1

INTRODUCTION TO PHARMACOKINETICS AND PHARMACODYNAMICS

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1.1 Introduction: Drugs and Doses
1.2 Introduction to Pharmacodynamics
  1.2.1 Drug Effects at the Site of Action
    1.2.1.1 Interaction of a Drug with Its Receptor
    1.2.1.2 Postreceptor Events
  1.2.2 Agonists, Antagonists, and Concentration–Response Relationships
1.3 Introduction to Pharmacokinetics
  1.3.1 Plasma Concentration of Drugs
  1.3.2 Processes in Pharmacokinetics
1.4 Dose–Response Relationships
1.5 Therapeutic Range
  1.5.1 Determination of the Therapeutic Range
1.6 Summary

Objectives
The material in this chapter will enable the reader to:

1. Define pharmacodynamics and pharmacokinetics
2. Understand the processes that control the dose–response relationship
3. Gain a general appreciation of how mathematical expressions in pharmacodynamics and pharmacokinetics can be used for the rational determination of optimum dosing regimens

1.1 INTRODUCTION: DRUGS AND DOSES

Drugs may be defined as chemicals that alter physiological or biochemical processes in the body in a manner that makes them useful in the treatment, prevention, or cure of diseases. Based on this definition, any useful drug must affect body physiology or biochemistry. By extension, any useful drug must, if used inappropriately, possess the ability to do harm. Drug action begins with administration of the drug (input) and concludes with the biological response (output, which can be a beneficial and/or an adverse effect). The inputs (dose, frequency of administration, and route of administration) must be selected carefully to optimize the onset, intensity, and duration of therapeutic effects for a particular disease condition. At the same time, the inputs selected must minimize any harmful effects of drugs.

The design of optimum dosing regimens requires a complete understanding of the processes and steps that translate the input into the output. It also requires an understanding of how the input–output relationship may be influenced by individual patient characteristics that may exist at the very beginning of therapy, as well as conditions that may arise during the course of drug therapy. These will include the age and weight of the patient, the presence of other diseases, genetic factors, concurrent medications, and changes in the disease being treated over time.

The material presented in this book will address and explain why, as shown in Table 1.1, there is such tremendous variability in the value of drug doses and dosing frequencies among therapeutic drugs. Additionally, it will address why different routes of administration are used for different drugs and different indications (Table 1.1).

The steps between drug input and the emergence of the response can be broken down into two phases: pharmacokinetic and pharmacodynamic. The pharmacokinetic phase encompasses all the events between the administration of a dose and the achievement of drug concentrations throughout the body. The pharmacodynamic phase encompasses all the events between the arrival of the drug at its site of action and the onset, magnitude, and duration of the biological response (Figure 1.1). The rational design of optimum dosing regimens must be based on a thorough understanding of these two phases and will, ideally, include the development of one or more mathematical expressions for the relationship between dose and the time course of drug response.

Optimum drug administration is important not only for ensuring good patient outcomes in clinical practice, but also in the design of clinical trials during drug development. The

<table>
<thead>
<tr>
<th>Drug</th>
<th>Daily Dose (mg)</th>
<th>Dose Frequency (h)</th>
<th>Route</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium carbonate</td>
<td>3000</td>
<td>2</td>
<td>Oral</td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>1600</td>
<td>6</td>
<td>Oral</td>
</tr>
<tr>
<td>Vancomycin (for MRSA&lt;sup&gt;a&lt;/sup&gt;)</td>
<td>2000</td>
<td>12</td>
<td>Intravenous</td>
</tr>
<tr>
<td>Amoxicillin</td>
<td>750</td>
<td>8</td>
<td>Oral</td>
</tr>
<tr>
<td>Vancomycin (for pseudomembranous colitis)</td>
<td>1000</td>
<td>6</td>
<td>Oral</td>
</tr>
<tr>
<td>Atenolol</td>
<td>100</td>
<td>24</td>
<td>Oral</td>
</tr>
<tr>
<td>Fluoxetine</td>
<td>20</td>
<td>24</td>
<td>Oral</td>
</tr>
<tr>
<td>Ramipril</td>
<td>10</td>
<td>12</td>
<td>Oral</td>
</tr>
<tr>
<td>Digoxin</td>
<td>0.250</td>
<td>24</td>
<td>Oral</td>
</tr>
<tr>
<td>Chloroquine</td>
<td>300</td>
<td>Weekly</td>
<td>Oral</td>
</tr>
</tbody>
</table>

<sup>a</sup>Methicillin-resistant *Staphylococcus aureus*.
cost of drug research and development is enormous, so it is critical that all drug candidates selected for human trials are evaluated in the most efficient, cost-effective manner possible. The application of pharmacokinetic and pharmacodynamic principles to this process has been shown to enhance the selection of optimum doses and optimum designs of phase II clinical trials.

1.2 INTRODUCTION TO PHARMACODYNAMICS

Pharmacodynamics (PD) is the study of the magnitude of drug response. In particular, it is the study of the onset, intensity, and duration of drug response and how these are related to the concentration of a drug at its site of action. An overview of some basic drug terminology and the drug response–concentration relationship is provided below.

