The nature of immunity

Key objectives

This chapter will enable you to:

✔ Outline the general purpose and properties of the immune system
✔ Distinguish the features of innate and adaptive immunity and the components of the immune system involved in each
✔ Give an overview of the processes that generate an immune response appropriate for defence of the body against the pathogen that instigated the response

Immunology is a relatively new science. It is a branch of biomedical science that covers the study of all aspects of immunity and the immune system in organisms. Immunity is a state of resistance to infection, and the immune system is composed of those constituents of the body (i.e. molecules, cells, tissues and organs) that contribute to generating this resistance. Immunology deals with the physiological functioning of the immune system in states of both health and disease. It incorporates malfunctions of the immune system, as well as the physical, chemical and physiological characteristics of the components of the immune system. These components can be studied individually (in isolation), in terms of their mutual interactions, and within the body as a whole. Its origin is usually attributed to events in the eighteenth century - firstly, to Lady Mary Wortley Montagu, the wife of the British ambassador in Constantinople (now known as Istanbul), who observed the positive effects of variolation on the native population in 1712. Variolation is the deliberate infection with smallpox, where dried smallpox scabs were blown into the nose of an individual who then contracted a mild form of the disease. She introduced it into England with royal patronage following initial experiments on condemned criminals and orphaned children. However, this procedure was not without risk of causing smallpox (variola) itself and the high morbidity and mortality associated with it made others look for less dangerous and more effective ways of controlling the disease.

Subsequently, Edward Jenner, a Gloucestershire family doctor, made the important observation that dairymaids, who frequently contracted cowpox (an infection of the hands acquired during milking), were remarkably resistant to smallpox and did not develop the disfigured pock-marked faces of those who had had smallpox infection.

Edward Jenner had suffered painfully from variolation performed when he was 8 years old. The increasing spread of smallpox throughout the population led him to develop the alternative technique of vaccination. This was first performed in 1796 when he inoculated material obtained from cowpox pustules into the arm of a healthy boy. Jenner was subsequently able to inoculate the boy with smallpox more than 20 times without any untoward effect. This courageous experiment aroused much criticism, but Jenner offered his new preventive treatment to all who sought it and performed many of his vaccinations in a thatched hut - which became known as the Temple of Vaccinia - in the grounds of his house at Berkeley. These are restored and now contain the Jenner Museum and a Conference Centre.
Although this vaccination was successful, it took almost two centuries (1796–1980) before the World Health Organization (WHO) was able to announce that smallpox was eradicated in 1980.

Interestingly, Jenner knew nothing of infectious agents that cause disease. Numerous scientific breakthroughs occurred, but it was not until Robert Koch first demonstrated, in 1876, that bacterial infectious agents (pathogens) cause disease. Any organism with the potential to cause disease is called a pathogen. There are five broad categories of pathogens, namely, viruses, bacteria, fungi, other relatively large and eukaryotic organisms (termed parasites) and prions (Table 1.1).

<table>
<thead>
<tr>
<th>Type of pathogen</th>
<th>Example pathogens</th>
<th>Diseases</th>
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<tbody>
<tr>
<td>Virus</td>
<td>Variola</td>
<td>Smallpox</td>
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<td></td>
<td>Human immunodeficiency virus</td>
<td>Acquired immune deficiency syndrome</td>
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<tr>
<td>Bacteria</td>
<td>Staphylococcus aureus or Streptococcus pneumoniae</td>
<td>Community-acquired pneumonia</td>
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<tr>
<td>Fungi</td>
<td>Candida albicans</td>
<td>Thrush</td>
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<tr>
<td>Parasite</td>
<td>Plasmodium malariae</td>
<td>Malaria</td>
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<tr>
<td>Prion</td>
<td>Creutzfeldt–Jakob disease (CJD) prion</td>
<td>CJD</td>
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Recognition and defence components

Before considering the complexity of the immune system as it exists, it is useful to consider some of the general design requirements of an immune system in order for it to protect the host organism. Clearly, the two important biological events are recognition of the target pathogen and effective defence against it. A major consideration is how many recognition specificities are required and how many kinds of defence, that is, methods of pathogen destruction, are necessary.

