportional to $|\Psi_B|^2$.

A wave or the sum of probability amplitudes? The wave concept explains the appearance of interference patterns best. However, the wave concept cannot explain the other phenomenon, the destruction of the interference pattern by “spying”. In other words, a wave can explain the *appearance* of quantum-mechanical superposition, but it cannot explain the *destruction* of the superposition in the process of observation.

Once convinced of this and the futility of the attempts to make “de Broglie’s waves” material, physicists admitted that these “waves” have nothing in common with really existing waves. This gave rise to a very expressive term of *probability waves*. Gradually, the term “wave mechanics” has been substituted everywhere by the term “quantum mechanics” while the term “wave function” has become more often replaced by the term “probability amplitude”.

Therefore, we should explain both the interference and diffraction of particles in terms of the necessity of summing up probability amplitudes instead of probabilities rather than in terms of enigmatic waves when the considered alternatives are indistinguishable. The probabilistic approach completely explains both the appearance and destruction of quantum-mechanical superposition.

In conclusion, let us consider a situation which illustrates the limited nature of the wave approach. We shall discuss the diffusion of very slow neutrons passing through a crystal.

Diffusion of neutrons in a crystal. A beam of neutrons with energies of only 0.1 eV is passed through a crystal. The neutrons diffused by the crystal’s nuclei are registered by a system of detectors (counters) along the x -axis (Figure 5.10). The crystal contains N nuclei, therefore, there are N alternatives. Each alternative corresponds to the diffusion of a neutron by a nucleus. Let us use $\Psi_j(x)$ to denote the probability amplitude that a neutron will arrive at the detector at point x after diffusing past the j -th nucleus.

It is interesting that the diffusion of a neutron by a nucleus may occur in two ways. In one case the neutron’s *spin is inverted* while there is no such inversion in the other case. Let me explain. A neutron can be represented as a rotating top. The top may rotate in either one direction or the other, the neutron’s spin being said to be either upwards or downwards, respectively. The crystal’s nuclei are also reminiscent of rotating tops,

i.e. they each have spin directions. When a neutron (top) collides with a nucleus, it may or may not change the direction of its rotation. In the former case, the neutron's spin remains unchanged while in the latter it is reversed. If a diffused neutron changes the direction of its rotation, the direction of rotation of the nucleus at which the act of diffusion occurred should somehow change as well. Therefore, if diffusion occurs with one neutron's spin inversion, we are dealing with a *distinguishable* alternative. We can state that diffusion occurred precisely at the nucleus which changed the direction of its rotation. If diffusion occurs without spin inversion, it is in principle impossible to indicate which nucleus diffused the neutron; here we deal with an *indistinguishable* alternative.

Suppose φ is the probability amplitude that a neutron will diffuse with spin inversion while χ is the probability amplitude without inversion. Let us use $\Phi(x)$ to denote the probability amplitude that a neutron with inverted spin will arrive at point x , and $X(x)$ the same for a neutron with noninverted spin. The distribution of diffused neutrons detected by the counters can be presented as:

$$w_x(x) = |\varphi|^2 |\Phi(x)|^2 + |\chi|^2 |X(x)|^2. \quad (5.11)$$

Naturally, the alternatives corresponding to different types of neutron diffusion are distinguishable; therefore, probability $w(x)$ consists of two terms (two probabilities are summed up). In turn, each term is the product of two probabilities.

Now let us express $|\Phi(x)|^2$ and $|X(x)|^2$ in terms of amplitudes $\Psi_j(x)$. If a neutron is diffused with spin inversion, the alternatives are distinguishable; therefore, the *probabilities are summed up*, and hence

$$|\Phi(x)|^2 = \sum_{j=1}^N |\Psi_j(x)|^2. \quad (5.12)$$

If the spin of a diffused neutron has not been inverted, the alternatives are indistinguishable; therefore the probability amplitudes are to be summed up (amplitude superposi-

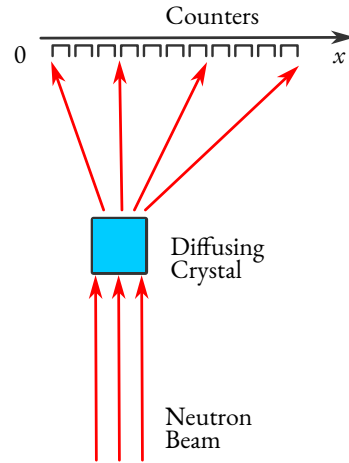


Figure 5.10: Diffraction of neutrons through a crystal.

tion occurs) and hence

$$|X(x)|^2 = \left| \sum_{j=1}^N |\Psi_j(x)|^2 \right|. \quad (5.13)$$

Substituting (5.12) and (5.13) into (5.11), we obtain:

$$w(x) = \left[|\phi|^2 \sum_{j=1}^N |\Psi_j(x)|^2 \right] + \left[|\chi|^2 \sum_{j=1}^N |\Psi_j(x)|^2 \right]. \quad (5.14)$$

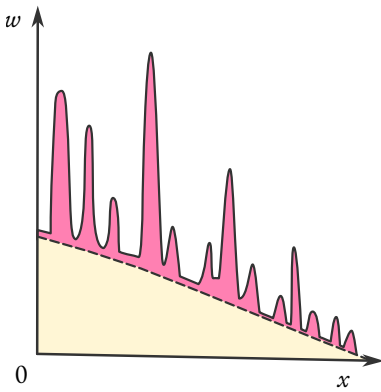


Figure 5.11: Distribution of diffraction of neutrons through a crystal.

The distribution of diffused neutrons $w(x)$ in experiment is shown in Figure 5.11. It consists of a smoothly varying “background” and a set of interference maxima. The “background” is defined in (5.14)) by the term in the first square brackets while the interference maxima give the term in the second square brackets.

Using wave concepts, we have to assume that a neutron has the wave properties while diffusing without spin inversion (the interference pattern appears). The same neutron does not show any wave properties in diffusion with spin inversion (the interference pattern does not appear). It is evident that this assumption is quite unnatural.

Probability and Causality

READER: “I think there is too much randomness in the microcosm. A neutron suddenly turns into three new particles at random, without any external influence. An atom may be at rest for many years and then suddenly, for no apparent reason, decays and turns into an atom of another chemical element. An electron randomly passes through a slit in the interferometer and quite as randomly arrives at a point on the screen. Doesn’t it mean that, in fact, there is no *causality* in the phenomena of the microcosm?”

AUTHOR: “No, it doesn’t. The phenomena of the microcosm show very explicitly the *dialectical unity of the random and the necessary*. Neutrons

decay in a random manner, but their quantity varies in time according to a certain law. An electron randomly arrives at a point on the screen, but the distribution of arrivals of many electrons is necessary. There are no grounds for doubting existence of causality in the microcosm. We should bear in mind that causality in the microcosm reveals itself unlike that in the macrocosm. In quantum mechanics, potential possibilities to realize events or, in other words, the *probabilities* of these events are only causally related, rather than individual realized events themselves. The probability amplitude (wave function) obeys a definite equation of motion. Knowing the probability amplitude at the initial moment and using this equation (it is called *Schrödinger's equation*), we can find the probability amplitude at an arbitrary moment in time."

READER: "It is not clear why a neutron should suddenly decay. Maybe, the particles in question are, in fact, more complex systems whose physical nature is not yet known?"

AUTHOR: "We touched on this in our first talk. I said that the search for *hidden parameters*, which would explain why, for instance, a neutron decays, eventually, at a given moment in time proved to be unsuccessful. But I would like to show what is behind the posed question. Asking it, you proceed from that probability in the microcosm is *not objective* but related with our lack of knowing *some details*. I think that both the examples from the microcosm and many of the examples from our macrocosm we cited convinced you that probability can be both subjective (related to a lack of knowledge) and objective. This is essential. It is only when probability is objective that we can say that probabilistic regularities are primary, or fundamental."

READER: "Please explain this idea."

AUTHOR: "If probability were reduced to a lack of information, it could be reduced in principle to dynamic relations supposing unambiguous prediction. This would mean that the probabilistic laws would conceal the dynamic ones. In this case it could be possible to assume that, in the last analysis, everything is strictly interrelated in the Nature."

READER: "But doesn't any event, any phenomenon have a cause in the long run?"

AUTHOR: "You're right to mention causality. However, why do you believe that the existence of objective probability means the absence of causality?"

READER: "Objective probability suggests objective randomness. And this randomness reveals itself without any cause, because it is related to chance."

AUTHOR: "I throw a die, and, say, the four comes up. You throw a die, and the three comes up. Are these events objectively random or not? What do you think?"

READER: "Each event has definite causes. The occurrence of an event depends, over a long stretch, on the position of the die in your hand, the wave of hand, the push, the air resistance, the distance from the hand to the floor, etc."

AUTHOR: "Right. And nonetheless, the events are not objectively random ones. Throwing a die, you are not interested in the way I threw mine. We are not interested in how a die is thrown at all, do not try to control and direct our actions. Therefore, the occurrence of the four on my die and the three on yours are objectively random events. The occurrence of the three is not related to the occurrence of the four just before it."

READER: "I don't quite understand."

