The incidence of adverse reactions to drugs

O. L. WADE

From the Department of Clinical Pharmacology, University of Birmingham

Concern about adverse reactions to drugs is not new. In 1887 at a meeting in Manchester the British Medical Association set up a small committee to examine the sudden and unexpected deaths which sometimes occurred during chloroform anaesthesia. This was the first collaborative investigation of an adverse reaction of a drug in human subjects. Professor McKendrick, Professor of Physiology in Glasgow, and Dr Coats, pathologist, and Dr Newman, pathological chemist, at the Western Infirmary, Glasgow (1880) reported that chloroform was dangerous in man not only because excessive doses depressed respiration but because small doses had a deleterious effect on the heart and might cause cardiac arrest. This report was contrary to the experience of many who had used chloroform extensively. Its findings were repudiated by Surgeon Major Lawrie, Residency Surgeon at Hyderabad in the Deccan, with such heat that it is surprising that the now yellowing pages of the Lancet of 1889 and 1890 are not a little charred. Major Lawrie appointed two Hyderabad Chloroform Commissions, on both of which he sat as President. Following the extensive administration of chloroform to dogs that Commission in its final report stated, in the words of a contemporary Anglo-Indian, that chloroform given in doses which did not cause respiratory depression was ‘as safe as whisky and water’ (Lawrie, Brunton, Bomford, and Hakin, 1890). But the report received little attention in Britain for here it was already indisputable that in human beings sudden death did sometimes occur in the early stages of chloroform anaesthesia.

This ancient controversy is worth recalling. It not only showed for the first time the enormous emotion that may be aroused in doctors when a drug which they value is found to cause serious adverse reactions, but it also demonstrated, although the lesson has yet to be fully appreciated, that the administration of a drug to animals may fail to reveal its hazards in man.

Towards the end of the first World War another collaborative investigation concerning an adverse reaction to a drug was instituted. The Medical Research Committee, later to become the present Medical Research Council, appointed a ‘Special Committee on the Manufacture, Biological Testing and Clinical Administration of Salvarsan and its Substitutes’. Its enquiries followed an outbreak of acute yellow atrophy due to neoarsphenamine benzoate at Cherry Hinton Hospital near Cambridge in 1917 and 1918. The Committee reported (Medical Research Council, 1922) that the most probable cause of the outbreak was the toxicity of organoarsenical compounds. But Professor Stuart McDonald, Professor of Pathology at Newcastle, suspected that there was an additional factor causing the epidemic of jaundice which was probably microbial infection (McDonald, 1918). The aetiology of this jaundice was in retrospect probably serum hepatitis. By a strange coincidence the Committee also made a report of a peculiar outbreak of malaria among patients treated with neoarsphenamine ‘606’ at Portobello Hospital, Dublin. Eight soldiers had died, and Professor A. C. O'Sullivan, Professor of Pathology at Trinity, found their organs full of malaria parasites, although only one of them was known to have served in a tropical country. The Committee found no reason to dissent from Professor O’Sullivan’s opinion that the infection had been conveyed from one or more carriers of the disease to others through the apparatus for injecting ‘606’, the blood of the carrier regurgitating into the last segment of the rubber tubing remaining in the crevices there and being washed into the veins of the next patient. If the hazard of crossinfection had been considered as a cause of the epidemic of jaundice at Cherry Hinton Hospital identification of an infectious cause of jaundice might have been made 20 years earlier than was the case.

