Part 1

Epidemiology
Epidemiology: defining disease and normality

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Learning objectives

In this chapter you will learn:
✓ what is meant by the term epidemiology;
✓ the concepts underlying the terms ‘normal, abnormal and disease’ from a (i) sociocultural, (ii) statistical, (iii) prognostic, (iv) clinical perspective;
✓ how one may define a case in epidemiological studies.

What is epidemiology?

Trying to explain what an epidemiologist does for a living can be complicated. Most people think it has something to do with skin (so you’re a dermatologist?) wrongly ascribing the origin of the word to epidermis. In fact the Greek origin is epidēmia – ‘prevalence of disease’ (taken from the Oxford online dictionary) – and the more appropriate related term is epidemic. The formal definition is

‘The study of the occurrence and distribution of health-related states or events in specified populations, including the study of the determinants influencing such states and the application of this knowledge to control the health problems’ (taken from the 5th edition of the Dictionary of Epidemiology)

An alternative way to explain this and easier to comprehend is that epidemiology has three aims (3Ws).

<table>
<thead>
<tr>
<th>Whether</th>
<th>To describe whether the burden of diseases or health-related states (such as smoking rates) are similar across different populations (descriptive epidemiology)</th>
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<tbody>
<tr>
<td>Why</td>
<td>To identify why some populations or individuals are at greater risk of disease (risk-factor epidemiology) and hence identify causal factors</td>
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<tr>
<td>What</td>
<td>To measure the need for health services, their use and effects (evidence-based medicine) and public policies (Public Health) that may prevent disease – what we can do to improve the health of the population</td>
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Population versus clinical epidemiology – what’s in a name?

The concept of a population is fundamental to epidemiology and statistical methods (see Chapter 3) and has a special meaning. It may reflect the inhabitants of a geographical area (lay sense of the term) but it usually has a much broader meaning to a collection or unit of individuals who share some characteristic. For example, individuals who work in a specific industry (e.g., nuclear power workers), born in a specific week and year (birth cohort), students studying medicine etc. In fact, the term population can be extended to institutions as well as people; so, for example, we can refer to a population of hospitals, general practices, schools etc.

Populations can either consist of individuals who have been selected irrespective of whether they have the condition which is being studied or specifically because they have the condition of interest. Studies that are designed to try and understand the causes of disease (aetiology) are usually population-based as they start off with healthy individuals who are then followed up to see which risk factors predict disease (population-based epidemiology). Sometimes they can select patients with disease and compare them to a control group of individuals without disease (see Chapter 5 for observational study designs). The results of these studies help doctors, health-policy-makers and governments decide about the best way to prevent disease. In contrast, studies that are designed to help us understand how best to diagnose disease, predict its natural history or what is the best treatment will use a population of individuals with symptoms or clinically diagnosed disease (clinical epidemiology). These studies are used by clinicians or organisations that advise about the management of disease. The term clinical epidemiology is now more often referred to as evidence-based medicine or health-services research. The same methodological approaches apply to both sets of research questions but the underlying questions are rather different.

One of the classical studies in epidemiology is known as the Framingham Heart Study (see http://www.framinghamheartstudy.org/about/history.html). This study was initially set up in 1948 and has been following up around 5200 men and women ever since (prospective cohort study). Its contribution to medicine has been immense, being one of the first studies to identify the importance of elevated cholesterol and high blood pressure in increasing the risk of heart disease and stroke. Subsequent randomised trials then went on to show that lowering of these risk factors could importantly reduce risk of these diseases. Furthermore, the Framingham risk equation, a prognostic tool, is commonly used in primary care to identify individuals who are at greater risk of future coronary heart disease and to target interventions (see http://hp2010.nhlbihin.net/atpiii/calculator.asp).

Regardless of the purpose of epidemiological research, it is always essential to define the disease or health state that is of interest. To understand disease or pathology, we must first be able to define what is normal or abnormal. In clinical medicine this is often obvious but as the rest of this chapter will illustrate, epidemiology has a broader and often pragmatic basis for defining disease and other health-related states.