1.2.1 Drug Effects at the Site of Action

Note that although some references and textbooks distinguish the terms drug effect and drug response, this distinction has not been adopted universally. In this book, effect and response are used interchangeably.

1.2.1.1 Interaction of a Drug with Its Receptor

Drug response is initiated by a chemical interaction between a drug and a special binding site on a macromolecule in a tissue. This macromolecule is known as a drug receptor. The drug–receptor interaction results in a conformational change in the receptor, which results in the generation of a stimulus that ultimately leads to a biochemical or physiological response (Figure 1.2). Most receptors (over 95%) are proteins; however, other types of receptors exist such as the DNA receptors of the alkylating agents used in cancer chemotherapy. The drug–receptor interaction involves chemical bonding, which is usually reversible in nature and can be expressed using the law of mass action (Figure 1.2). Thus, at the site of action, the drug binds to its receptor and equilibrium is established between the bound and the unbound drug. As the drug is eliminated from the body and removed from its site of action, it dissociates from the receptor, which is left unchanged, and the response dissipates.
In contrast, a few drugs form irreversible covalent bonds with their receptors. For example, aspirin inhibits platelet aggregation by inhibiting the formation of thromboxane in the platelets. It accomplishes this by binding covalently to and blocking the catalytic activity of cyclooxygenase, the enzyme that produces thromboxane. The effect of a single dose of aspirin will persist long after the drug has been removed from its site of action and will continue until new cyclooxygenase molecules are synthesized, which can then resume the production of thromboxane. Other examples of drugs that bind irreversibly to their receptors include the alkylating agents mentioned above and proton pump inhibitors, such as omeprazole, which block the secretion of gastric acid by binding irreversibly to the H⁺, K⁺-ATPase pumps of parietal cells.

The drug–receptor interaction is highly dependent on the chemical structure of both the drug and the receptor and, therefore, small changes in the structure of the drug can reduce or destroy activity. For example, the drug–receptor interaction can distinguish between the R- and S-isomers of drugs that have chiral carbon atoms. Usually, one isomer is much more active than the other. The S-isomer of warfarin, for example, is two to five times more active than the R-isomer. The development and promotion of S-omeprazole (Nexium) is based on the premise that the S-isomer has the higher affinity for the binding site and thus offers therapeutic advantages over preparations containing racemic mixtures (equal quantities of each isomer) of omeprazole, such as Prilosec and its generic equivalents. Receptors are assumed to exist for all active endogenous compounds (natural ligands) such as neurotransmitters and hormones. The interaction between natural ligands and their receptors controls and/or regulates physiological and biochemical processes in the body. In most cases, drugs mimic or antagonize the actions of endogenous ligands by interacting with their cognate receptors. For example, epinephrine is a natural ligand that interacts with β₂-adrenergic receptors in bronchial smooth muscle to bring about bronchial dilation. Albuterol, a drug, also interacts with this receptor to produce bronchial dilation. Acetylcholine transmits signals through a synapse by interacting with its nicotinic receptor found on postsynaptic neuronal membranes. This interaction, which is mimicked by the drug nicotine, results in the production of a response called an action potential.

It should be noted that there are a few drugs that do not act on receptors but that exert their action by bringing about physicochemical changes in the body. For example, conventional
antacids, such as calcium carbonate, act as buffers to reduce acidity in the stomach and polyethylene glycol, an osmotic laxative, acts by preventing the absorption of water in the large intestine.

1.2.1.2 Postreceptor Events

Drugs almost always bring about some type of change in the *intracellular environment* of cells, but the lipophilic cell membrane presents a physical barrier to most drugs and endogenous ligands. As a result, most receptors are located on the cell membrane itself. The stimulus generated from the interaction of the drug with the membrane bound receptor has to be relayed to the inside of the cell. The relaying of the initial stimulus, known as *coupling* or *signal transduction*, often involves a cascade of different steps during which the initial signal may be amplified or diminished. Some important transduction mechanisms are summarized below (see Figure 1.3).

1. Interaction of a drug with a receptor can lead directly to the opening or closing of an *ion channel* that lies across a cell membrane. In this case, the signal is relayed by changes in the ion concentration within the cell. For example, the interaction of acetylcholine with its nicotinic receptor results in the opening of an ion channel allowing $\text{Na}^+$ to move into the cell thus, initiating the production of an action potential.