There exists an enormous variety of infectious pathogens, including many types of bacteria, viruses, fungi, parasites and prions, each with its own mechanisms of transmission, infection and reproduction. This means that no single recognition or defensive strategy is effective against all pathogens and therefore a wide variety of cellular and secreted components are present within the body that collectively constitute the immune system. Examples of the main cells and molecules of the immune system are given in Tables 1.2 and 1.3, respectively. These components vary in terms of whether their main role is recognition or defence, although most possess a combination of these properties.

Innate and adaptive immunity

The cellular components that mediate recognition and defence can be categorized by various criteria, including their developmental lineage from stem cells in the bone marrow (myeloid or lymphoid) and their morphology as mature blood leucocytes (Table 1.2). Polymorphonuclear leucocytes (PMNs) are distinguished from mononuclear cells by their lobulated nuclei, and they largely coincide with the granulocytes defined by distinctive cytoplasmic granules. The immune system’s cellular components can also be considered as mediators of either innate or adaptive immunity (Table 1.2).

The recognition properties associated with innate immunity may have evolved to recognize chemical structures that are characteristic of infectious pathogens and differ from constituents of host organisms. These include various microbial lipids, carbohydrates, proteins and even nucleic acids that are collectively termed pathogen-associated molecular patterns (PAMPs). They are bound by secreted proteins (e.g. mannose-binding lectin and C-reactive protein) and by cell surface and cytoplasmic proteins (e.g. macrophage mannose receptor and Toll-like receptors) called pattern recognition molecules that are inflexible in their specificities and identical between cells; these are considered in detail in Chapter 2. Innate immunity is rapidly activated in the early stages of an infection, and its defensive properties can limit the
proliferation and spread of a pathogen within the body. However, it is only moderately efficient in clearing infection, and its capabilities remain the same on repeated exposure to the same microbe.

The resolution of an infection usually requires an additional adaptive immune response by T lymphocytes and B lymphocytes (often referred to simply as T cells and B cells). Each lymphocyte specifically recognizes an individual antigen (usually a protein but also other types of chemical for B lymphocytes), and there are mechanisms for enhancing the specificity of recognition. Thus, the antigen receptor expressed by a particular lymphocyte is different from that of virtually all other lymphocytes in the body. In addition, the B lymphocytes produce and secrete a soluble form of their antigen receptors called antibodies or immunoglobulins. An adaptive immune response takes longer to activate than innate immunity but generates more effective defence which improves upon repeated exposure to the same microbe. The details of antigen recognition are considered in Chapter 2, and the development, activation and functions of lymphocytes are described in Chapters 3 and 4; immunoglobulins are considered in Chapter 5.

It is the cardinal features of adaptive immunity mediated by lymphocytes that Edward Jenner recognized in immunity to smallpox and utilized in the development of vaccination. Furthermore, an individual who is immune to smallpox will not be protected against diphtheria unless she/he has also met the *Corynebacterium diphtheriae* on a previous occasion. This illustrates the specificity of the adaptive immune response. Lymphocytes can detect remarkably small chemical differences between antigens, for example, subtly differing strains of influenza virus, minor substitutions of a benzene ring or the difference between dextro and laevo isomers. Were it not for the fact that cowpox and smallpox viruses share important antigens, the experiments of Jenner would have been a dismal failure (although he would not have attempted them without the evidence of the milkmaids).

Another feature of adaptive immune responses is the memory that develops from previous experiences of foreign material — a characteristic that enables immunization to be of clinical value. This altered

<table>
<thead>
<tr>
<th>Table 1.2 Cells of the immune system.</th>
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<tr>
<td><strong>Cell type</strong></td>
</tr>
<tr>
<td>Neutrophils</td>
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<tr>
<td>Eosinophils</td>
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<tr>
<td>Basophils</td>
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<tr>
<td>Mast cells</td>
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<tr>
<td>Monocytes/macrophages</td>
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<tr>
<td>Dendritic cells</td>
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<tr>
<td>Natural killer cells</td>
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<tr>
<td>Cytotoxic T lymphocytes</td>
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<td>Helper T lymphocytes</td>
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<tr>
<td>B lymphocytes</td>
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PMN, polymorphonuclear.