The introduction of the sulphonamides in the late 1930s brought familiarity with adverse reactions to all physicians. But these drugs, and later penicillin, streptomycin, and the adrenocorticosteroids, led to such advances in medical treatment that adverse reactions, although recognized, caused no great anxiety. This complacency was shattered in 1961. At a congress of gynaecologists at Kiel on 19-20 October 1961, von Massenbach from Lubeck, Lenz from Hamburg, and Wiedemann from Kiel drew
attention to the large number of children who had recently been born in Germany with hypoplastic or aplastic limb deformities. A month later, at a paediatrics meeting in Dusseldorf, Lenz first suggested that the hypnotic drug thalidomide might be the responsible agent (Taussig, 1962). Thalidomide was marketed in Germany under a number of proprietary names and often was compounded with analgesics for the treatment of pain, cough, and insomnia. Not only were these preparations widely prescribed by doctors but they were purchased over the counter by the public. The profusion of preparations and names made retrospective inquiries about the medicines women had used in pregnancy difficult but Lenz's suspicions were soon confirmed. It is now believed that more than 6000 deformed babies were born in West Germany and some 500 in Britain as a result of the use of thalidomide. Many of these children still live, and remain a grim reminder of a tragedy that shocked the world.

One result of the public outcry after the thalidomide tragedy was that governments in many countries established organizations to ensure that adequate and appropriate toxicity tests were carried out before new drugs were used in human beings. However, as it became recognized that the results of tests in animals cannot be directly extrapolated to man, an urgent need developed for monitoring the use of drugs, especially newly introduced drugs, to identify adverse reactions as soon as possible. In Britain a report on the assessment of drug toxicity which was prepared for the Medical Research Council (1963) not only gave advice about the way in which a system of notification of adverse reactions by doctors using drugs should be established but stated: 'Early recognition alone is not always enough. The purpose of determining the toxic effects (of a drug) in man will usually be to obtain intelligent restriction of its further use. To make such a decision possible it is necessary not only to recognize the toxic effect but also to estimate its incidence and to compare that with the danger of the diseases for which the drug is being used.'

These wise words bear repetition for hasty and unnecessary decisions to ban the use of a drug have been a feature of recent years and are often the result of undue, unbalanced, and sometimes premature publicity by the press which lead to overreaction of the public or politicians. It is, however, now widely recognized that there is a need not only to identify the adverse reactions to a drug but to determine their incidence in relation to the use of the drug. The data available to answer such questions have come from sources each of which has had its limitations.

Medical Literature

Although there was increasing awareness of the serious nature of adverse reactions to certain drugs following the introduction of the sulphonamides in 1935, it was considered sufficient throughout the 1950s to leave the duty of reporting the toxic effects of drugs to individual physicians or pharmacologists who usually reported them in articles or letters in medical journals. It is, however, usually difficult for a physician to establish the relationship between an unexpected toxic effect and a drug. The patient may be receiving many drugs and a single physician may seldom see the same adverse reaction to a drug more than once or twice in his professional life. Nevertheless since 1957 Dr L. Meyler and his colleagues have published a number of most valuable surveys of reports of adverse reactions to drugs occurring in the world literature (Meyler, 1957; Meyler and Herxheimer, 1968). These surveys are of great value as works of reference, for the reader can rapidly ascertain what reactions have been reported with any given drug. However, they seldom provide reliable information about the incidence of reactions and indeed may give a false impression of the incidence, for many reports may appear about certain reactions of special interest to physicians or pharmacologists.

The Registries Reports of Adverse Reactions

In 1951 two American haematologists, Wintrobe and Stur-geon, found by accident that each had seen two or three cases of aplastic anaemia in patients who had been treated with chloramphenicol. It was Dr Wintrobe's idea to establish a Registry of Blood Dyscrasias and between 1953 and 1962 reports of 1195 patients with blood dyscrasias suspected as due to drugs were received. The number of reports received was small: this was due to poor publicity, the novelty of the scheme, and the dauntingly detailed form which had to be filled in. There was also anxiety, which persists in the United States, that by reporting an illness as due to a drug he has prescribed, a physician might expose himself to a charge of negligence by his patient. The establishment of this registry by Dr Wintrobe marks the beginning of systemic monitoring of adverse reactions to drugs (Erselev and Wintrobe, 1962).