AUTHOR: "I can give you another example. Suppose the events are telephoned taxi orders. Each order conceals a chain of causes. However, the arriving orders are objectively random events for the taxi-depot dispatcher. And this is not because he does not know the chain of causes but because of an objective circumstance, namely the lack of connection between the actions of the people making orders for taxi. The events are considered, as it were, in two different planes. In one, they are objectively random, while in the other each of them has definite causes. As you see, objective probability agrees with causality."

READER: "Your example is from practice. And what about microphenomena? Let us once again take the example with neutron decay. Suppose this event is objectively random in a 'plane'. But in what plane should we look for the causes for the neutron decay?"

AUTHOR: "Neutron decay is indeed objectively random. We cannot control the lifetime of a given neutron in principle because of deep reasons and not a lack of knowledge about some details. There is no internal "clock" in a neutron. As was noted above, neutrons "do not get old". This can be seen in that a neutron may live for some time irrespective of how long it has already lived by the moment we start counting time. Because it is

objectively random, neutron decay is not a causeless event. I want to note that when we speak of the *spontaneous* behaviour of a particle, we are being inaccurate. Strictly speaking, only a hundred per cent *isolated* particle can behave *spontaneously*. And here we come close to a fundamental point which we haven't discussed yet."

"The point is that a particle is *not isolated*, it interacts with the world around it. It is in essence dependent on the conditions of each concrete situation. The term 'interaction' should be understood here in a wider meaning than it is understood when considering usual (force) interactions."

READER: "New puzzles of quantum mechanics."

AUTHOR: "I do not mean any puzzles. At a certain level of investigation of physical phenomena, isolation is lost in principle. For instance, the distinct boundary between the field and the matter is erased. The mutual transformations of particles become apparent. The idea of the *unity of the world and the universal interrelation of the phenomena in it* acquires a special meaning on the level of the microcosm."

READER: "How can we imagine in a demonstrative way that a decaying neutron is not isolated?"

AUTHOR: "A *vacuum* in quantum mechanics is not a void but a space in which particles are randomly born and annihilated. The neutron interacts with them."

Chapter 6

Probability in Biology

Mutations that bring about changes seem to be a random phenomenon at first glance, but they have regularity in the long run.

N.I. Vavilov

The genetic code as it is passed from generation to generation changes randomly due to many causes and without any definite direction, and these changes only randomly turn to be fit to survive.

B.M. Mednikov

Introduction

Jean Baptiste Lamarck (1744-1829). In 1809, the French scientist Jean Baptiste Lamarck published *Philosophy of Zoology*. It was the first attempt to produce a theory of evolution for all species, but it was unsuccessful. In his work on the theory, Lamarck started from two erroneous axioms. Firstly, he believed that the tendency to improvement is inherent in all living beings. He saw here the drive for evolution. Naturally, there is no mysterious inner drive which makes all species evolve and advance. Secondly, Lamarck believed that the environment can directly induce changes in the shape of living being's organs. For instance, there was a time when giraffes with short necks existed. For some reason, their habitat changed and their food rose high above the ground (the leaves of high trees). In order to reach the food, giraffes had to extend their necks. This occurred from generation to generation. As a result of long-term exercise, the necks of giraffes became much longer.

One of Lamarck's proofs was the generally known fact that a physically weak person could become an athlete by being regularly in sport. He formulated the following law:

"In each animal that has not yet completed its development, more frequent and prolonged exercise of some organ reinforces the organ, develops it, increases, and gives it more strength, in proportion to the duration of its usage; while a constant lack of exercise gradually weakens any organ, brings its decline, continuously decreases its ability, and finally, makes it disappear."

Lamarck was utterly wrong. It is known that trained muscles, like other acquired abilities, cannot be inherited. Using modern terminology, we can say that Lamarck did not understand the difference between phenotype and genotype. The *genotype* is the genetic constitution of an organism, usually in respect to one or more genes responsible for a particular ability. Parents transfer a set of hereditary elements to their progeny. The *phenotype* is the entire physical, biochemical, and physiological make-up of an individual as determined both genetically and environmentally, the set of internal and external features of the organism. The phenotype varies during the organism's life as it interacts with the environment. Regular physical exercise, persistent learning, a correct organization of labour and rest help everyone improve their own phenotype. However, this does not influence the genotype.

Charles Darwin (1809-1882). The correct theory of evolution of the species was developed by the English scientist Charles Darwin, and his theory became known as Darwinism. Darwin presented the theory in *The Origin of Species by Means of Natural Selection, or the Preservation of Favoured Races in the Struggle for Life*, which was published in 1859.

Darwin emphasized three factors: *variability, inheritance, and natural selection*. The environment, which influences an organism, may bring about random changes in its genotype. These changes can be inherited and gradually accumulated in the progeny. The nature of the changes varies. Some of them are randomly more favourable from the viewpoint of the organism's adaption to the environment while others are less favourable or even bad. When the progeny accumulate these random changes, natural selection reveals itself. The organisms that are least fit produce less offspring, die prematurely, and are forced out by the more fit individuals in the long run.

In describing Darwin's theory, I emphasize the role of the random on purpose. The reader may recognize the familiar idea of the *selection of information from noise*.

In his consideration of the evolution of species, Lamarck in fact only *recognized necessity*. Once the environment changes, the organism would necessarily change by exercising or not exercising the relevant organs. Lamarck's "evolution" would only

necessitate a complication in the organism's organization if each species had an inner drive to advance.

Darwin considered evolution from the positions of the dialectical unity of the necessary and the random. The indifferent Nature causes random hereditary changes in the organism. Then, by natural selection, it mercilessly throws off those which randomly prove to be less fit and keeps those which randomly prove to be adapted to the environment. The result is that the evolution of a species occurs by necessity. The development proceeds through the selection of the fittest, the Nature being indifferent as to whether the organism becomes more or less complicated. The possibilities for adaptation are diverse. The result is the diversity of the plant and animal species we observe. Earth is thought to accommodate about 1.5 million animal species and about 0.5 million plant species.

Darwin's theory has become universally recognized. However, there was a "soft spot" in it, which was pointed out in 1867 by Fleming Jenkins, a teacher from Edinburgh. Jenkins noted that Darwin's theory is not clear about the mechanism by which the changes in the progeny accumulated. At first, changes in a trait only occur in a limited number of individuals. These individuals crossbreed with normal ones. The result, as Jenkins asserted, should be *dissipation* of the changed trait in the progeny and not its *accumulation*. The trait should dilute out and gradually eliminate ($1/2$ of the change in the first generation, $1/4$ of the change in the second generation, $1/8$ in the third, $1/16$ in the fourth, etc.)

Darwin contemplated Jenkins's objection for the remaining fifteen years of his life. He could not find a solution.

However, a solution was already found in 1865 by Gregor Johann Mendel, a teacher in the monastery school in Brunn (now Brno, Czechoslovakia). Alas, Darwin did not know about Mendel's investigations.

Gregor Johann Mendel (1822-1884) Mendel started his famous experiments on peas three years before the publication of *The Origin of Species*. When Darwin's book appeared, he read it thoroughly and was very interested in Darwin's work. Mendel is said to have remarked with respect to Darwin's theory: "It is not yet complete. Something is missing." Mendel's investigation was directed to mending the "flaw" in Darwin's theory. Mendel was a plant breeder and he wanted to follow the change in the genotype over successive generations of a crossing. He picked the pea as the subject of investigation.

Mendel took two varieties of pea, one with yellow seeds and one with green seeds. By crossing the two varieties, he found that the first generation only had yellow seeds. The green pea trait had vanished. Then Mendel crossed the first generation with itself and

grew a second generation. This time individuals with green seeds appeared, although there were noticeably fewer of them than there were individuals with yellow seeds. Mendel counted the number of both and took the ratio, i.e.

$$x : y = 6022 : 2001 = 3.01 : 1.$$

Mendel carried out six other experiments simultaneously. In each experiment, he used two varieties of pea each with a different trait. For instance, in one of his experiments, he crossed a pea variety with smooth seeds with one with wrinkled seeds. He found only smooth-seed individuals in the first generation. Individuals with wrinkled seeds appeared in the second generation. The ratio of the number of individuals with smooth seeds to the number of individuals with wrinkled seeds was

$$x : y = 5474 : 1850 = 2.96 : 1.$$

In the other five experiments, Mendel crossed varieties which differed in skin colour or seed shape or colouration when immature or the location of flowers or the size of the individuals (dwarfs and giants).

	In each experiment, the first generation consisted of individuals with one of the two opposite parental traits. Mendel called this trait the <i>dominant</i> one, and the other trait, which disappeared for a generation, he called the <i>recessive</i> one. Yellow seeds was a dominant trait, while the green-seed trait was recessive in the first of the experiments we mentioned. In the second experiment, the smooth-seed trait was dominant, and the wrinkled-seed was recessive. We gave the ratio $x : y$, i.e. the ratio of the number of individuals with the dominant trait to the number of individuals with the recessive one in the second generation for the two of Mendel's experiments. Mendel obtained the following ratios from the other five experiments as shown in the Table 6.1.
$x : y = 705 : 224 = 3.15 : 1,$	
$x : y = 882 : 299 = 2.95 : 1,$	
$x : y = 428 : 152 = 2.82 : 1,$	
$x : y = 651 : 207 = 3.14 : 1,$	
$x : y = 787 : 277 = 2.84 : 1.$	

In each case, the $x : y$ ratio is close to $3 : 1$. So Mendel could maintain that when individuals with opposite traits are crossed, *one trait is suppressed by the other* and not diluted out (as Jenkins believed). Thus Mendel asserted the existence of dominant and recessive traits such that individuals in the first generation only have the dominant trait, while the recessive one is completely suppressed (the *law of uniformity of first generation individuals*). When the first generation is crossed with one another, individuals bearing

both the dominant and recessive traits appear in the second generation, their ratio being approximately 3 : 1.