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<th>Table 1.3 Secreted mediators of immunity.</th>
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<tr>
<td><strong>Antimicrobial</strong></td>
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<tr>
<td>Antibodies/immunoglobulins (IgM, IgG, IgA, IgE, IgD)</td>
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<tr>
<td>Pentraxins (e.g. C-reactive protein)</td>
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<tr>
<td>Collectins (e.g. mannan-binding lectin)</td>
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<tr>
<td>Complement proteins</td>
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<tr>
<td>Defensins</td>
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<tr>
<td>Lytic enzymes</td>
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<tr>
<td>Interferons</td>
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<tr>
<td>Cytotoxins (perforins, granzymes)</td>
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<tr>
<td><strong>Regulatory/inflammatory</strong></td>
</tr>
<tr>
<td>Cytokines (e.g. interleukins, interferons, tumour necrosis factors)</td>
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<tr>
<td>Chemokines (and other chemoattractants)</td>
</tr>
<tr>
<td>Eicosanoids (e.g. prostaglandins, leukotrienes)</td>
</tr>
<tr>
<td>Histamine</td>
</tr>
</tbody>
</table>
reactivity may last for the entire lifespan of the individual. The ability of an organism to respond more rapidly and to a greater degree when confronted with the same antigen on a second occasion is illustrated in Figure 1.1. This compares the speed and magnitude of the human response to an antigen that the subjects had not previously encountered (bacteriophage φX174). In the first or primary response, there is a delay of at least 10 days before the antibody level in the circulation reaches its maximum, and this level shows considerable variation between individuals, rarely exceeding a titre of 1000. In the secondary response, all individuals respond maximally within 10 days, and in all cases, the levels attained are of a titre of 10,000 or more. The outcome of an acute infection is often a close race between the activities of the replicating pathogen and the adaptive immune response, and it is for this reason that prior exposure, for example, to a vaccine, can give the host a considerable advantage.

A third important feature of adaptive immune responses is self-discrimination, which is illustrated in Figure 1.2. If split-skin grafts are placed on the flanks of rodents, it is possible to observe within 2 weeks whether they have healed well and been accepted (Fig. 1.2a) or whether they have been rejected (Fig. 1.2b). In this experiment, the successful graft was obtained from another animal of identical genetic composition (i.e. another member of the same inbred strain). The rejected graft came from an unrelated member of the same species. These chemical differences are relatively minor and demonstrate not only the recognition ability of lymphocytes but also the efficient way in which they fail to react against tissue of ‘self’ origin. Previously, it was thought that components of the immune system failed to recognize self at all, but it is now clear that self-recognition does occur in a controlled and regulated manner such that – except in the special circumstance of autoimmune disease – tissue damage does not take place.

**Stages of an immune response**

The different properties of innate and adaptive immunity mean that they are complementary, and they cooperate with each other in order to give the best possible defence. This can be exemplified by considering the stages of a generalized response to a bacterial skin infection (Fig. 1.3; Clinical Case Scenario 1.1). The skin itself constitutes an effective barrier to infection

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**Figure 1.1** Primary and secondary antibody responses following intravenous injection of bacteriophage φX174 used as a test antigen in humans (Data kindly provided by Drs Peacock and Verrier Jones).

**Figure 1.2** Discrimination between self and non-self illustrated by skin grafting in a rodent. (a) The graft was of ‘self’ type. (b) The graft was from an unrelated animal.
because most microbes cannot penetrate the hard, keratinized surface of the epidermis (Fig. 1.3a), but if this is breached (e.g. by a cut), then microbes may infiltrate and start to replicate in the softer underlying dermal tissues. If this is a primary response to infection (because it is the first time this particular microbe has infected the body), then there will be no immunological memory to generate an early adaptive response, but components of the innate immune system that are resident in the infected tissues can be rapidly activated, including complement proteins in the tissue fluid and macrophages (Fig. 1.3b). The activation of a range of complement proteins triggered by interactions with bacterial surface molecules may result in bacterial lysis by the membrane attack complex of complement and/or opsonization.

Figure 1.3 Stages of an immune response to bacterial skin infection. (a) Bacteria are unable to penetrate the intact keratinized epithelial barrier. (b) Bacterial entry (e.g. via a cut) stimulates an immediate local innate response by tissue macrophages and complement proteins. (c) An early induced inflammatory response is stimulated by inflammatory mediators produced by macrophages, complement proteins and mast cells, leading to infiltration by neutrophils from the blood and plasma containing more complement. (d) Bacterial antigens are carried in the lymph, and associated with dendritic cells (DC), to draining lymph nodes where specific T and B lymphocytes are activated. These lymphocytes, and antibodies produced by the B cells, return to the infected tissues via the blood circulation.
Darren, a fit and healthy 27-year-old, attended a walk-in clinic, concerned about a swelling on his left calf. Six days earlier, he had taken part in a cross-country race and had grazed his calf on some brambles. Although the skin had been broken and the wound bled, he had thought nothing more of it at the time and completed the race. However, he was now worried that the lesion might be infected.