2. Signal transduction for a large number of drugs involves the *activation of a G-protein* (guanine nucleotide-binding protein). The drug–receptor interaction on the membrane triggers the activation of a G-protein on the cytoplasmic side of the membrane, which then initiates a series of events that culminate in the biological response. Activated G-protein can produce a variety of effects, including stimulation or inhibition of enzymes, and the opening or closing of ion channels. These events usually result in changes in the concentration of an intracellular compound known as the

![Diagram](image-url)

**FIGURE 1.3** Diagrammatic representations of how a drug receptor interaction brings about intracellular events. The intracellular relay of the initial signal resulting from the interaction of a drug with a membrane-bound receptor can be accomplished in one of three ways: (1) the direct opening of ion channels; (2) the activation of a G-protein that may lead to the activation of another enzyme or to a modulation of an ion channel; (3) the activation of protein kinase. Alternatively, (4), some drugs are able to penetrate membranes and directly activate intracellular receptors.
second messenger. Examples of second messengers include cyclic adenosine-3′,5′-monophosphate (cAMP), calcium, and phosphoinositides. The second messengers then relay the response further through a series of complex steps. For example, the interaction of catecholamines such as norepinephrine with certain β-receptor subtypes involves G-protein activation. This then stimulates adenylyl cyclase to convert adenosine triphosphate to cAMP, which acts as the second messenger. Subsequent events include the stimulation of specific protein kinases, activation of calcium channels, and modification of cellular proteins. Other examples of G-protein–coupled receptors are the action of acetylcholine on its muscarinic receptors and the action of serotonin on its 5-HT receptors.

3. The interaction of a drug with its receptor can also result in the stimulation of a receptor-associated enzyme, tyrosine kinase. The activated tyrosine kinase phosphorylates key macromolecules, which are often a part of the receptor itself, to relay the signal. Insulin and peptide growth factors, for example, use this form of signal transduction.

Some drugs are lipophilic enough to penetrate the cell membrane, while others may be transported across the cell membrane by uptake transporters. Drugs that are able to enter a cell can interact directly with intracellular receptors. Examples of drugs that act on intracellular receptors include many steroids such as glucocorticoid steroids, sex hormones, and thyroid hormones. The HMG-CoA reductase inhibitors (commonly known as statins) and metformin also act within the cell (hepatocyte) and both are dependent on uptake transporters to deliver them to the intracellular space and their site of action.

### 1.2.2 Agonists, Antagonists, and Concentration–Response Relationships

A drug that mimics the endogenous receptor ligand to activate the receptor is referred to as an agonist. The typical relationship between the drug effect and the agonist concentration at the receptor site is shown in Figure 1.4a. Note that as the concentration of the drug increases, the effect increases. At low concentrations, there is a linear relationship between concentration and effect (i.e., the response is proportional to the concentration). At higher drug

![FIGURE 1.4](image_url)  
**FIGURE 1.4** Plots of response versus drug concentration: (a) on a linear scale and (b) on a semilogarithmic scale.
concentrations, increases in concentration bring about much smaller changes in effect (the *law of limited returns*). Eventually, at very high concentrations, the effect achieves a maximum value and then remains constant and independent of concentration. In this area of the curve, increases in concentration will not result in further increases in response. This relationship is observed because response is generated by a saturable, capacity-limited process. For example, the response may be limited by the number of receptors that a tissue contains. At low drug concentrations, there are many free receptors and as the drug concentration increases, the drug can bind to the free receptors and response can increase proportionally. At higher concentrations, more and more of the receptors are occupied. As a result, increases in the drug concentration produce much less increase in effect. Eventually, all of the receptors are occupied (or saturated) and a maximum effect is observed. To accommodate a wide range of concentrations, the relationship between effect and concentration is usually plotted on a semilogarithmic scale, which transforms the plot to a sigmoidal shape (Figure 1.4b).

Many agonists are able to produce the system’s maximum response without fully occupying all the receptors. In these systems, the maximum response of the drug must be the result of some other saturable, capacity-limited process that occurs after receptor binding. These tissues or systems are said to have *spare receptors*. Experimentally, the presence of spare receptors can be demonstrated by destroying some of the receptors. If an agonist is still able to produce a maximum response, the system must contain spare receptors.

The efficiency with which a drug’s interaction with the receptor is converted into the initial stimulus or biosignal is a function of the number of receptors at the site of action and a drug’s *intrinsic efficacy*. Intrinsic efficacy can be defined as the magnitude of the stimulus produced per unit receptor occupied. The value of the stimulus that results from a specific concentration of a drug is also a function of the drug’s affinity for its receptors. Affinity can be defined as the extent or fraction to which a drug binds to receptors at any given drug concentration. Drugs that have high affinity require less drug to produce a certain degree of binding and to elicit a certain response compared to drugs with low affinity. Affinity is one of the factors that determines *potency* (see Chapter 19).

A drug that binds to a receptor but does not activate it is referred to as an *antagonist*. The presence of an antagonist at the receptor site blocks the action of the agonist (Figure 1.5). Higher concentrations of the agonist are needed to displace the antagonist and to produce the effect that is elicited when the antagonist was absent. The antagonist shifts the concentration–response curve of an agonist to the right (Figure 1.6). At sufficiently high concentrations of the antagonist, the agonist’s action may be blocked completely and the effect of even high concentrations of the agonist is reduced to zero. Some drugs bind to

![Diagrammatic representation of the action of an antagonist. The antagonist (ATG) binds to the receptor but does not produce a signal. Its presence on the receptor blocks the action of agonists (AG), including the natural ligand.](image-url)
receptors, but the binding is less efficient and a full response cannot be achieved even when the drug’s concentration is very high and all the receptors are occupied (Figure 1.7). These drugs are referred to as partial agonists. A partial agonist will block the effect of a full agonist. In the presence of high concentrations of a partial agonist, the action of a full agonist can be reduced to the maximum response elicited by the partial agonist. Clinically, partial agonists are used to act as buffers to avoid full stimulation of a system. Examples of partial agonists include several β-blockers, including pindolol, and the opioid buprenorphine. The latter is a partial agonist on the μ-opioid receptors and is considered a safer alternative to morphine because it does not produce as much respiratory depression (see Chapter 19).