What is dis-ease?

Doctors generally see a central part of their job as treating people who are not ‘at ease’ – or who in other words suffer ‘dis-ease’ – and tend not to concern themselves with people who are ‘at ease’. But what is a disease? We may have no difficulty justifying why someone who has had a cerebrovascular accident (stroke), or someone who has severe shortness of breath due to asthma, has a disease. But other instances fit in less easily with this notion of disease. Is hypertension (high blood pressure) a disease state, given that most people with raised blood pressure are totally unaware of the fact and have no symptoms? Is a large but stable port wine stain of the skin a disease? Does someone with very protruding ears have a disease? Does someone who experiences false beliefs or delusions and imagines her/him-self to be Napoleon Bonaparte suffer from a disease?

The discomfort or ‘dis-ease’ felt by some of these individuals – notably those with skin impairments – is as much due to the likely reaction of others around the sufferer as it is due to the intrinsic features of the problem. Diseases may thus in some cases be dependent on subjects’ sociocultural environment. In other cases this is not so – the sufferer would still suffer even if marooned alone on a desert island. The purpose of this next section is to offer a structure to the way we define disease.
A sociocultural perspective

Perceptions of disease have varied greatly over the last 400 years. Particular sets of symptoms and signs have been viewed as ‘abnormal’ at one point in history and ‘normal’ at another. In addition, some sets of symptoms have been viewed simultaneously as ‘abnormal’ in one social group and ‘normal’ in another.

Examples abound of historical diseases that we now consider normal. The ancient Greek thinker Aristotle believed that women in general were inherently abnormal and that female gender was in itself a disease state. In the late eighteenth century a leading American physician (Benjamin Rush) believed that blackness of the skin (or as he termed it ‘negritude’) was a disease, akin to leprosy. Victorian doctors believed that women with healthy sexual appetites were suffering from the disease of nymphomania and recommended surgical cures.

There are other examples of states that we now consider to be diseases, which were viewed in a different light historically. Many nineteenth-century writers and artists believed that tuberculosis actually enhanced female beauty and the wasting that the disease produces was viewed as an expression of angelic spirituality. In the sixteenth and seventeenth centuries gout (joint inflammation due to deposition of uric acid) was widely seen as a great asset, because it was believed to protect against other, worse diseases. Ironically, recent research interest has suggested a potential protective role of elevated uric acid, which may cause gout, for both heart and Parkinson’s disease.

In Shakespeare’s time melancholy (what we would now call depression) was regarded as a fashionable state for the upper classes, but was by contrast stigmatised and considered unattractive among the poor. The modern French sociologist Foucault points out that from the eighteenth century onwards those who showed signs of what we would now call mental illness were increasingly confined in institutions, as tolerance of ‘unreason’ declined. Whereas previously ‘mad’ people had often been viewed as having fascinating and desirable powers (and were legitimised as holy fools and jesters), increasingly they were seen as both disruptive and in need of treatment. Other examples exist of the redefinition of socially unacceptable behaviour as a disease. Well into the second half of the last century single mothers were viewed as being ill and were frequently confined for many years in psychiatric institutions.

As some diseases have been accepted as part of the normal spectrum of human behaviours so new ones have been labelled. Newly recognised diseases include alcoholism (previously thought of simply as heavy drinking), suicide (previously thought of as a criminal offence, it was illegal in the UK until the 1960s so that failed suicides were prosecuted and successful suicides forfeited all their property to the State), and psychosomatic illness (previously dismissed as mere malingering).

Some new disease categories have arisen simply because new tests and investigations allow important differences to be recognised among what were previously thought of as single diseases. For example people died in past times of what was believed to be the single disease of dropsy (peripheral oedema), which we now know to be a feature of a wide range of diseases ranging across primary heart disease, lung disease, kidney disease and venous disease of the legs. There are still disagreements in modern medicine about the classification of disease states. For example, controversy remains around the underlying pathophysiology of chronic fatigue syndrome (myalgic encephalomyelitis) and Gulf War syndrome.