When the doctor examined the affected area on Darren’s calf, she noted that it was swollen and red; Darren told her that the area felt warm and slightly painful when touched. The swelling was circular, with a diameter of about 6 cm and a diffuse edge. There was a small amount of pus in the centre of the wound. Darren’s temperature was normal and there were no systemic signs of infection (e.g. fever, chills, malaise).

The doctor concluded that this was an uncomplicated case of cellulitis (inflammation of the skin and subcutaneous tissues associated with acute infection) in a healthy young adult and that the site of inflammation on the calf was associated with a normal, acute immune response to localized infection. She considered that the infection was most probably streptococcal or possibly staphylococcal (but unlikely to be methicillin-resistant *Staphylococcus aureus* (MRSA)). The doctor prescribed an oral course of high-dose flucloxacinil antibiotic that inhibits bacterial cell wall synthesis and is effective against both streptococci and staphylococci. She advised Darren to complete the course of antibiotics. Within a week, the swelling, redness and tenderness resolved and Darren experienced no further problems.

(For further information relevant to this case, see Chapter 10, p. 118.)

(i.e. coating) of the bacteria by complement proteins that help to adhere the bacteria to the macrophages, which express complement receptors as well as the pattern recognition molecules mentioned in the ‘Innate and Adaptive Immunity’ section. Macrophages are phagocytes that can engulf microbes and bring about their digestion.

The activation of complement proteins and macrophages not only results in microbial destruction directly but also induces amplifying events (i.e. inflammation). In addition, tissue-resident mast cells, which are a major source of inflammatory mediators, are activated by complement-derived peptides. These amplifying events can be divided into several categories: local vasodilation and increase in vascular permeability; adhesion of inflammatory cells to the blood vessel wall; their chemical attraction, that is, chemotaxis; immobilization of cells at the site of infection; and activation of the relevant cells and molecules to liberate their lytic products (Fig. 1.4). In the present example, the inflammatory mediators induce the influx of leucocytes (particularly neutrophils that, like macrophages, are phagocytes) and plasma containing further supplies of complement proteins (Fig. 1.3c).

While the innate response is being established during the first few hours and days of the infection, the processes are being set in train to generate the adaptive response. This involves the transport of microbial components (i.e. antigens) from the site of infection to neighbouring lymphoid tissues, which is where the majority of lymphocytes reside transiently during their circulation around the body. Lymphocytes develop in primary lymphoid organs, consisting of bone marrow and thymus, in the adult. Lymphocytes circulate through lymph nodes, the white pulp of the spleen and mucosa-associated lymphoid tissue: these locations are referred to as secondary lymphoid organs (Fig. 1.5). The total weight of these various lymphoid components can exceed that of the liver. It is at these sites that large numbers (i.e. hundreds of millions) of the different varieties of lymphocyte come into intimate contact with each other and with specialized antigen-presenting cells so as to provide an optimal environment for the activation of the small proportion of the body’s lymphocytes that specifically recognize the antigens of a particular microbe. This is why antigens are carried to lymphoid tissues to induce lymphocyte activation rather than these interactions occurring initially within the site of infection. For example, tissue fluid that drains from infected tissues into the lymphatic system may carry microbial antigens to draining lymph nodes where they can be recognized by specific B cells. In addition, microbial antigens are captured and processed by antigen-presenting cells, called dendritic cells, which are present in most tissues. The dendritic cells then migrate to the draining lymph nodes where they present the antigens to T cells (Fig. 1.3d). The activated T and B cells return to the blood circulation whence they enter the inflamed, infected tissues,
together with antibodies secreted by terminally differentiated B cells called **plasma cells**, in a similar manner to the earlier influx of other leucocytes and complement proteins (Fig. 1.3d). The efficiency of bacterial elimination will then be enhanced by antibodies that opsonize the bacteria, thereby augmenting complement activation and phagocytosis, and regulatory proteins called **cytokines** produced by the T cells that increase the antimicrobial activity of the phagocytes.

**Figure 1.4** Amplifying events involved in the local recruitment of inflammatory cells and molecules from the circulation into an extravascular site of infection.