After the thalidomide catastrophe registries of adverse reactions were established in a number of countries. Their main purpose is to provide an early warning that a drug causes an adverse reaction. It is clear, however, that only a small proportion of adverse reactions that occur are reported. In the United Kingdom in 1966 when the relationship
The incidence of adverse reactions to drugs

between the oral contraceptive pill and thrombosis
was of wide public interest, it was found that in a
sample of 53 women who had died of thrombotic
illnesses and who were known by their family
doctor to be on oral contraceptives only eight had
been reported to the Committee on Safety of Drugs
(Inman and Vessey, 1968). It is known, too, that
many deaths of asthmatic patients using isoprenaline
aerosols were never reported, but this was because
the relationship between these aerosols and sudden
death was unsuspected (Inman and Adelstein,
1969).

Yet despite serious defects these registries still
constitute one of our most valuable sources of
information about adverse reactions. The Com-
mite on Safety of Medicines (the name was changed
when the Medicines Act was passed in 1968) receives
about 350 reports a month. It is now possible to
characterize a profile of the reactions of a given drug
or group of drugs. When a new but related drug
is marketed suspicions may be aroused if reports of
an unexpected reaction such as jaundice begin to
arrive (Wade, 1970).

The Committee's policy has always been to ask
for a simple report from doctors and a supply of
prepaid yellow postcards is sent to every medical
practitioner. They are asked to report all unexpected
and all serious adverse reactions to drugs, especially
those suspected to be due to new drugs. It has been
a disappointment that so few reports of adverse
reactions to drugs have come to the Committee from
hospitals. Reporting from a hospital can be greatly
increased if a physician on the staff takes a major
interest in the problem and if junior doctors, nurses,
or pharmacists can be employed as 'drug safety
officers'. In the West Midlands Region a Midlands
Adverse Reactions Study Group has been formed and
reporting from a number of the hospitals has
increased greatly. It may be possible to stimulate the
interest of family doctors working in the community,
and this is desirable because the use of drugs by them
is very different from the use in hospitals.

### Intensive Monitoring

The seriousness of an adverse reaction depends not
only on its nature but also on its frequency in
relation to the use of a drug. In the US Army
Custer (1946) determined the incidence of aplastic
anaemia due to the antimalarial drug mepacrine
with a precision which will seldom be equalled. He
showed that the incidence was 2.84 per 100 000
men taking mepacrine compared with 0.1 per
100 000 men not taking mepacrine. In spite of this
low incidence, Custer's evidence that mepacrine can
cause aplastic anaemia is accepted because he
surveyed millions of troops with excellent medical
records.

Intensive monitoring has been developed in
hospitals. Patients are kept under surveillance, all
drugs administered are recorded, and any event
which might be an adverse reaction to a drug is
noted. In a survey carried out at the Belfast City
Hospital, Hurwitz and Wade (1969) showed that
about 10% of patients in our wards had adverse
reactions to drugs (table I), that these tended to be
commoner in women than men (table II), and were
more frequent in the aged than in the young (table
III). It was interesting to find that patients who on
admission had a history of previous drug reactions
or of allergic illness were at greater risk than those
who had no such history (table IV) and it was
possible to identify certain drugs (table V) which
cauied a high incidence of reactions and certain
combinations of drugs (table VI). Similar findings
have been reported by others.

<table>
<thead>
<tr>
<th>No.</th>
<th>Adverse Reaction to Drugs</th>
<th>Rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients admitted</td>
<td>1268</td>
<td>118</td>
</tr>
<tr>
<td>No. given drugs</td>
<td>1160</td>
<td>118</td>
</tr>
</tbody>
</table>

Table I  Adverse reactions to drugs in hospitals

<table>
<thead>
<tr>
<th>Sex</th>
<th>No. of Patients</th>
<th>Rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Given Drugs</td>
<td>With Reactions</td>
</tr>
<tr>
<td>Males</td>
<td>682</td>
<td>50</td>
</tr>
<tr>
<td>Females</td>
<td>478</td>
<td>68</td>
</tr>
<tr>
<td>Total</td>
<td>1160</td>
<td>118</td>
</tr>
</tbody>
</table>