The sociocultural context of health, illness and the determinants of health-care-seeking behaviour as well as the potential adverse effects of labelling and stigma are main topics of interest for medical sociologists and health psychologists and the interested reader may wish to read further in other texts (see Further reading at the end of this chapter).

Abnormal as unusual (statistical)

In clinical medicine – especially in laboratory testing – it is common to label values that are unusual as being abnormal. If, for example, a blood sample is sent to a hospital haematology laboratory for measurement of haemoglobin concentration the result form that is returned may contain the following guidance (the absolute values will differ for different laboratories and units will differ by country):
This **reference range** is derived as follows: a large number (several hundred) of **samples** from people believed to be free of disease (usually blood donors) are measured and the reference range is defined as that central part of the range which contains 95% of the values. By definition, this approach will result in 5% of individuals who may be completely well, being classified as having an **abnormal** test result.

### Normal (Gaussian) distributions

In practice, as with haemoglobin concentration above, many distributions in medical statistics may be described by the **Normal**, also known as **Gaussian distribution**. It is worth noting that the statistical term for ‘Normal’ bears no relation to the general use of the term ‘normal’ by clinicians. In statistics, the term simply relates to the name of a particular form of frequency distribution. The curve of the Normal distribution is symmetrical about the **mean** (see Chapter 2) and bell-shaped.

The theoretical Normal distribution is continuous. Even when the variable is not measured precisely, its distribution may be a good approximation to the Normal distribution. For example in Figure 1.1, heights of men in South Wales were measured to the nearest cm, but are approximately Normal.

### Abnormal as increased risk of future disease (prognostic)

An alternative definition of abnormality is one based on an increased risk of future disease. A biochemical measure in an asymptomatic (undiagnosed) individual may or may not be associated with future disease in a **causal** way (see Chapter 7). For example, a raised C-reactive protein level in the blood indicates infection or inflammation. Whilst noncausally related, epidemiological studies demonstrate that C-reactive protein can also predict those at an increased future risk of coronary heart disease (CHD). Treatments focused on lowering C-reactive protein will not necessarily reduce the risk of CHD.

In a man of 50 years a systolic blood pressure of 150 mm Hg is well within the usual range and may not produce any clinical symptoms. However, his risk of a fatal myocardial infarction (heart attack) is about twice that of someone with a low blood pressure.

- Does he have a disease, and should he be treated?
- What factors might influence this decision?

These are important questions to consider when we come to think of disease in terms of increased risk of future adverse health outcomes.

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### Figure 1.1 Heights of 1,000 men in South Wales.

Note: This figure is known as a **histogram** and is used for displaying grouped numerical data (see Chapter 2) in which the relative frequencies are represented by the areas of the bars (as opposed to a **bar chart** used to display categorical data, where frequencies are represented by the heights of the bars).

The superimposed continuous curve denotes the theoretical Normal distribution.
Thresholds for introducing treatment for blood pressure have changed over the years, generally drifting downwards. This is due to two main factors:

(1) researchers have gradually extended their limits of interest as they have become more confident that blood pressure well within usual limits may have adverse effects in the future.

(2) newer drugs have tended to have fewer and less dangerous side effects, making it reasonable to consider extending treatment to lower levels of blood pressure, where the benefits – though present – are less striking.

Blood glucose levels provide similar problems to blood pressure levels – specifically, for type II diabetes which is treated with diet control, tablets and occasionally insulin (rather than type I which requires insulin as a life-saving measure). At what blood glucose level should one attach the label ‘diabetic’ and consider starting treatment? To address these questions large prospective studies (called cohort studies) are required. In such studies, subjects have a potential risk factor such as blood glucose levels measured at the beginning of the study. They are then followed up, sometimes for many years, to examine whether rates of disease differ according to levels of blood glucose at the start of the study.

Does a fasting glucose in a healthy individual have any implication for their future health?