In summary, drug action is mediated primarily by the interaction of a drug with membrane-bound receptors at its site of action. This produces conformational changes in the receptor, which lead to the generation of an initial signal. The signal is then relayed to the intracellular environment by means of a variety of transduction processes. The response increases with increases in drug concentration until enough receptors are occupied to generate the maximal response. The response to a specific concentration of drug is dependent on drug-specific properties (e.g., intrinsic efficacy and affinity) and tissue-specific properties (e.g., number or density of receptors and amplification or diminution of the initial signal during transduction). An important goal in a study of pharmacodynamics is to derive
a mathematical expression for the magnitude of drug response as a function of drug concentration:

$$E = f_{PD}(C)$$ (1.1)

where $E$ is the drug effect or response, $C$ is the drug concentration, and $f_{PD}$ is a pharmacodynamic function that links these two variables and contains the drug-specific parameters of intrinsic efficacy and affinity. In equation (1.1), $E$ is the dependent variable because it is dependent on all the other components of the equation. The drug concentration at the site of action ($C$) is the independent variable because it is independent of all the other components of equation (1.1). This expression would allow the effect to be estimated at any drug concentration and allow the required concentrations for optimum response to be identified.

1.3 INTRODUCTION TO PHARMACOKINETICS

Pharmaco- comes from the Greek word for “drug,” pharmakon, and kinetics comes from the Greek word for “moving,” kinetikos. Pharmacokinetics (PK) is the study of drug movement into, around, and out of the body. By extension, it involves the study of drug absorption, distribution, and elimination (metabolism and excretion) (ADME).

Pharmacokinetics involves the study of how drugs enter the body, distribute throughout the body, and leave the body. It is concerned with the driving forces for these processes and the rate at which they occur. Pharmacokinetics is the study of the time course of drug concentrations in body compartments. From a therapeutic perspective, the drug concentration at the site of action is by far the most important: Concentrations should be sufficiently high to produce a response but not so high as to produce toxicity. Since it is not possible to routinely measure this concentration clinically, the plasma concentration of the drug is the main focus in pharmacokinetics. It is often assumed that the plasma concentration reflects the drug concentration at the site of action. This is generally true and the relationship is often linear. Increases or decreases in the plasma concentration will be reflected by proportional increases or decreases at the site of action, respectively. However, as discussed in subsequent chapters, this is not always the case and a more complex relationship between these two concentrations may exist. It is important to note that although changes in the plasma concentration will usually result in proportional changes in the drug concentration at the site of action, the reverse is not true. Because the amount of drug that is delivered to the site of action is usually such a very small fraction of the total amount of drug in the body (in other tissues and the systemic circulation), local changes in the amount of drug at the site of action are generally not reflected by noticeable changes in the plasma concentration.

1.3.1 Plasma Concentration of Drugs

As stated above, pharmacokinetics is concerned with the body’s exposure to a drug and how drug concentrations change over time. For the most part, drug concentrations in the plasma are the focus in pharmacokinetics. The rationale for this is twofold. First, blood is one of the few body fluids that can be obtained and analyzed repeatedly for drug concentrations at specified times after the administration of a dose. The concentration of drug in whole blood is not commonly used in pharmacokinetics because blood is a complex physical system that consists of red blood cells, white blood cells, and platelets suspended in plasma water. Blood with the cellular elements removed, either by centrifugation (plasma) or clotting
INTRODUCTION TO PHARMACOKINETICS AND PHARMACODYNAMICS

Drug in tablet

G.I. membrane

Liver

Drug in tissues

Drug metabolism

Drug in systemic circulation

Drug in tissues

Drug excretion

Drug removed from body

Absorption

Distribution

Disposition

Elimination

FIGURE 1.8 Processes of drug absorption, distribution, and elimination (metabolism and excretion) (ADME). Drug contained within the tablet must undergo absorption. It must penetrate the gastrointestinal membrane and pass through the liver before reaching the systemic circulation. Once in the blood, it has the opportunity to distribute to the tissues, including the site of action. As soon as drug is present in the systemic circulation, it is subject to elimination. This occurs primarily in the liver and kidneys, where drugs undergo metabolism and/or excretion, respectively. The fate of a drug in the systemic circulation (distribution and elimination) is referred to as drug disposition.

(serum), is preferred. The collection of plasma requires the use of an anticoagulant such as heparin. However, heparin can interfere with the assay of some drugs. In these cases (e.g., for measuring digoxin concentration), serum rather than plasma is used as the reference fluid. In this book, no distinction will be made between plasma and serum, and the term plasma concentration will be used almost universally.