**Figure 1.5** The lymphoid system in humans, showing the distribution of primary and secondary lymphoid organs and tissues.
Some of the T and B cells activated by antigens of the infecting microbe revert to a resting state and constitute the body’s population of memory lymphocytes specific for that microbe. A subsequent infection with the same, or a closely related (i.e. antigenically similar), microbe would induce a faster and bigger secondary response by these lymphocytes, as described earlier in this chapter.

**Immunological defence strategies**

The nature of the defensive strategy that the immune system employs in order to eliminate a microbe is determined not only by the biological nature of the microbe but also by the tissue compartment in which the infection is concentrated. In particular, it is critical whether the microbe remains *extracellular* (i.e. in fluids or at the surfaces of cells of the tissues it infects) or enters the cytoplasm of cells to become *intracellular*. Extracellular pathogens (including many types of bacteria and parasitic worms), which do not cross the plasma membrane of cells, are vulnerable to opsonization by antibodies and complement proteins; bacteria can then be phagocytosed by macrophages and neutrophils, and parasitic worms are attacked by eosinophils (Fig. 1.6a). Antibodies and complement proteins are considered in Chapters 5 and 6, respectively, phagocytes in Chapter 7 and eosinophils in Chapter 8. However, some phagocytosed microbes are resistant to intracellular digestion and can survive and replicate in cytoplasmic vesicles of macrophages where they are no longer exposed to antibodies and complement: mycobacteria that cause tuberculosis and leprosy are important examples of this. The macrophages must then be stimulated into a heightened state of activation by

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**Figure 1.6** Immunological defence strategies. (a) Extracellular pathogens (e.g. bacteria and parasitic worms) are directly exposed to antibodies, complement, phagocytes (macrophages and neutrophils) and eosinophils. (b) Microbes that are resistant to digestion (e.g. mycobacteria) can survive intracellularly in macrophage vesicles, but Th1-derived cytokines (e.g. γ-interferon) can enhance the digestive activity of the macrophages. (c) Intracellular pathogens, like viruses that generate cytosolic antigens, are targeted by interferons that block their replication and killer cells (NK and Tc cells) that induce apoptosis of the infected cells.
cytokines derived from helper T lymphocytes (Th cells) in order to overcome the microbes’ resistance to digestion (Fig. 1.6b). Some microbes deliberately invade cells; this applies to all viruses, which hijack the metabolic machinery of the cells they parasitize in order to replicate. In order to combat intracellular viruses, interferons induce an antiviral state in cells, which inhibits viral replication. In addition, natural killer (NK) cells and cytotoxic T lymphocytes (Tc cells), which are described in Chapter 9, deliberately kill infected cells, thus inhibiting viral replication (Fig. 1.6c).

### An overview

The frontispiece gives a schematic overview of the cells and secreted mediators of immunity that have been introduced in this chapter and that are discussed in more detail in the following chapters. This shows how these components of the immune system interact and cooperate to generate the various defensive options that are effective against different categories of infective agents.

Pathogens are the source of the PAMPs and antigens necessary for the generation of innate and adaptive immunity, respectively, and these pathogens then become the target of the integrated innate and adaptive defensive response that is generated. The activities of the immune components that generate rapid innate responses (e.g. macrophages, granulocytes, NK cells and complement proteins) are greatly enhanced by the addition of the adaptive response by T and B lymphocytes. Dendritic cells are a pivotal link between innate and adaptive immunity; they are activated by microbial PAMPs that interact with their pattern recognition molecules, together with ‘danger signals’ released by stressed and damaged cells (e.g. ‘DAMPs’ discussed in Chapter 2). This activation of dendritic cells enables them to efficiently activate T cells by presenting antigens.

The dendritic cells and other cell types also provide ‘polarizing signals’ that promote naive helper T-cell (Th0 cell) differentiation into various T-cell subsets that are characterized by the production and secretion of different regulatory proteins called cytokines that stimulate different cellular activities. The frontispiece shows these main functional subsets (Th1, Th2, Th17 and Treg), some of the key cytokines they produce and the cell types on which they act; this is discussed in detail in Chapter 4. Furthermore, although all B cells are initially programmed to produce classes of immunoglobulins called IgM and IgD, the different T-cell subsets promote immunoglobulin class switching in B cells to produce other classes of antibodies (IgG, IgE or IgA) that have different functional properties; this is discussed in detail in Chapter 5.