However, Mendel did not stop there. He crossed the second generation with itself and obtained individuals in the third and then in the fourth generation. Mendel discovered that second-generation individuals with the recessive trait did not produce different progeny in either the third or fourth generation. About one third of the second-generation individuals with the dominant trait behaved in the same way. Two thirds of the second-generation individuals with the dominant trait produced different third-generation progeny, the ratio being 3 : 1 again. Third-generation individuals with the recessive trait and one third of the individuals with the dominant trait did not produce different progeny in the fourth generation, while the other individuals in the third generation did produce different progeny, the ratio of individuals with each trait being 3 : 1 again.

Note that the production of different progeny demonstrates an essential point: individuals with identical external features may possess different hereditary trait, which is revealed in the external features of their progeny. We see that one cannot use the phenotype to make generalizations about the genotype. If an individual does not produce different progeny, then it is called *homozygotic*, otherwise being termed *heterozygotic*. All the individuals with the recessive trait in the second generation are homozygotic.

Mendel's results can be seen in Figure 6.1 where the yellow circles are individuals with the dominant trait, while the green circles are individuals with the recessive trait. We see a definite pattern. Mendel discovered this pattern and therefore discovered the mechanism by which hereditary traits are passed down from generation to generation. Mendel understood that the pattern had a probabilistic nature.

The pattern of crossings had been observed before Mendel. Suffice it, for instance, to cite the diary of Mendel's contemporary, a gardener at the Paris Botanical Gardens:

"Starting from the second generation, the outward appearance changes noticeably. The perfect uniformity of the first generation individuals is usually replaced by an extreme diversity of progeny, some of them being close to the species type of the father and the other close to that of the mother ..."

But nobody before Mendel had attempted to investigate the change in separate traits, or count the number of individuals with different traits in consecutive generations. Mendel was the first person to do this, spending eight years on his experiments. Therefore, unlike his predecessors, Mendel came to understand the *pattern behind the hereditary transmission of traits*.

It is good to pause here, to discuss the laws governing crossbreeding which Mendel

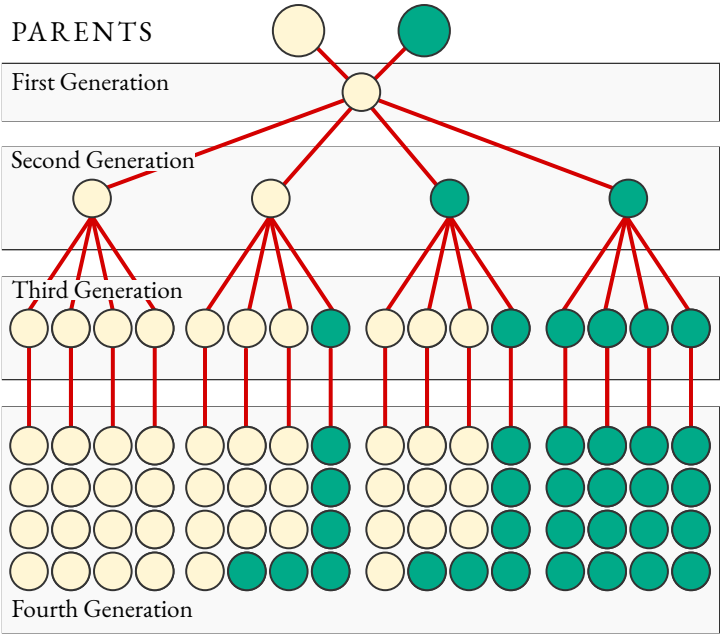


Figure 6.1: Results of Mendel's experiment with yellow and green peas till fourth generation.

discovered. We shall do this in the next section from the viewpoint of modern genetics. Let me only tell the reader that Mendel presented his results first in February 1865 to the Society of Natural Scientists in Brünn. The audience did not understand the exceptional importance of the presentation, nor could they guess that it would cause a revolution in the study of heredity. In 1866, Mendel's paper was published in the Brünn Bulletin and was sent to some 120 listed scientific institutions in many countries. Unfortunately, Darwin did not receive a copy.

The world now recognizes Mendel as the founder of modern genetics. However, the recognition only came in 1900, fifteen years after his passing.

The Patterns After the Random Combination of Genes in Crossbreeding

Chromosomes and genes. Perhaps you can recall some data on *cytology*, the branch of biology dealing with the structure, behaviour, growth, and reproduction of cells, and the functions and chemistry of the cell components. There are two types of cell: *germ* cells (*gametes*) and *somatic* cells. The nucleus of each cell contains threadlike structures, *chromosomes*, which carry linearly arranged genetic units in gigantic molecules of deoxyribonucleic acid (DNA) or combination with protein molecules. The chromosomes, or, to be more accurate, the DNA molecules are the carriers of genetic information, which is encoded in the sequence of bases, defining the genotype of the organism. The separate parts of a chromosome, responsible for a hereditary trait, are the basic units of heredity, or *genes*. Each chromosome contains several hundred genes. Sometimes, a chromosome is viewed as a thread with beads for the genes.

Each species has a fixed *set of chromosomes*. For instance, oats possess 42 chromosomes, *Drosophila* possess 8 chromosomes, chimpanzees possess 48 chromosomes, and human beings have 46 chromosomes. The nucleus of every somatic cell contains all the chromosomes needed for the individual of that species. This means that *each cell* in the organism contains all the individual's *genetic information*.

The numbers of chromosomes we gave for several species characterize the chromosomes in the somatic cell, rather than in germ cells. Each germ cell (gamete) has half the number of chromosomes than a somatic cell.

Let us start with the chromosome set of a somatic cell. This set includes two *sex chromosomes*. Female individuals have two identical sex chromosomes (two *X-chromosomes*) while male individuals have two different sex chromosomes (one *X-chromosome* and one *Y-chromosome*). The somatic chromosomes in a somatic cell come in pairs; the chromosomes in each pair (they are called *homologous*) are very much like each other. Each contains the same number of genes at the same loci on both chromosome threads, and the main point is that they are responsible for the same kind of trait. For instance, the pea has a pair of homologous chromosomes each of which contains a gene for seed colour. This gene, like any other gene, has two forms (they are called *alleles*), dominant and recessive. The dominant form of the colour gene (the *dominant allele*) corresponds to yellow while the recessive one (the *recessive allele*) corresponds to green. If the genes on both homologous chromosomes contain the same allele, the individual is *homozygotic* with respect to the trait in question. If a chromosome contains an allele which is different from the one contained in the homologous chromosome, the individual is *heterozygotic*. Its phenotype shows the trait

corresponding to the dominant allele.

Now let us consider the chromosome set of a gamete (a germ cell). A gamete has only one sex chromosome. It is always an *X-chromosome* for a female individual. A male individual may contain either an *X-chromosome* (in some gametes) or a *Y-chromosome* (in the other gametes). Besides the single sex chromosome, a gamete contains one chromosome from each pair of homologous chromosomes.

Suppose there are only two pairs of homologous chromosomes, and a certain trait corresponds to each pair. Moreover, assume the given individual is heterozygotic with respect to both traits. This individual will have four types of gamete, which can be seen in Figure 6.2 (a) (the red colour in the figure is for the chromosomes with the dominant alleles and the blue colour for the recessive alleles). The individual in Figure 6.2 (b) is homozygotic with respect to one trait and heterozygotic with respect to the other. There are only two types of gamete in this case.

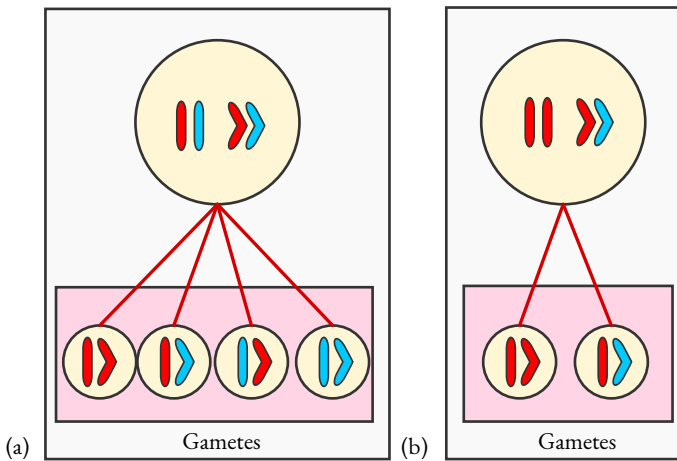


Figure 6.2: Combination of chromosomes and their results.