The second rationale for focusing on plasma concentrations in pharmacokinetics is that the circulatory system is the central fluid for the receipt and distribution of drugs (Figure 1.8). All drug input processes conclude when drug reaches the plasma, and all disposition (distribution and elimination) processes begin once drug is present in the plasma. Thus, drugs at absorption sites such as the gastrointestinal tract or subcutaneous tissue are absorbed into the circulatory system. Once in the blood, drugs undergo distribution to various tissues in the body and undergo elimination primarily through the liver and/or kidneys.

Plasma or plasma water consists of small dissolved molecules (e.g., glucose, ions, nutrients, and drugs) and suspended substances such as proteins, which are too large to dissolve. Many drugs can bind or associate with the plasma proteins. The binding is reversible and may be expressed according to the law of mass action:

\[
[D] + [P] \overset{k_1}{\underset{k_2}{\rightleftharpoons}} [DP] \tag{1.2}
\]
where \( D \) is the free drug concentration, \( P \) is the concentration of the protein not involved in binding, \( DP \) is the concentration of the drug–protein complex, and \( k_1 \) and \( k_2 \) are the rate constants for the forward and backward reaction, respectively.

Thus, many drugs exist in the plasma in an equilibrium between two forms: one component dissolved in the plasma water (free drug) and one component associated with or bound to plasma proteins (bound drug). The term plasma concentration \( (C_p) \) in pharmacokinetics refers to the total drug concentration of the drug, that is, the bound plus the free drug. Total drug concentrations are reported routinely because they are much easier and less expensive to measure than free drug concentrations. However, as presented in subsequent chapters, the free concentration is the clinically important component: Only free unbound drug is able to pass biological membranes, interact with the receptor, and generate a pharmacological response.

1.3.2 Processes in Pharmacokinetics

Pharmacokinetics involves the study of the processes that affect the plasma concentration of a drug at any time after the administration of a dose. These processes are summarized in Figure 1.8. Most drugs are administered orally as tablets. A tablet is a compressed powder mass that consists of the active drug, which usually comprises only a small portion of the overall tablet, and other compounds required for either the manufacture of the tablet (i.e., diluents and lubricants) or to optimize the characteristics of the finished product (i.e., color, taste, and hardness). Once a tablet is swallowed, it enters the stomach, where the drug contained within the hard powder mass must be exposed and released. The tablet must first disintegrate into small particles to enable the drug to dissolve in the gastrointestinal fluid. These initial processes of disintegration and dissolution are part of biopharmaceutics, which may be defined as the study of how a drug’s chemical and physical properties influence both the administration of the drug and the pharmacokinetic behavior of the dosage form in vivo. When the drug is dissolved in the gastrointestinal fluid, it has the opportunity to pass across the epithelial cell lining of the gastrointestinal membrane and get taken up into the blood on the other side. Once in the circulatory system, the drug has to pass through the liver, which is a major organ of drug elimination. The absorbed drug may undergo elimination by metabolism during its first pass through the liver. After passing through the liver, the drug is taken to the heart, which pumps the drug throughout the entire circulatory system. At this point, the drug has been absorbed. The rate and extent of absorption of a drug are very important determinants of the early plasma concentrations of a drug. Rapid rates of absorption will promote high early plasma concentrations. Once the heart pumps the drug around the body, the drug is given the opportunity to distribute to all the tissues, including the biophase or site of action. A drug’s distribution pattern, particularly the rate and extent to which it distributes to the tissues, is also an important determinant of the early plasma concentrations. If a drug distributes extensively to the tissues, little drug will be left in the plasma and the plasma concentration will be low. The plasma concentration will also be influenced by drug elimination, which occurs as soon as the drug is in the plasma. The main pathways of elimination are hepatic metabolism and renal excretion. The process of drug elimination will continue to affect the plasma concentration until the drug has been removed from the body completely.

In summary, a drug’s pharmacokinetics are determined by the simultaneous processes of ADME (Figure 1.8). The combined processes of drug elimination and drug distribution or the fate of a drug once it is present in the body is referred to as drug disposition.
12 INTRODUCTION TO PHARMACOKINETICS AND PHARMACODYNAMICS

TABLE 1.2 Pharmacokinetic Processes That Control the Dose–Plasma Concentration Relationship after the Consumption of a Tablet

<table>
<thead>
<tr>
<th>Process</th>
<th>Type of Process</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Release of drug: tablet disintegration</td>
<td>Biopharmaceutics</td>
</tr>
<tr>
<td>2 Dissolution of tablet</td>
<td>Biopharmaceutics</td>
</tr>
<tr>
<td>3 Absorption of drug through gastrointestinal membrane into the blood</td>
<td>Absorption</td>
</tr>
<tr>
<td>4 Passage through the liver</td>
<td>Absorption</td>
</tr>
<tr>
<td>5 Entry to systemic circulation</td>
<td>Absorption</td>
</tr>
<tr>
<td>6 Distribution to the biophase</td>
<td>Biophase distribution</td>
</tr>
<tr>
<td>7 Distribution throughout the body</td>
<td>Distribution</td>
</tr>
<tr>
<td>8 Elimination (metabolism and excretion)</td>
<td>Elimination</td>
</tr>
</tbody>
</table>

(Figure 1.8). The individual pharmacokinetic steps associated with the administration of a tablet are summarized in Table 1.2.