The glucose tolerance test is commonly used as a diagnostic aid for diabetes. In one of the very early epidemiological studies, conducted in Bedford UK (Keen et al., 1979), 552 subjects had their blood glucose measured when fasting and again two hours after a 50 g glucose drink. On the basis of this they were classified as having high, medium or low glucose levels. The cohort was then followed up for ten years, at which point the pattern of deaths that had occurred was as illustrated in Table 1.1.

Amongst both men and women, those with high levels of glucose following the glucose tolerance test had an increased risk of all causes and cardiovascular death. In addition, the female medium glucose group had an increased risk compared to the low glucose group. This additional risk is far less dramatic amongst the men in this study. Basing a definition of abnormality on future 10-year risk of death, treatment might be considered for women with a medium glucose level in addition to those with a high glucose level.

Based on studies such as this, the World Health Organisation (WHO) recommends levels of blood glucose, which should be regarded as indicating diabetes and therefore considered for treatment (fasting glucose \( \geq 7.0 \) mmol/L (126 mg/dl) and/or 2 hour post-load glucose \( \geq 11.1 \) mmol/L (200 mg/dl)). It also identifies an intermediate risk group who are said to have Impaired Glucose Tolerance or borderline diabetes (fasting glucose \( < 7.0 \) mmol/L and 2 hour post-load glucose \( \geq 7.8 \) mmol/L but \( < 11.1 \) mmol/L). Such individuals are not generally treated but may legitimately be kept under increased surveillance. However, the increased risk of cardiovascular disease appears to show a linear relationship with fasting glucose with no obvious threshold. A recent WHO report concluded ‘there are insufficient data to accurately define normal glucose levels, the term normoglycaemia should be used for glucose levels associated with low risk of developing diabetes or cardiovascular disease’ (WHO/IDF, 2006).

Abnormal as clinical disease

It is better to define values of a particular test as abnormal if they are clearly associated with the presence of a disease state – rather than simply being unusual. However this is often less than straightforward.

The range of values describing diseased individuals is rarely clearly and completely separated from that for healthy individuals. The nice bell shaped curve described above may actually be bimodal with a second superimposed distribution either at the top (see Figure 1.2) or bottom end or both. This overlap means that there will be healthy people with ‘abnormal’ results and people with disease with apparently ‘normal’ results (see Chapter 9 on diagnostic tests for more details).

For example, it is widely believed by many doctors that chronic (i.e. of long duration) mildly reduced haemoglobin (Hb) levels (of 100–110 g/L) or anaemia, such as might be seen in menstruating females, may account for fatigue and tiredness. In a study of 295 subjects in South Wales no association was found between Hb level and fatigue until the Hb level fell to well below 100 g/L (Wood
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Table 1.1 Glucose tolerance and mortality in the Bedfordshire cohort.

<table>
<thead>
<tr>
<th>Glucose group</th>
<th>Number</th>
<th>All deaths</th>
<th>Cardiovascular deaths</th>
<th>Number</th>
<th>All deaths</th>
<th>Cardiovascular deaths</th>
</tr>
</thead>
<tbody>
<tr>
<td>High glucose</td>
<td>51</td>
<td>19 (37.2%)</td>
<td>15 (29.4%)</td>
<td>63</td>
<td>25 (39.7%)</td>
<td>18 (28.5%)</td>
</tr>
<tr>
<td>Medium glucose</td>
<td>130</td>
<td>29 (22.3%)</td>
<td>19 (14.6%)</td>
<td>119</td>
<td>35 (29.4%)</td>
<td>25 (21.0%)</td>
</tr>
<tr>
<td>Low glucose</td>
<td>104</td>
<td>20 (19.2%)</td>
<td>12 (11.5%)</td>
<td>85</td>
<td>9 (10.6%)</td>
<td>4 (4.7%)</td>
</tr>
</tbody>
</table>

*Oral glucose tolerance test: After an overnight fast the participant is asked to drink a solution containing 1.75 g/kg body weight (maximum 75 g) of glucose dissolved in 250 ml of water within 2–3 minutes. Blood samples are taken just before and two hours after ingestion of the glucose solution.

and Elwood, 1966). Fatigue is common in the population generally for a wide range of reasons and is only strongly associated with Hb level among severely anaemic individuals. A longstanding Hb of between 100 and 115 g/L (which it should be noted is outside the laboratory reference range, whose lower limit is 115 in women and 130 in men) in an otherwise healthy person who is complaining only of fatigue shouldn’t therefore generally be considered as responsible for this symptom.