During fertilization, a female gamete fuses with a male gamete. The fertilized egg (called a *zygote*) has a complete chromosome set. Each pair of homologous chromosomes receives one chromosome from the father and one from the mother. The organism develops from a zygote through a series of divisions. The division of the cell is preceded by the *replication* of all the chromosomes contained in the nucleus of the cell. The result is that the nucleus of each somatic cell of the organism contains the same set of chromosomes and genes that the zygote had. When the organism reaches sexual

maturity, a special process occurs leading to the production of gametes. We shall discuss this process below.

The law of segregation. Let us consider one particular trait, for instance the colour of pea seeds, as in one of Mendel's experiments. Let us consider the results of this experiment from the point of view of modern cytology.

All the individuals in the first generation are heterozygotic for the trait. Each somatic cell contains both alleles for seed colour: yellow (dominant allele) and green (recessive allele). Naturally, every seed belonging to these individuals is yellow. Each first-generation individual has two types of gamete: some with the dominant allele (A -gametes) and the others with the recessive allele (a -gametes). It is clear that there must be both female and male A -gametes and a -gametes.

Now let us consider the second generation. Each new organism develops from a zygote which is formed when a male gamete (A or a) fuses with a female gamete (A or a). Clearly, four alternatives are possible (Figure 6.3):

*AA or a male A -gamete fuses with a female A -gamete,
 Aa or a male A -gamete fuses with a female a -gamete,
 aA or a male a -gamete fuses with a female A -gamete, and
 aa or a male a -gamete fuses with a female a -gamete.*

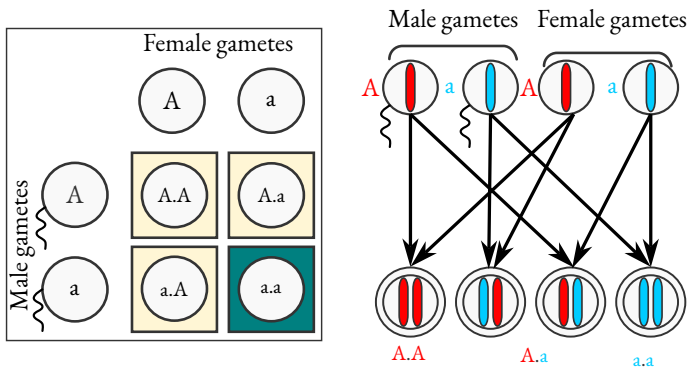


Figure 6.3: Analysing Mendel's experiment using modern cytology.

All these alternatives are equally probable. Therefore, if we take a large enough number of zygotes, a quarter of them will be composed of AA -zygotes, a quarter will contain aa -zygotes, and finally, a half will contain Aa -zygotes (the variants Aa and aA

are equal from the viewpoint of trait heredity). If a zygote contains at least one dominant allele, the phenotype will reveal the dominant feature (yellow seeds). Therefore, individuals (plants) developing from AA - or Aa -zygotes will have yellow seeds while individuals developing from aa -zygotes will have green seeds. We see, therefore, that the probability that *an individual will have a dominant trait is $3/4$ while the probability that an individual will have the recessive trait is $1/4$* . Hence the ratio 3 : 1 Mendel obtained, which quantitatively characterizes the segregation of a trait in the transition from the first generation of the crossing to the second. Mendel both found this ratio and correctly explained it using the notion of *probability*. This was *Mendel's first law*, which is also known as the law of segregation.

I want to emphasize: a zygote is formed as the result of the *random union* of male and female gametes. A large number of such random unions will *necessarily* lead to a definite pattern, which is expressed in the Mendel's first law.

Note that AA - and aa -zygotes produce homozygotic individuals with respect to the trait while Aa -zygotes produce heterozygotic individuals, and in the next generation the heterozygotic individuals will produce a 3 : 1 split of traits again.

The law of independent assortment of genes. Suppose we look at the second generation of a crossing involving two traits at the same time. Let us assume (this is essential) that the genes responsible for the traits are on different pairs of homologous chromosomes. An example of this combination is the colour of pea seeds and the shape of the seeds. Let us use A to denote the dominant allele of colour (yellow), a to denote the recessive allele of colour (green), B to denote the dominant allele of shape (smooth seeds), and b to denote the recessive allele of shape (wrinkled seeds).

Each first-generation individual has four types of male and four types of female gamete: AB , Ab , aB , and ab (recall Figure 6.2 (a)). A zygote is formed when two gametes (male and female) of any of the four types fuse. There are 16 possible alternatives; they are presented in Figure 6.4. *Each alternative is equally probable*. Therefore, the ratio of the number of zygotes of different types (with respect to the total number of zygotes, which should be large) is:

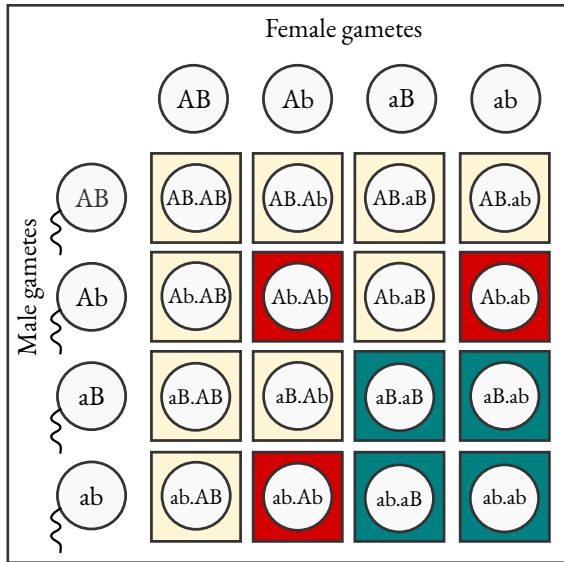


Figure 6.4: Analysing Mendel's experiment using modern cytology.

$1/16$ for zygotes $AB \cdot AB$,
 $1/16$ for $Ab \cdot Ab$,
 $1/16$ for $aB \cdot aB$,
 $1/16$ for $ab \cdot ab$,
 $1/8$ for $AB \cdot Ab$ (including $Ab \cdot AB$),
 $1/8$ for $AB \cdot aB$ (including $aB \cdot AB$),
 $1/8$ for $AB \cdot ab$ (including $ab \cdot AB$),
 $1/8$ for $Ab \cdot aB$ (including $aB \cdot Ab$),
 $1/8$ for $Ab \cdot ab$ (including $ab \cdot Ab$), and
 $1/8$ for $aB \cdot ab$ (including $ab \cdot aB$).

Regarding the suppression of recessive alleles by the corresponding dominant alleles, we can conclude that the probability that an individual will have yellow smooth seeds in the second generation equals the sum of probabilities for the zygotes $AB \cdot AB$, $AB \cdot Ab$, $AB \cdot aB$, $Ab \cdot ab$, and $Ab \cdot ab$, i.e. $1/16 + 1/8 + 1/8 + 1/8 + 1/8 = 9/16$. The probability that an individual will have yellow wrinkled seeds equals the sum of probabilities of the formation of zygotes $Ab \cdot Ab$ and $Ab \cdot ab$, i.e. $1/16 + 1/8 = 3/16$. The probability that an individual will have green smooth seeds equals the sum of probabilities of the formation of zygotes $aB \cdot aB$ and $aB \cdot ab$, i.e. $1/16 + 1/8 = 3/16$.

And finally, the probability that an individual will have green wrinkled seeds equals the probability of the formation of the zygote $ab \cdot ab$, i. e. $1/16$. Therefore, the numbers of different phenotypes (with these traits) in the second generation are in the ratio $9 : 3 : 3 : 1$. This is the essence of *Mendel's second law*, according to which the segregation by one trait is *independent* from the segregation by another trait.

Morgan's Law. The law of the independent assortment of genes is valid when the genes are on different chromosomes in a gamete (and on different pairs of homologous chromosomes in a somatic cell). If the genes belong to the *same* chromosome, they will be inherited together. This is the explanation for deviations from Mendel's second law. The deviation was discovered and investigated by the American biologist Morgan and is observed whenever traits are defined by linked genes, i.e. the genes are on the same chromosome. The joint inheritance of linked genes became known as *Morgan's law*.

Thomas Hunt Morgan (1866-1945) was the founder of the chromosome theory of inheritance. By introducing the idea of a chromosome, he substantiated Mendel's laws and pointed out under which conditions they are applicable. Besides, he obtained a number of new results. These results include Morgan's law and the phenomenon of chromosome crossing over, which he discovered.

Chromosome crossing over. In an investigation of the inheritance of traits defined by linked genes, Morgan discovered that the linkage *is not absolute*: some of the second-generation individuals inherit some of the linked genes from one parent and the rest from the other. Carrying out his investigations on *Drosophila*, Morgan could explain this fact. He showed that the formation of germ cells in an organism (this process is called *meiosis*) starts with a "farewell dance" of homologous chromosomes.

Imagine two elongated homologous chromosome threads, which, before they leave each other and join different gametes, tightly embrace each other (each gene in contact with the corresponding gene) and then wind around each other several times. This winding of the chromosomes (crossing over) results in the intracellular forces which arise to pull the chromosomes apart, *break* them. The site where the break occurs varies randomly from one pair of crossed-over chromosomes to another. The result is that one gamete receives *complementing* parts of both homologous chromosomes rather than an intact chromosome, and the other parts of these chromosomes are received by the other gamete. The process is illustrated in Figure 6.5. Let me emphasize that corresponding genes on both chromosomes (I mean the alleles) are in contact with each other at the moment of break. Therefore, wherever the break might be, an allele from one chromosome gets into one gamete while an allele from the other chromosome gets into the other gamete. In other words, either gamete gets an allele with the considered gene. This can be thought of as "dancing" pairs of chromosomes exchanging equivalent

parts of themselves before leaving each other. All the same, each gamete has a complete set of genes characterizing the given chromosome. And there is a *random combination* of paternal and maternal alleles.