The goal of pharmacokinetics is to study each of the ADME processes with the aim of:

1. Identifying the drug and patient factors that determine the rate and extent of each process. Topics to be considered include:
   - How does a drug’s lipophilicity influence absorption, distribution, and elimination?
   - What factors determine a drug’s distribution pattern?
   - Is the whole of a dose absorbed into the body?
   - Does a drug get to every tissue in the body?
   - To what extent, do drugs undergo renal as opposed to hepatic elimination?
   - How are pharmacokinetic processes affected by patient characteristics, such as the age of the patient, renal or hepatic impairment, ethnicity, and genetics?

2. Identifying a way to quantify or summarize each process in ADME using a single parameter. Issues to be considered include:
   - How can the extent of absorption of a drug be quantified?
   - How can the extent to which a drug distributes the tissues be quantified?

3. Deriving a mathematical expression for the rate of each process in ADME and for the overall relationship between a drug’s plasma concentration and time after any dose:

\[
C_p = f_{PK} \text{ (dose, time)} \tag{1.3}
\]

where \( C_p \) is the plasma concentration, and \( f_{PK} \) is a function that contains expressions and parameters for ADME. In equation (1.3), \( C_p \) is the dependent variable because it is dependent on all the other components of the equation, time is the independent variable, and dose is a constant in a given situation.

1.4 DOSE–RESPONSE RELATIONSHIPS

It will become apparent in subsequent chapters that for most drugs, the drug concentration in the body at any time is proportional to the dose. As a result, plots of response at a certain time as a function of dose (Figure 1.9) resemble the plots of response versus concentration.
DOSE–RESPONSE RELATIONSHIPS

Response

Logarithm of dose

FIGURE 1.9 Graph of response versus logarithm of dose.

(Figure 1.4): A hyperbolic plot is often observed on the linear scale, and a sigmoidal plot is observed on the semilogarithmic scale. Thus, dose–response curves are analogous, but not identical, to pharmacodynamic concentration–effect curves.

In contrast to the plots of response versus concentration, which are purely dependent on a drug’s pharmacodynamics, a dose–response curve is a function of both the drug’s pharmacodynamic characteristics (intrinsic efficacy and affinity) and its pharmacokinetic characteristics (the fraction of the dose absorbed, the extent to which a drug distributes throughout the body, etc.). Note that low doses produce no effect, and as the maximum response is approached, increasing the dose produces little change in the response (limiting returns). Based on the characteristics shown in Figure 1.9, doses must be selected to avoid the subtherapeutic areas of the plot and to avoid doses that approach or lie on the plateau that provide little or no additional benefit over lower doses. Most drugs also produce toxicity at higher concentrations, and it is important that doses are selected that minimize this toxicity. The toxicity may be an extension of the drug’s pharmacological action (e.g., the major adverse effects of warfarin, digoxin, and anticholinergic drugs), in which case it is important to avoid areas on the dose–response curve close to the maximum effect. Alternatively, the toxicity may arise because the drug may interact with multiple receptors of different types, particularly at higher concentrations, to produce undesired effects. Examples of this type of toxicity include muscle toxicity associated with the statins and drowsiness associated with first-generation antihistamines. The development of models and mathematical expressions of the pharmacokinetic and pharmacodynamic phases of drug response provides an opportunity for the rational selection of optimum dosing regimens.

The expression for a drug’s pharmacokinetics [equation (1.3)] can be combined with the expression for a drug’s pharmacodynamics [equation (1.1)] to produce a complete expression for the dose–response relationship:

$$E = f_{PD}(f_{PK}(\text{dose}, \text{time}))$$

(1.4)

Note that in this equation, the plasma concentration of the drug ($C_p$) has been substituted for the drug concentration at the site of action ($C$) in the pharmacodynamic equation. This assumes that the concentration at the site is always proportional to the plasma concentration. The validity and limitations of this are discussed in subsequent chapters. Equation (1.4) enables the full time course of drug response to be estimated after any dose. It could also be used to estimate the dose and dosing interval to produce optimum response. If these relationships are identified early in the course of drug development, they can be used to
INTRODUCTION TO PHARMACOKINETICS AND PHARMACODYNAMICS

determine optimum doses for clinical trials. This in turn will increase the efficiency of trials, reduce the time for drug development, and decrease the price of these highly costly studies. The expressions can also be used to simulate response data for situations not yet studied clinically. For example, if a drug’s pharmacokinetics and pharmacodynamics are known after a single dose, it is possible to use a combined PK–PD equation to simulate the response that may be expected during multiple dosing therapy. Simulations can be performed using different dosing regimens to try to obtain an estimate of what may be the most effective dosing regimen.