Table II  Sex and drug reaction

0.01 > p > 0.001

<table>
<thead>
<tr>
<th>Age of Patients (years)</th>
<th>No. Given Drugs</th>
<th>No. with Reactions</th>
<th>Rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10–19</td>
<td>64</td>
<td>2</td>
<td>3.1</td>
</tr>
<tr>
<td>20–29</td>
<td>100</td>
<td>3</td>
<td>3.0</td>
</tr>
<tr>
<td>30–39</td>
<td>122</td>
<td>7</td>
<td>5.7</td>
</tr>
<tr>
<td>40–49</td>
<td>159</td>
<td>12</td>
<td>7.5</td>
</tr>
<tr>
<td>50–59</td>
<td>222</td>
<td>18</td>
<td>8.1</td>
</tr>
<tr>
<td>60–69</td>
<td>252</td>
<td>27</td>
<td>10.7</td>
</tr>
<tr>
<td>70–79</td>
<td>178</td>
<td>38</td>
<td>21.3</td>
</tr>
<tr>
<td>80–89</td>
<td>59</td>
<td>11</td>
<td>18.6</td>
</tr>
<tr>
<td>90–99</td>
<td>4</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>1160</td>
<td>118</td>
<td>10.2</td>
</tr>
</tbody>
</table>

Table III  Age and drug reactions

Rank correlation coefficient = + 0.86
Table IV  Adverse reactions and a history of previous reactions, allergic disease, and jaundice

<table>
<thead>
<tr>
<th>Drug</th>
<th>No. of Patients</th>
<th>Rate (%)</th>
<th>Given Drugs</th>
<th>With Reactions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ampicillin</td>
<td>103</td>
<td>8</td>
<td>7.8</td>
<td></td>
</tr>
<tr>
<td>Other penicillins</td>
<td>167</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Orciprenaline</td>
<td>74</td>
<td>6</td>
<td>8.1</td>
<td></td>
</tr>
<tr>
<td>Choline theophyllinate</td>
<td>100</td>
<td>5</td>
<td>5.0</td>
<td></td>
</tr>
<tr>
<td>Methoxyphenamine</td>
<td>133</td>
<td>1</td>
<td>0.8</td>
<td></td>
</tr>
<tr>
<td>Morphine</td>
<td>90</td>
<td>5</td>
<td>5.6</td>
<td></td>
</tr>
<tr>
<td>Methadone</td>
<td>78</td>
<td>2</td>
<td>2.6</td>
<td></td>
</tr>
<tr>
<td>Opium</td>
<td>57</td>
<td>1</td>
<td>1.8</td>
<td></td>
</tr>
<tr>
<td>Pethidine</td>
<td>200</td>
<td>1</td>
<td>0.5</td>
<td></td>
</tr>
<tr>
<td>Dihydrocodeine</td>
<td>76</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Paracetamol</td>
<td>128</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

Table V  Drugs and adverse reactions

<table>
<thead>
<tr>
<th>Drug</th>
<th>Patients Given</th>
<th>Reactions to Digitalis</th>
<th>Rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Digitalis alone</td>
<td>53</td>
<td>5</td>
<td>9</td>
</tr>
<tr>
<td>Digitalis and frusemide</td>
<td>79</td>
<td>16</td>
<td>20</td>
</tr>
<tr>
<td>Digitalis and hydroflumethazide</td>
<td>23</td>
<td>8</td>
<td>33</td>
</tr>
<tr>
<td>Digitalis, frusemide and hydroflumethazide</td>
<td>18</td>
<td>6</td>
<td>33</td>
</tr>
<tr>
<td>Digitalis and other diuretics</td>
<td>24</td>
<td>4</td>
<td>17</td>
</tr>
<tr>
<td>Digitalis and all diuretics</td>
<td>144</td>
<td>34</td>
<td>24</td>
</tr>
</tbody>
</table>

Table VI  Reactions to digitalis and diuretics

The Therapeutic Audit

The thoughtful doctor will realize that it is not enough to establish and improve the monitoring of adverse reactions to drugs. If sensible decisions are to be made much needs to be known about the use of the drug. How widely is it used? Who prescribes it? Which patients receive it? For what illnesses? What good does it do? What other drugs are available?