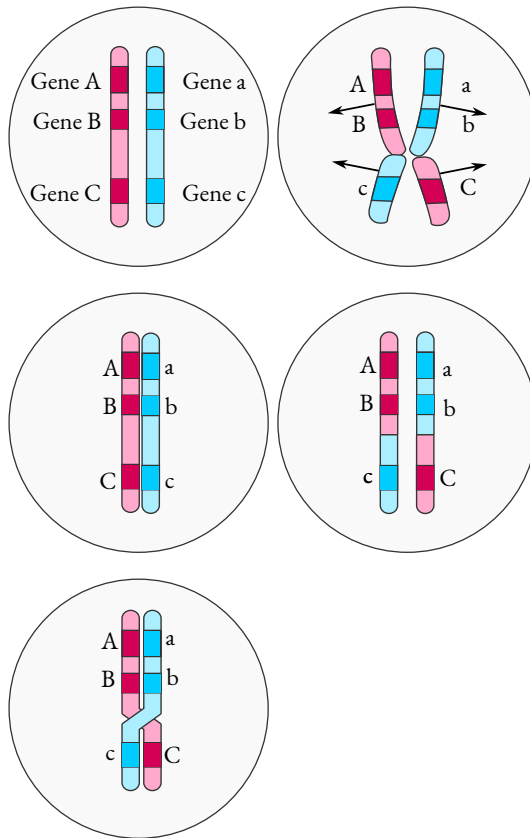


Figure 6.5: “Farewell dance” of the genes combines genes from the two parents.

Chance plays an essential role in the phenomenon of chromosome crossing over. The site of the break is *random* in a pair of chromosomes, and therefore, the combination of parental alleles is *random*.

By expanding the domain of the random, the phenomenon of chromosome crossing over enhances intra-species development, creating additional possibilities for “shuffling” the parental genes. At the same time, the phenomenon, as it were, protects the species

from random genetic “infringements”. Suppose individuals from two different species cross at random and hybrids appear. Each “homologous pair” in the hybrids unites chromosomes that are very unlike in their gene structure (because the chromosomes come from parents of different species). When the time comes to produce the germ cells, these chromosomes are unable to carry out the “farewell dance” because of fundamental differences. They consequently are unable to form gametes, and therefore, no second-generation hybrids appear. This is why mules (the hybrid offspring of a male ass and a female horse) do not have any progeny.

A boy or a girl? I have already noted that the sex chromosomes of a female are both the same: they are X -chromosomes. By contrast, the sex chromosomes of a male are different, each male having one X -chromosome and one Y -chromosome. Half of all male gametes carry one X -chromosome and the rest carry one Y -chromosome. If a female gamete joins a male X -gamete, an XX -zygote is produced, and a female offspring develops from it. But if a female gamete fuses with a male Y -gamete, an XY -zygote is produced, and a male offspring develops from it. This is the answer to the question: a boy or a girl?

Mutations

We have considered random changes in the genetic code that might occur when a combination of parental genes is crossed over. All these changes are limited by the available gene pool. New genes cannot be created in the process. However, random inheritable changes do occur which are not related to the combination of genes. They are caused by the action of the environment on the genetic structure of the chromosomes and random disorders in the biological mechanism that maintains the genetic information during meiosis and the division of the somatic cells. These genetic changes are called *mutations*.

The appearance of mutations. There is a serious human disease in which a sufferer’s blood is unable to clot. This disease is called *hemophilia*. It is inherited and occurs in men only. It has been found out that hemophilia is the consequence of a *mutation* in a gene that is located on the X -chromosome. Since women have two X -chromosomes, the mutated gene, which is recessive, on one chromosome is matched by a normal gene on the other, which suppresses the illness. This is why women do not suffer from hemophilia. This is not the case in men. The set of sex chromosomes in men consists of two *different* chromosomes: one X -chromosome and one Y -chromosome. There is no normal paired gene which can suppress the hemophilia gene. Consequently

a man receiving an *X*-chromosome with the mutated gene from a phenotypically healthy mother suffers from hemophilia.

Fortunately, mutations are mostly harmless. A short-fingered hand, a sixth finger, and the heart on the right are relatively rare mutations. More frequent mutations show themselves as, for instance, different eye colours, baldness (including the shape of the bald spot), and unusual hair colour in animals. Mutations often occur in plants and appear in a great variety of ways, such as changes in the shape of the stem, leaves, and flowers.

The causes of mutations. A mutation is a rather rare event. For instance, the probability that a gamete with an *X*-chromosome taken at random will contain the mutation related to hemophilia is only one in 10^5 . Other mutations occur even less often, with the probability of about one in 10^6 on the average. However, we should take into account the diversity of mutations. They can be associated with very different genes of which there is an enormous number in each gamete. We should also take into account that mutations are inherited and thus *accumulate*. The result is that mutations per se are not too rare events. It has been calculated that one in ten human gametes carries a mutation.

The appearance of each mutation is a *random* event. However, the event results from objective causes. An organism develops from a zygote due to the cell divisions. The process of cell division begins with replication of chromosomes, and therefore, DNA molecules in the cell nucleus. Each DNA molecule recreates an exact copy of itself with the same set of genes. The complicated process of replication of a DNA molecule sometimes occurs with random deviations. We know that genetic information is recorded in DNA very economically on the molecular level. When the data is copied, various kinds of “misprint” are possible due to the thermal movement of molecules. The “misprints” appear due to the unavoidable *fluctuations* in the behaviour of matter. For instance, when a DNA molecule replicates, there might be a random increase in the number of hydrogen ions in the vicinity of some nitrogen base. This fluctuation may cause the detachment of the base from the DNA, i.e. to a disturbance in the structure of the gene.

In every sexually reproducing species, the progeny only receive the mutations in the germ cells. Therefore, the random disordering that occurs in the formation of the germ cells, in meiosis, is essential. These disorders may cover both separate genes and chromosomes as a whole. Individual gametes may receive a chromosome with a distorted gene structure or not receive a chromosome at all. The formation of gametes with extra chromosomes is also possible.

The thermal movement of matter molecules is not the only cause of mutation. Special investigations have revealed a number of *external* factors which cause mutations and are called *mutagenic* factors. Certain chemicals and various kinds of radiation, e. g. X-rays, neutron beams, fast charged particles, are all mutagenic.

Advantages and disadvantages of mutations. From the viewpoint of evolution, mutations are certainly advantageous. Moreover, they are necessary. The vast diversity of genes in each species and the diversity of species existing on the Earth are a consequence of mutations having occurred over many millions of years, and they still occur. From the point of view of an individual, as a rule, mutations are harmful and even lethal more often than not. Being the result of long-term evolution, each organism is a complex genotype and adapted to its habitat. A random change in the genotype would more likely disrupt its smoothly running biological mechanism.

Therefore, we see that mutations are at the same time both useful (even necessary) and harmful. If mutations occur too frequently in a given species (for instance, because its habitat is radioactively contaminated), this will increase the mortality rate and, as a consequence, cause the decline or possibly the extinction of the species. By contrast, if mutations occur too rarely in a given species, it may not be able to adapt and may also become extinct should its habitat change considerably. For instance, the dinosaurs could not adapt to a cooling in the climate and became extinct. Thus, it is disadvantageous for there to be too many mutations or for them to be too frequent. It is also disadvantageous for there to be practically no mutations or for them to occur too rarely.

The Organism and mutations. The adaptation of an organism to its habitat also supposes the adaptation to mutations, owing to which the degree of harm brought about by mutations can be essentially reduced. This adaptation is natural because the development of species is directly related to its survivability.

Let us discuss this problem from the positions of genetics. Suppose a zygote appears when a normal and a mutated gamete combine. We shall call a gamete mutated if one of its chromosomes has a faulty (mutated) gene. Suppose this gene is responsible for a vital process, and so we are dealing with a dangerous mutation. The mutated gene is opposed by the normal gene in the paired chromosome. Now mutated gene may either be dominant or recessive with respect to the normal gene, and we shall consider both possibilities.

If the mutated gene is *dominant*, it immediately starts its “harmful activity”, and the organism may die as an embryo. Darwinian selection here carries out its sanitary mission long before the dominant mutation can propagate to future progeny. The result is that there is no accumulation of dominant mutated genes. This is not so if

the mutated gene is *recessive*. It is suppressed by the normal gene, and therefore, the organism will be phenotypically healthy. Moreover, there will be healthy organism phenotypes in the progeny. It is only in rare cases that the recessive mutated gene reveals itself, i.e. when a descendant gets the gene simultaneously through the paternal and maternal gametes.

I would very much like to say that the Nature has taken care to decrease the danger of harmful mutations. However, recall that the Nature never takes care of anything or anybody. The principle is the selection of the fittest. There is no “wisdom” in the Nature.