1.5 THERAPEUTIC RANGE

In vivo pharmacodynamic studies aimed at developing mathematical expressions of drug response are relatively new. Historically, in vivo pharmacodynamic studies have been very difficult to perform. Some reasons for this are presented below:

1. It is difficult to obtain precise measurements of drug response. Meaningful models and mathematical expressions for drug response require that response data be collected on a continuous scale. The data must also possess a reasonable degree of precision. All-or-none responses and subjective data, based largely on a patient’s or a physician’s opinion, have limited value in this application. The response to only a handful of drugs (e.g., anticoagulants and hypoglycemic agents) meets these criteria. In the last 10–20 years, this problem has been overcome by the development and use of biomarkers (see Chapter 19) of drug response. Biomarkers are parallel changes in the levels or intensities of concrete measurable biological molecules or other effects that have been found to be predictably associated with a drug’s biological response. Examples of biomarkers may include cells, proteins, antibodies, body temperature, or features of an electroencephalograph.

2. The mathematical expressions derived from pharmacodynamic models are mainly nonlinear and could not be applied to clinical data until computer software became available for nonlinear regression analysis.

3. Each drug or drug class has a unique mechanism of action and way of relaying or coupling the initial drug effect. Signal transduction may take less than a second for some drugs, several minutes for others, or up to several hours for others. As a result, summarizing the characteristics of the concentration–response relationship can be complex.

4. In many cases, a drug’s response lags behind the plasma concentration. This can confound the concentration–response relationship and add an additional layer of complexity to modeling response as a function of plasma concentration.

By contrast, pharmacokinetic studies are relatively simple to perform. Blood is easily sampled, drug assays for most drugs are fairly easily developed, and the data analysis is relatively straightforward and could be performed even before the wide availability of computers and software for pharmacokinetic analysis, by linearizing the mathematical expressions and analyzing the data using simple linear regression analysis. Furthermore, the pharmacokinetics of most drugs can be modeled using one of about three basic well-established models. As a result, pharmacokinetic studies and modeling have been a central part of the drug development process for decades. In order to use pharmacokinetic models for the design of dosing regimens, it is necessary to have target-optimal plasma concentrations
or some idea of the concentration–response relationship. In the absence of mathematical expressions for this relationship, a very simple approach for linking drug concentrations to response was developed and termed the therapeutic range. The therapeutic range is defined as the range of plasma concentrations that are associated with optimum response and minimal toxicity in most patients. Most commonly, the goal of therapy is to maintain drug concentrations within the therapeutic range at all times. There are a small number of drugs for which this is not desirable, such as certain antibiotics and drugs like nitroglycerin, where tolerance develops with continuous exposure to the drug.

The therapeutic range is illustrated in Figure 1.10, which shows:

- The **minimum effective concentration** (MEC) is the lower boundary for effective drug concentrations; plasma concentrations below the MEC have a high probability of being subtherapeutic.
- The **maximum tolerated concentration** (MTC) is the upper boundary for optimum drug concentrations; plasma concentrations above the MTC have a high probability of producing adverse effects or toxicity.
- The **onset of action** of a drug, which may be estimated as the time it takes for plasma concentrations to reach the MEC.
- The **duration of action** of a drug, which may be estimated as the time during which plasma concentrations remain within the therapeutic range.

### 1.5.1 Determination of the Therapeutic Range

To apply the therapeutic range appropriately, and to understand both its value and limitations, it is necessary to appreciate how it is typically derived. It is usually determined by studying the effects of a drug in a large population and noting the plasma concentrations at which patients:

- experience therapeutic effects;
- experience side effects or toxicity.
The cumulative plot of the percentage of all patients who experience a therapeutic response is then plotted as a function of plasma concentration (Figure 1.11). The cumulative plot of the percentage of patients experiencing adverse effects at the various concentrations is then added to the same graph (Figure 1.11). Similar sigmoidal shapes are obtained for both curves, but the plot for toxicity is always displaced to the right. Higher concentrations are needed for adverse compared to therapeutic effects (if this were not the case, the drug would not be of therapeutic value). A frequent characteristic of these plots is that although 100% of patients experience toxicity if concentrations are high enough, fewer than 100% of patients experience therapeutic effects even at high concentrations. Patients who do not respond therapeutically even to high concentrations are referred to as nonresponders.

This plot is then used to estimate a drug’s therapeutic range. The MEC and MTC are usually chosen at concentrations where a high percentage of patients experience a therapeutic effect and a small percentage of patients experience toxicity, respectively. The specific concentrations selected for the MEC and the MTC will depend on the margin of safety and the risk–benefit ratio acceptable for a given indication. For example, the MTC for an over-the-counter analgesic or nonsteroidal anti-inflammatory drug will be chosen at a concentration associated with much less toxicity than that of a drug used to treat a life-threatening condition such as cancer. In Figure 1.11, the MEC was selected as the concentration at which 70% of the population experienced a therapeutic benefit, and the MTC was selected as the concentration at which 10% of the population experienced some adverse effects.