Studies on the use of drugs in the community are very much in their infancy. Dr Helen Hood and I had the fortunate opportunity to have access to details of all prescriptions written by family doctors in Northern Ireland from 1966 onwards (Wade, 1970). All the data are recorded on computer tape to allow payments to be made to pharmacists and we have been able to follow the change of prescribing of individual drugs over time. Figure 1 shows the changes in the prescribing of hypnotic drugs between 1966 and 1970 (Wade and Hood, 1972a). The great increase in the prescribing of Mandrax was probably related to intense and skilful advertising of this preparation; its use decreased after 1969 and nitrazepam (Mogodon) is now the group leader. Of special interest, in view of the deaths caused by...
The incidence of adverse reactions to drugs

October 1967 (4024 prescriptions) October 1969 (6101 prescriptions)

October 1971 (8177 prescriptions) October 1973 (9544 prescriptions)

excessive use of isoprenaline aerosols, is a study of the use of bronchodilator aerosols. This shows (fig 2) a considerable increase in the use of chromoglycate (Intal) and salbutamol (Ventolin) while the use of the isoprenaline aerosols decreased (Wade and Hood, 1972b).

It has also been possible to study the geography of drug use. One of our first studies was of the use of insulin and oral hypoglycaemic drugs (Wade, Hadden, and Hood, 1972). The prescribing of insulin was remarkably evenly distributed throughout the province. But there were great variations in the prescribing of oral hypoglycaemic drugs (fig 3).

At the time of the survey (1966) the drugs being mainly used were tolbutamide and chlorpropamide. A detailed survey was made of the use of these drugs in Londonderry and Newry, low- and high-use areas respectively (table VII). It was found that in each city almost exactly the same proportion of persons were receiving oral hypoglycaemic drugs.

The difference in overall use was due to the low daily doses prescribed for each patient in Londonderry and the high doses in Newry. I suspect that the main reason for this difference was that in Londonderry the diabetic clinic had a dietician who gave advice and detailed supervision of patients. In Newry there was no dietician. These observations may be important in relation to the anxiety raised by the University Diabetic Group Project (1970) in the USA. The incidence of cardiovascular complications found in that survey was higher in patients on the oral antidiabetic drugs than in those on insulin. This might be investigated in areas where the use of the oral drugs differs greatly.

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**Fig 2** The prescribing of bronchodilator aerosols in Northern Ireland 1967-1973.

**Fig 3** The prescribing of oral hypoglycaemic drugs in Northern Ireland 1966.

The circles indicate the population in different towns and rural areas. The prescribing density is represented by the shading and hatching.

---

<table>
<thead>
<tr>
<th>Table VII</th>
<th>Prescribing of oral antidiabetic drugs in two towns</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Newry</td>
</tr>
<tr>
<td>Diabetic patients</td>
<td>39</td>
</tr>
<tr>
<td>Diabetics per 1000 patients</td>
<td>1.21</td>
</tr>
<tr>
<td>Units of antidiabetic drugs/1000 patients/month</td>
<td>411</td>
</tr>
<tr>
<td>Units of antidiabetic drugs/month/per doctor</td>
<td>837</td>
</tr>
</tbody>
</table>
Recently it has been possible to carry out a study of the prescribing of the oral hypoglycaemic drugs in Northern Ireland, Norway, and Sweden (Crooks, Elmes, Friebel, Halse et al, 1974). The analysis is not yet complete but it is clear that the differences in prescribing found within Northern Ireland are very much smaller than the differences which exist between these three countries (fig 4). It seems to me that if large differences in the prescribing of drugs are occurring it may be possible not only to learn more about the incidence of adverse reactions to the drug but also to assess what value its use is, by comparing the morbidity, span of life, and mortality of communities in which the drug is used with those where it is not used. It has been particularly interesting to find that in one area of Sweden, the county of Jamtland, it is possible not only to enumerate all the citizens by age and sex but also to identify who in the community gets a particular drug. Cardiac glycosides, antihypertensive drugs, and oral hypoglycaemic drugs are found to be prescribed predominantly to the over 60s (fig 5).