Unfortunately, people sometimes increase the danger of mutations. The probability that two recessive genes will combine in a descendant increases if close relatives marry or a *small group* of people, for instance, a small religious sect, small community, or the population of a hamlet in the mountains, intermarry. Wherever this practice is common, various types of genetic disease are unavoidable (they are called *recessive diseases*). There are about five hundred such diseases known so far. They may bring about idiocy, debility, deaf-mutism, constitutional inferiority, etc. Therefore, any artificial separation or division of people into closed groups increases the genetic danger and leads to a higher probability of recessive disease.

In the second half of this century, the mutation danger drastically increased due to nuclear weapon testing. Radioactivity is very mutagenic. Therefore, it is impossible to overestimate the importance of the international treaty banning the testing of nuclear weapons in the atmosphere, space, and underwater, which was concluded at the initiative of the Soviet Union. In 1963, the treaty was signed by the USSR, USA, and Great Britain. Over a hundred countries have signed it so far.

The law of homologous series in hereditary variability. Each individual mutation is a random, undirected, and unpredictable event. If a given species sustains relatively many mutations (this is seen in plants), the picture of mutations on the whole shows some *regularity*, or necessity. This is substantiated by the *law of homologous series* in mutations discovered by the Soviet biologist Nikolai I. Vavilov (1887-1943). Generalizing a great deal of data, Vavilov concluded that genetically close species should be characterized by similar (homologous) series of hereditary variability. For instance, if mutations cause a number of rather frequently occurring hereditary traits in rye, a similar series of traits should also be observed in wheat, barley, oats, etc.

Vavilov’s law is sometimes compared to Mendeleev’s periodic table, thus emphasizing that like the periodic table it can be used to predict new members, or mutants. In 1917, during a scientific expedition in the Pamir, Vavilov found a variety of wheat

with leaves without a ligule, a small growth at the base. At the time, biologists were not aware of rye or barley varieties without ligules. However, Vavilov's law required that they exist, and in 1918 a variety of rye was found without ligules, while in 1935, a barley variety without ligules was obtained after irradiating common barley with X-rays.

Evolution Through the Eyes of Geneticists

There was a time when some biologists tried to oppose the theories of Darwin and Mendel. This should be regarded as a frustrating mistake and seems absurd today. It is generally recognized that genetics have put Darwin's theory of the origin and evolution of species on a sound scientific basis, and explained the heritability of changed traits. Darwinism is a logical and authoritative science capable of giving valuable practical recommendations. Modern genetics is deeply rooted in Darwinism.

Undirected hereditary variability. The Soviet biologist Ivan Shmalgausen (1884-1963) once said that each species and each of its populations contain a "pool of hereditary variability". This pool can be utilized by natural selection in a changed habitat.

There are two basic "mechanisms" for the appearance of undirected hereditary variability. Firstly, there is *mutation variability*. Mutations underlie the diversity of species and the diversity of genes within a species. Mutation changes occur very slowly, but they occur continuously and have done so since the time immemorial. The "mechanism" by which hereditary variability appears as the result of the random crossing of *parental* genes is faster. Here we should distinguish between the combination of genes as the result of fusing random pairs of gametes and the combination of genes as the result of "shuffled" parts of paired chromosomes getting randomly into a gamete (the phenomenon of chromosome crossing over).

Naturally, the changes in the combination of genes are limited by the volume of the gene pool. However, the pool is enormous. It has been calculated that the gene pools of a father and a mother make it possible in principle to construct up to 10^{50} different human genotypes. This is a rather hard number to imagine. Less than 10^{10} people live on the Earth. Therefore, there is practically no chance that two individuals will be genetically identical (unless, of course, they are twins developing from the same zygote). Each person is *genetically* unique; a person possesses a genotype which is unlike any other genotype.

Darwin's demon versus Maxwell's demon. We discussed the Maxwell's demon in Chapter 4. Without getting outside information, the demon could not in principle select

faster molecules and direct them into the other half of the vessel. This hapless demon demonstrated the fundamental *impossibility of selection at the atomic or molecular level*, as was demanded by the second law of thermodynamics.

In a discussion on natural selection in the *Nature*, the American biochemist and science-fiction writer Isaac Asimov (1920-1992) used the term “Darwin’s demon”. Unlike Maxwell’s hapless demon, Darwin’s demon operates very successfully, selecting organisms with a better chance for survival and letting them reproduce and move into the next generation. The major distinction between the Darwin’s and Maxwell’s demons is that they operate on *different* levels. Anything begins at the atomic or molecular level. Random, undirected mutation and the random combinations of genes occur at this level. If Maxwell’s demon could operate, he would start by selecting the most “advantageous” mutations and the most “successful” combinations of genes. This does not occur because selection is impossible at atomic or molecular level.

And here is where the *principle of reinforcement* starts. Suppose that a mutated gene has got into a zygote. While the organism develops, the cells divide, and the result is that the mutated gene is replicated about 10^{15} times. The combination of genes in the zygote has also been replicated. Therefore, *random changes* in the genetic code in the process of the development of the phenotype becomes *reinforced*. And this is a transition from the atomic or molecular level to the level of *macrophenomena*. Selection at this level is possible. I want to emphasize: Darwin’s demon does not try to select different genetic codes, and in this sense it is not quite like Maxwell’s demon. It influences the organism’s phenotypes, where any change in the genetic code is amplified about 10^{15} times.

There should be no need to explain how Darwin’s demon operates. The way natural selection is realized is described in every textbook on biology. Let me only note that the “demon” is rather merciless. It operates severely: it eliminates phenotypes which have randomly proved unfit. Taking those which are randomly less or more fit to the habitat, it gives preference to the more fit while the less fit are, as a rule, eliminated.

However, Darwin’s demon does not operate directly and gives the less fit a chance to survive. Changes in the genetic code which may not be used today may be utilized tomorrow. They are useless and even harmful today, but they may become useful later. It means that we should not hurry and render the verdict. Let the random variation in the genetic code “sleep”, stay dormant for a while, for several generations of phenotypes, masked as a recessive gene. It may suddenly be helpful later.

Naturally, the effect of Darwin’s demon or, in other words, natural selection does not oppose the second law of thermodynamics in any way. We noted above, that living beings only exist due to the inflow of negentropy from the environment, i.e. due to the

rise of entropy in this environment. This increase in entropy is the “fee” for the service provided by Darwin’s demon.

Diversity of species. The diversity of species on the Earth, where *Protozoa* coexist with very complicated and organized species, is the result of evolution proceeding for about two thousand million years. Two thousand million years ago the Earth was only inhabited by bacteria and blue-green algae. Several hundred million years later, unicellular organisms with a cellular nucleus appeared. After a period of several hundred million years more, *Coelenterata*, worms, and molluscs appeared. About five hundred million years ago, fish appeared, followed by amphibia, and still later by reptiles. Mammals appeared about a hundred million years ago. Note that there is no mere transition from less complicated species to more complicated ones in this evolutionary process. Naturally, many species became extinct; nevertheless, today a tremendous number of simple species exist alongside complicated ones. Evolution has been directed *from the less fit to the more fit* rather than *from the simple to the complicated* because natural selection operates in this direction and no other one. The characteristic feature of this process is the *increase in the number of species* and their growing diversity. Higher species will appear, which is an advance for the evolution process.

We could give a number of reasons why evolution increases the number of species. Firstly, hereditary variability increases in time, i.e. mutations accumulate and the gene pool extends. Secondly, there are a great number of ways to adapt to any given change in the environment. Natural selection approves of any acceptable versions. The selected variants may have either a more or less complicated organization. Thirdly, once it has appeared, a species has a certain stability. In particular, it resists the danger of being incorporated by other species. Recall that hybrids produced by crossing between different species cannot form germ cells, and therefore, cannot have any progeny. Naturally, when we consider the increase in the number of species, we have to take into account the reverse processes, such as the elimination of a species due to an interspecific struggle or the extinction of a species because of its inability to adapt to sudden severe changes in the environment.

Unpredictability of new species. We considered fluctuations in an ensemble of gas molecules in Chapter 4 and saw how the fluctuations of the variables for an individual molecule are great. They are comparable to the means of the variables. On the contrary, fluctuations of the variables for a macrosystem are extremely small. Therefore, a macro system could be described on the basis of dynamic laws rather than probabilistic laws. This is done in thermodynamics. This means that the transition from the atomic or molecular level of consideration to the macrolevel brings about, as it were, a reciprocal compensation of numerous random deviations in the behaviour of individual molecules.

The result is that the behaviour of the macrosystem as a whole becomes unpredictable unambiguously.

As to Nature, we encounter a qualitatively different situation. The individual fluctuations characterizing random changes in the genetic code are reinforced 10^{15} times and can be revealed on the macro level, in the organism phenotype. There is no reciprocal compensation here. *Each fluctuation grows to macroscopic dimensions.* Therefore, we can assert that the process of evolution in the Nature is *fundamentally unpredictable* in the sense that no one can foresee the emergence of concrete species. In other words, each species proves to be a random phenomenon. It can be eliminated, a new species can be created, but an extinct species cannot be restored. Each existing species is unique in this sense.