The therapeutic range has been enormously useful clinically, particularly in helping clinicians determine optimum doses of drugs that have both narrow therapeutic ranges and wide interpatient variability in dose requirements. Examples of these drugs are shown in Table 1.3. A dose that is optimum for one patient (i.e., a dose that gives plasma concentrations in the therapeutic range) may produce concentrations below the MEC in a second patient and produce concentrations above the MTC in a third patient. As a result, doses are
TABLE 1.3 Therapeutic Ranges of Example Drugs [1]

<table>
<thead>
<tr>
<th>Drug</th>
<th>Therapeutic Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cyclosporine</td>
<td>100–400$^a$ μg/L, whole blood HPLC$^b$ analysis</td>
</tr>
<tr>
<td>Digoxin</td>
<td>0.5–2$^c$ μg/L</td>
</tr>
<tr>
<td>Lithium</td>
<td>0.6–1.5 mEq/L</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>10–20 mg/L</td>
</tr>
<tr>
<td>Tacrolimus</td>
<td>5–20$^a$ μg/L, whole blood</td>
</tr>
<tr>
<td>Theophylline</td>
<td>5–15 mg/L</td>
</tr>
</tbody>
</table>

$^a$Depending on the time after transplant, the type of transplant, and the preference of the center.

$^b$High-performance liquid chromatography.

$^c$Depending on the indication.

frequently individualized by measuring plasma concentrations achieved by a typical dose and then applying pharmacokinetic principles to calculate a dose that will provide concentrations in the therapeutic range.

It is, however, important to recognize that the therapeutic range has limitations, which include:

1. It represents the range of concentrations that are optimum for most people. Certain patients will, however, experience therapeutic effects at concentrations below the MEC, and others will experience toxicity below the MTC. Some patients never respond therapeutically to a drug even at concentrations well above the MTC.

2. It does not incorporate a graded concentration-related response (i.e., a response that increases with increases in concentration). It is an all-or-nothing response: Patients are predicted to respond when the plasma concentration is within the established therapeutic plasma concentration range and not to respond when the plasma concentration is below the MEC.

3. It only applies to plasma concentrations that are in equilibrium with the drug concentrations at the site of action. It can take a long time for some drugs to distribute to their site of action. For example, it takes about 6–8 h for digoxin to fully distribute to its site of action (the myocardium of the heart). During this distribution period, the therapeutic range will not apply. For example, serum concentrations above the MTC in this period will not necessarily be associated with toxicity.

**Therapeutic Index (TI) or Therapeutic Ratio**  
Like the therapeutic range, the TI or therapeutic ratio is a way to express the safety margin offered by a drug. It is the ratio of the dose of the drug that produces toxicity in 50% of patients to the dose of the drug that produces therapeutic response in 50% of patients:

$$TI = \frac{TD_{50}}{ED_{50}}$$  (1.5)

where $TD_{50}$ is the dose that produces toxicity in half the patients, and $ED_{50}$ is the therapeutic or effective dose in half the patients. If, for example, a drug has a TI of 100, the toxic dose is about 100 times larger than the effective dose and the drug has a wide safety margin. Conversely, a TI of 3 would indicate a small margin of safety. A drug with a small therapeutic ratio will have a narrow therapeutic range.
1.6 SUMMARY

In summary:

- **Pharmacokinetics** may be defined as a study of the relationship between drug concentration and time after the administration of a given dose. It involves the study of all the processes that affect this relationship: that is, a drug’s ADME. Pharmacokinetics represents the first stage in the process of drug response.

- In pharmacokinetics, the plasma concentrations of a drug are usually studied. A goal is to derive a mathematical expression for the relationship between the plasma concentration, dose, and time:

\[
C_p = f_{PK} (\text{dose, time}) \tag{1.6}
\]

where \(C_p\) is the plasma concentration of the drug, and \(f_{PK}\) is a function that describes the relationships among \(C_p\), dose, and time. The function incorporates the drug’s pharmacokinetic parameters.

- **Pharmacodynamics** may be defined as a study of the relationship between drug concentration at the site of action and the onset, duration, and intensity of response to the drug. The pharmacodynamic phase constitutes the second and final step in drug response.

- A goal is to derive a mathematical expression for the relationship between the response and the drug concentration:

\[
E = f_{PD} (C) \tag{1.7}
\]

where \(E\) is the drug effect or response, \(C\) is the concentration at the site of action, and \(f_{PD}\) is a function that describes the relationship between the two and incorporates a drug’s pharmacodynamic parameters.

- Integrating pharmacokinetics and pharmacodynamics covers the entire dose–response relationship. Mathematical expressions for the pharmacokinetic and pharmacodynamic phases can be combined to provide a complete mathematical expression of the dose–response relationship:

\[
E = f_{PD} (f_{PK} (\text{dose, time})) \tag{1.8}
\]

- Equation (1.8) provides a complete expression for the time course of drug response. It will allow the drug response to be calculated at any time after any dose. It will allow optimum dosing regimens to be determined and can be used to simulate drug response data in situations not studied clinically.

**REFERENCE**