It is likely that studies in other countries would show other important differences in the use of drugs. Some eight years ago a study of the sales of antibiotics in six European countries showed that at that time a quarter of all antibiotics used in Germany was chloramphenicol (table VIII) (Engels and Siderius, 1968).

**Death and Drugs**

It has always seemed to me that the most important adverse reactions of drugs are those which cause or contribute to death. With Dr Tesh and with the cooperation of Professor Curran and his colleagues, I have recently arranged an investigation of the part that drugs may have played in the death of 100 patients coming to necropsy at the Queen

![Diagram showing prescribing of oral antidiabetic drugs in Jamtland, 1970.](http://jcp.bmj.com/)

**Table VIII** Sales of antibiotics in Europe. As percentage of total sales within each country

<table>
<thead>
<tr>
<th></th>
<th>France</th>
<th>UK</th>
<th>West Germany</th>
<th>Belgium</th>
<th>Sweden</th>
<th>Holland</th>
</tr>
</thead>
<tbody>
<tr>
<td>Penicillins</td>
<td>20</td>
<td>35</td>
<td>45</td>
<td>11</td>
<td>55</td>
<td>30</td>
</tr>
<tr>
<td>Ampicillins</td>
<td>1</td>
<td>15</td>
<td>5</td>
<td>13</td>
<td>15</td>
<td>30</td>
</tr>
<tr>
<td>Tetracyclines</td>
<td>40</td>
<td>35</td>
<td>20</td>
<td>20</td>
<td>25</td>
<td>30</td>
</tr>
<tr>
<td>Macrolides</td>
<td>8</td>
<td>5</td>
<td>1</td>
<td>30</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Combined tetracyclines and macrolides</td>
<td>8</td>
<td>2</td>
<td>1</td>
<td>15</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Chloramphenicol</td>
<td>5</td>
<td>1</td>
<td>25</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Others</td>
<td>18</td>
<td>7</td>
<td>3</td>
<td>10</td>
<td>2</td>
<td>6</td>
</tr>
</tbody>
</table>

Fig 5 Prescribing of oral antidiabetic drugs in Jamtland, 1970.

The age and sex of the community is shown in open histograms and superimposed is the hatched and shaded histogram of patients receiving oral antidiabetic drugs.
The incidence of adverse reactions to drugs

Important Factors Contributing to Death  No. of Patients

<table>
<thead>
<tr>
<th>After operation</th>
<th>12</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diagnostic procedures</td>
<td>4</td>
</tr>
<tr>
<td>Drugs</td>
<td>20</td>
</tr>
<tr>
<td>Cessation of therapy</td>
<td>4</td>
</tr>
<tr>
<td>Bed rest</td>
<td>3</td>
</tr>
<tr>
<td>None of above</td>
<td>57</td>
</tr>
</tbody>
</table>

Table IX  Analysis of 100 necropsies

Elizabeth Hospital, Birmingham (table IX). Although the assessment is rather subjective and the analysis incomplete, it would appear that the use of drugs, or the cessation of drug therapy, has played a part in the death of at least a quarter of the patients. To me this suggests that a comprehensive multicentre survey carried out by pathologists might be of considerable value. It might lead to a more cautious use of drugs by physicians and it might identify drug hazards which are not yet recognized.

References


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