Conclusion We have discussed a number of problems in biology related to genetics and evolution theory. These problems clearly show the *fundamentality of probabilistic laws* and the *fundamental role of chance*. However, the topic of probability in biology is much wider. It also includes a number of problems that could not be treated in this book, such as the origin of life on the Earth, the change in the sizes of populations of species, the simulation of the nervous system, and the creation of a model of the human brain.

A Concluding Conversation

It is only when we finish writing that we find what we should have begun from.

Blaise Pascal

AUTHOR: "This book on the world of probabilities has come to an end. I hope that it gave some food for thought."

READER: "I have to admit that some points do not fit in with my own views. For instance, it is hard for me to see how randomness can be used to solve problems. I mean the perceptron, the Monte Carlo method, and the principle of homeostat. These are very much like 'miracles'."

AUTHOR: "In the meantime, they are just as 'miraculous' as the random number table."

READER: "I do not understand."

AUTHOR: "Each new digit in the table is independent of its predecessors. In spite of that, the table as a whole has stability. The digits appear independently from each other, but the frequency in which any digit appears is determinate."

"Besides, it is useless to try and write down a set of random digits 'by hand'. For instance, you might write 8, 2, 3, 2, 4, 5, 8, 7 ... And naturally, you see that perhaps you should write a 1 or a 6 because the digits are not in the sequence. And against your will, you correct your actions as a result of your preceding ones. The result is that you won't have a table of truly random numbers."

"It is essential to see that the occurrence of each random event is in no

way related to the preceding ones. Therefore, the stability observed in the picture of a large number of random events seems to be ‘miraculous’. In the long run, the ‘miracle’ is responsible for the properties of the perceptron or the Monte Carlo method.”

READER: “I can agree that the ‘root of the evil’ hides, in the long run, in a random number table. How can you explain the puzzling properties of this table?”

AUTHOR: “The explanation is in the word ‘symmetry’.”

READER: “Please explain.”

AUTHOR: “Having found a digit to add to your table, you take care to provide *symmetry* with respect to the occurrence of all the other digits. In other words, any digits from 0 to 9 should have the *same chance* of appearing.”

READER: “Suppose I have a bag and draw out balls labelled with different digits. What kind of symmetry do you mean here?”

AUTHOR: “For instance, the symmetry with respect to the exchange of the balls. Imagine that all the balls suddenly change places. If the symmetry exists, you will not notice the exchange. But this is not all. Once you return the balls to the bag and mix them, you restore the initial situation and take care to make the system symmetrical with respect to each act in which a ball is drawn. As you can see, the explanation is deep enough. Symmetry and asymmetry are related to the most fundamental notions. These notions underlie the scientific picture of the universe.”

READER: “I have read your book *This Amazingly Symmetrical World*. I was really amazed how far symmetry penetrates into every phenomenon occurring in this world. Now I see that the same can be said about randomness.”

AUTHOR: “Thank you. You refer to my book *This Amazingly Symmetrical World*, in which I attempted to set forth the notion of *symmetry* and show how the concepts of symmetry and asymmetry underlie our physical picture of the world.”

“In fact, the point of that book was not just symmetry but the *dialectical unity of symmetry and asymmetry*. Here I was not just considering randomness but the *dialectical unity of necessity and randomness*, which is, by the way, expressed in terms of probability.”

READER: "Judging from the remarks above, there seems to be a relation between necessity-randomness and symmetry-asymmetry."

AUTHOR: "Yes, and a very profound one. The principles of symmetry-asymmetry control both the laws of Nature and the laws of human creativity. And the role of probabilistic principles is no less fundamental."

READER: "I'd like to discuss the relation between symmetry and probability in more detail."

AUTHOR: "The classical definition of probability is underlain by the idea of equally possible outcomes. In turn, *equally possible outcomes always have a certain symmetry*. We dealt with equally possible outcomes when we discussed throwing a die or tossing a coin. Recall the definition of the statistical weight of a macro-state in terms of the number of equally possible micro=states (Chapter 4), and recall our discussion of equally possible alternatives while considering Mendel's laws (Chapter 6). In each case, the probability of an event was defined as being proportional to the number of equally possible (I can now say, symmetrical) outcomes, in each of which the given event is realized. In other words, the probability of an event is the sum of the probabilities of the respective equally possible outcomes."

READER: "I begin to think that the very rule of the *summation of probabilities* is based on a certain symmetry."

AUTHOR: "An interesting idea."

READER: "Given we are looking for the probability that one of two events will occur, it is irrelevant which one does because either of them brings about a result. The symmetry here is related to the independence with which the result is obtained with respect to the substitution of one event for the other."

AUTHOR: "We can go further. Suppose there is a deeper symmetry related to the *indistinguishability* between the first and the second event (similar situations were discussed in Chapter 5). The rule of the *summation of probabilities* is replaced in this case by the rule of the *summation of the probability amplitudes*."

READER: "True, I can clearly see here the relation between symmetry and probability."

AUTHOR: "This relation can be represented even more clearly if we use the

notion of *information*. Of course, you remember that information is underlain by probability in principle (see Chapter 3). Now the relation between information and symmetry is as follows: *less information corresponds to a more symmetrical state.*”

READER: “Then it is possible to believe that an increase in the symmetry of a state should result in a rise in its entropy.”

AUTHOR: “Exactly. Have a look at Figure 4.12. The state with the greatest statistical weight, and therefore, with the greatest entropy is the state corresponding to the uniform distribution of molecules in both halves of the vessel. Evidently, this is the most symmetrical arrangement (there is a mirror symmetry with respect to the plane separating the vessel in two).”

READER: “That is something here to think over. It means that human creativity reduces symmetry. However, symmetry is widely used in art. Is this not a contradiction?”

AUTHOR: “No. We use symmetry-asymmetry rather than only symmetry in art. We have already discussed it elsewhere, in my book on symmetry. Of course, these problems require special consideration. We can only touch on the problems here and not go into any detail.”

“I emphasized in my book on symmetry that symmetry operates to *limit the number of possible variants of structure or variants of behaviour*. Obviously, necessity operates in the same direction. On the other hand, asymmetry operates to increase the number of possible variants. Chance acts in the same direction. I have repeatedly drawn your attention to the fact that chance *creates new possibilities* and gives rise to new alternatives.”

READER: “This means that we can speak of the ‘composition of forces’ as follows. There are symmetry and necessity on the one side, and asymmetry and chance are on the other side.”

AUTHOR: “Yes, this ‘composition of forces’ is correct. Please recall the parable about the ‘Buridan’s ass’. I started with it my first conversation ‘between the author and the reader’ in *This Amazingly Symmetrical World.*”

READER: “I know this parable. The legend has it that a philosopher named Buridan left his ass between two heaps of food. The ass starved to death because he could not decide which heap to start with.”

AUTHOR: “The parable was an illustration of mirror symmetry. There were

two identical heaps of food and the ass at the same distance between them. The ass was unable to make his choice.”

READER: “As I see it, the ass starved to death because of symmetry.”

AUTHOR: “As the parable has it, he did. In reality, however, the ass lived in the ‘symmetrical world built on probability’ rather than in the ‘symmetrical world’ without any randomness. Any chance occurrence (a fly could bother the ass, he could jerk or move a little) might easily *destroy the symmetry*: one of the heaps could become a bit closer, and the problem of choice is ‘null and void’. As physicists say, a *spontaneous violation of symmetry* could easily occur.”

READER: “Is it possible to conclude that symmetry is harmful while chance is beneficial?”

AUTHOR: “I’m sure you realise that such a question is too far reaching. We have seen that symmetry decreases the number of versions of behaviour and reduces a number of alternatives. It is logical to admit that this reduction may lead to a hopeless situation, to a blind alley. And then chance becomes essential. On the other hand; too many chances, an abundance of alternatives and disorder may also be harmful. And then, order comes to rescue, i.e. symmetry and necessity.”

READER: “The danger of randomness is understandable. But what might be the danger of symmetry? If of course we exclude the situation the ‘Buridan’s ass’ was in.”

AUTHOR: “Firstly, the ‘Buridan’s ass’ was not an illustration from the life of animals but rather the presentation of a problem. Secondly, it is quite easy to give a practical example of the *danger of symmetry*. Designers of bridges, towers, and skyscrapers know that they must not be too symmetrical because of the danger of resonance oscillation, which can destroy a construction. There are well-known accidents when bridges have been destroyed due to resonance caused, for example, by a company of marching soldiers, rhythmic bursts of wind, or other seemingly inoffensive causes. Therefore, when large constructions are built, the symmetry is always violated in some way by randomly placed asymmetric beams, panels, etc.”

READER: “True, symmetry may be dangerous. As far as I understand, it is quite easy to destroy symmetry, be it a fly bothering an animal or an extra beam in a construction.”

AUTHOR: "Your attention has been drawn to an essential point. The *instability of symmetry* makes it easily upset and, in particular, allows for the possibility of spontaneous violation."

READER: "Symmetry is unstable. This is something new to me."

AUTHOR: "The investigation of unstable symmetry has not been going long, only a decade. It has led to the appearance of a new scientific discipline called *catastrophe theory*. This theory studies the relationship between symmetry and chance from the point of view of the development of various processes and phenomena."

READER: "The very name of the theory is somewhat dismal."