The frontispiece summarizes the particular combinations of immune effector cells and secreted mediators that are orchestrated by T-cell-derived cytokines and B-cell-derived antibodies to generate the combinations of defensive and inflammatory activities appropriate for the nature of the infections generated by particular pathogens. Overall, the purpose of the polarizing signals is to ensure that the qualitative nature of the immune response generated is appropriate to the nature of the pathogen whose PAMPs and antigens triggered the response.

The abundance of means by which recognition and defence can be achieved is necessary to meet the enormous task that confronts the immune system, that is, the constant threat to the survival of the host from a universe of pathogenic organisms ranging from the smallest viruses through bacteria, protozoa and fungi and to metazoan parasites with their often complex life cycles. The remarkable ability of successful pathogens to evolve mechanisms by which they can evade the immune response adds a further dimension, which is considered in detail in Chapter 10.

### Immunopathology

The outcome for the host is often ‘survival at a price’ and damage to host tissues by the immune system is a common finding during the course of most infectious diseases. When the reaction is excessive or inappropriate, major tissue damage can ensue and this is referred to as hypersensitivity. When the driving force for the response is from a non-infectious agent (i.e. innocuous), this leads to allergy. However, when the source is from self-components (i.e. self-tissue), this leads to the development of autoimmunity. Some pathogens are also able to initiate various forms of lymphoproliferative disease and can cause immunodeficiency. Immunopathology is composed of these various deviations from the ideal, many examples of which are found in human disease. These are described in Part 2.
Chapter 1 The nature of immunity

**KEY POINTS**

- Immunity provides protection against pathogenic organisms, and the functions of the immune system are recognition of these foreign pathogens and defence of the body against them.
- A wide variety of cells and secreted molecules necessarily constitute the immune system in order to deal with the vast array of pathogenic organisms that can infect the body.
- Innate and adaptive immunity constitute the two main arms of the immune system which differ in their recognition and activation properties. Adaptive immunity improves upon repeated exposure to the same antigens.
- Lymphocytes are responsible for the specificity, memory and self-discrimination of adaptive immune responses, as exemplified in vaccination.
- The early innate response to infection stimulates an inflammatory response and initiates the processes that result in the adaptive lymphocyte response.
- Defence is mediated by a range of innate and adaptive cells and molecules that stimulate inflammation and cause digestion or lysis of foreign pathogens or infected cells. Different defence mechanisms are required to deal with extracellular pathogens and with intracellular pathogens either in macrophage vesicles or in the cytosol.
- Polarizing signals generated in response to the stimulation of the immune system by a particular pathogen promote the type of immune response appropriate to that pathogen.
- Different subsets of T cells (e.g. Th1, Th2, Treg and Th17) stimulate the production of different classes of antibodies and the activation of different immune effector cells and molecules, thereby generating qualitatively different immune responses that are required for the eradication of different pathogens.
- Activation of the immune system can also lead to damage to host tissues, and some pathogens can induce abnormal proliferation of the lymphoid system or cause it to become deficient. These together constitute the various forms of immunopathology.

**SELF-ASSESSMENT QUESTIONS**

(Answers to these questions are given on p. 215.)

1.1 From the following list of cell types, choose the most appropriate in answer to the questions below:

- B cells
- Basophils
- Dendritic cells
- Eosinophils
- Macrophages
- NK cells
- Neutrophils
- Tc cells
- Th cells

(a) Which three cell types generate adaptive immunity?
(b) Which cell type can differentiate into antibody-producing plasma cells?
(c) Which two cell types are particularly associated with killing virus-infected cells?
(d) Which two cell types are phagocytes?

(e) Which cell type specializes in transporting antigens to secondary lymphoid tissues for presentation to T cells?
(f) Which three cell types are defined as granulocytes?

1.2 Which two of the following are primary lymphoid organs?

- Bone marrow
- Lymph node
- Mucosa-associated lymphoid tissue
- Spleen
- Thymus

1.3 Which of the following statements refer to innate immunity, and which refer to adaptive immunity?

(a) It is induced by pathogen-associated molecular patterns.
(b) It involves recognition of antigens.
(c) It generates immunological memory.
(d) It is rapidly activated in the initial phase of an immune response.
NOTES

1 Further information can be obtained from the Custodian, The Chantry, Church Lane, High Street, Berkeley, Gloucestershire GL13 9BH, UK.

2 The titre is the reciprocal of the weakest dilution of serum at which antibody can still be detected.