In general, the definition of abnormality as clinical abnormality is both logical and clear. It is nevertheless an approach that usually involves thinking in terms of the probability of disease being present, rather than the certainty.

Defining a case in epidemiological studies

Before an epidemiologist is able to study any disease s/he needs to develop and agree upon a case definition: a definition of disease that is as free as possible of ambiguity. This should enable researchers to apply this definition reliably on a large number of subjects, without access to sophisticated investigations. Because epidemiological case definitions are not used as a guide to the treatment of individuals they may differ from the sorts of definitions used in routine clinical practice.

Chronic Fatigue Syndrome provides a good example of the problems of agreeing on a case definition for a rather ill-defined condition. At a meeting in Oxford in 1990, 28 UK experts met to agree a case definition for Chronic Fatigue Syndrome (Sharpe *et al*., 1991). They came up with the following:

- Fatigue must be the principal symptom.
- There must be a definite point of onset (fatigue must not have been lifelong).
- Fatigue must have been present for at least 6 months and present for at least 50% of that time.
- Other symptoms may be present – e.g. myalgia (muscle pain), mood and sleep disturbance.
- Certain patients should be excluded: those with medical conditions known to produce chronic fatigue (such as severe anaemia); patients with a current diagnosis of schizophrenia, manic-depressive illness, substance abuse, eating disorder.

What is being attempted here is to produce a reasonably reliable definition (one that will classify the same person in the same way when used repeatedly by different observers) that can be applied without recourse to sophisticated tests, that excludes already well recognised causes of fatigue such as anaemia but which encompasses relevant patients.

This has now been updated in the UK by NICE guidelines (2007) that state a diagnosis should be
made after other possible diagnoses have been excluded and the symptoms have persisted for 4 months in an adult and 3 months in a child or young person (a shorter duration than previously stated). They suggest guidelines based on expert consensus opinion (see Box 1.1).

The use by both UK and American epidemiologists of the descriptive term ‘Chronic Fatigue Syndrome’ rather than ‘Post-viral Fatigue Syndrome’ is deliberate. The term implies no particular etiology (cause) unlike ‘Post-viral Fatigue Syndrome’, which presupposes that a viral cause is established and which may therefore inhibit exploration of other possible causes.

The NICE definition is intended to be used by clinicians and often ‘research case definitions’ are stricter so that some true cases are missed but you are less likely to include any false positive cases. So for example the USA Centre for Disease Control and Prevention case definition still has a requirement for a 6-month minimum period of symptoms.

Box 1.1 Symptoms that may indicate CFS/ME.

Consider the possibility of CFS/ME if a person has:

- fatigue with all of the following features:
  - new or a specific onset (i.e. not lifelong)
  - persistent and/or recurrent
  - unexplained by other conditions
  - has resulted in a substantial reduction in activity level characterised by post-exertional malaise and/or fatigue (typically delayed, e.g. by at least 24 hours, with slow recovery over several days)

and

- one or more of the following symptoms:
  - difficulty with sleeping, such as insomnia, hypersomnia, unrefreshing sleep, a disturbed sleep–wake cycle
  - muscle and/or joint pain that is multi-site and without evidence of inflammation
  - headaches
  - painful lymph nodes without pathological enlargement
  - sore throat
  - cognitive dysfunction, such as difficulty thinking, inability to concentrate, impairment of short-term memory, and difficulties with word-finding, planning/organising thoughts and information processing
  - physical or mental exertion makes symptoms worse
  - general malaise or ‘flu-like’ symptoms
  - dizziness and/or nausea
  - palpitations in the absence of identified cardiac pathology

The symptoms of CFS/ME fluctuate in severity and may change in nature over time.


REFERENCES


**FURTHER READING**