AUTHOR: "The catastrophes considered in the theory occur on different levels. Suppose a particle causes a violent process in a Geiger-Müller counter. The result is that the particle is registered. The process is a catastrophe on the scale of the microcosm. An enormous bridge or a jet plane may be suddenly brought down due to resonance oscillations. This is a catastrophe on our common scale. Catastrophes occur in a diversity of situations: sudden crystallization in a supercooled liquid, a landslide, the start of laser generation, etc. In each case, the system has an unstable symmetry, which may be upset by a random factor. These random factors may be very slight in influence, but they destroy the symmetry and therefore *trigger violent processes in an unstable system*, and these processes are called catastrophes."

READER: "Catastrophe theory appears to show up the deep relationship between symmetry-asymmetry and necessity-randomness quite clearly."

AUTHOR: "I quite agree with you. However, it is a theme for another book."

Recommended Literature

Chapter 1

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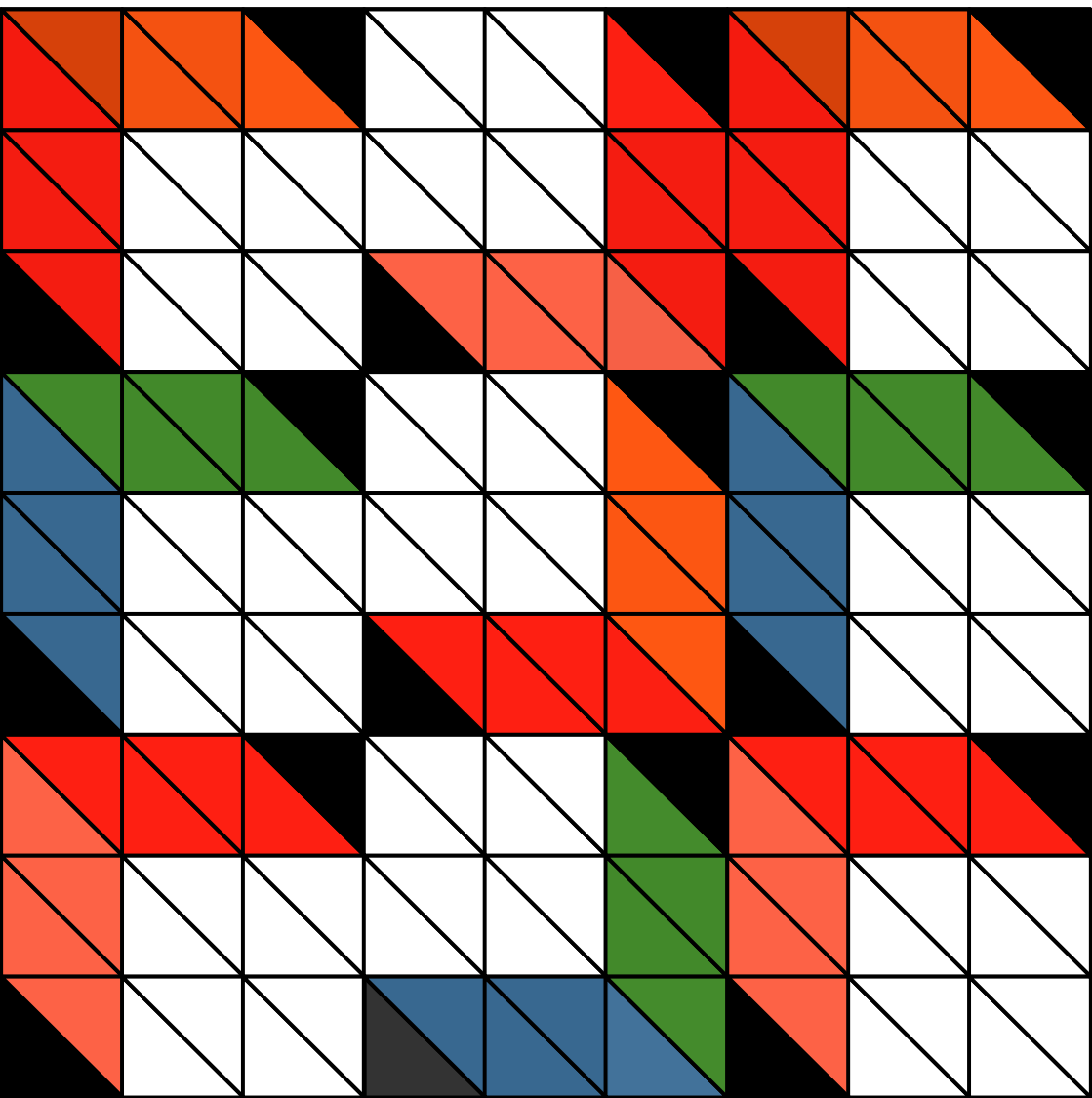
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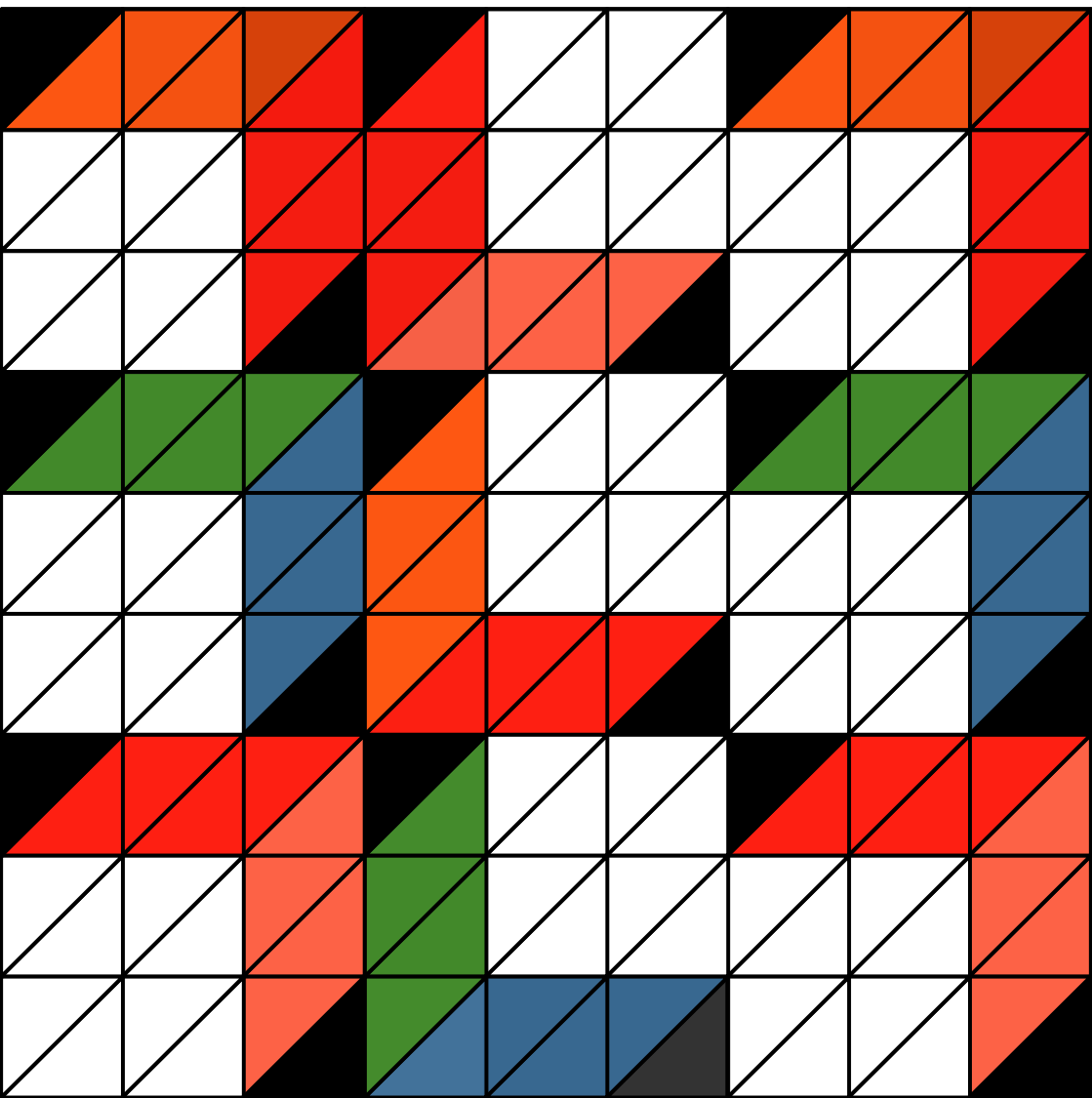
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Lev Tarasov

The World Is Built on Probability

This text is divided into two major parts.

The aim of the first part is to convince the reader that the random world begins directly in his or her own living room because, in fact, all modern life is based on probability. The first part is on the concept of probability and considers making decisions in conflict situations, optimizing queues, games, and the control of various processes, and doing random searches.

The second part of this text shows how fundamental chance is in nature using the probabilistic laws of modern physics and biology as examples. Elements of quantum mechanics are also involved, and this allows the author to demonstrate how probabilistic laws are basic to microscopic phenomena. The idea is that the reader, passing from the first part of the book to the second one, would see that probability is not only around us but it is at the basis of everything.



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