The general practitioner

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Summary

Despite the instruction he receives in medical school, the general practitioner prescribes mainly by trade names. He receives much information, for example advertising and visits from representatives, in which drugs are identified by trade names. Of greatest importance, these at present provide the most convenient and satisfactory method of ensuring that the patient receives a product which will be consistently effective.

A recent editorial in the Journal of the Royal College of General Practitioners (1973) reminds us that William Osler observed that one of the differences between man and animals is man's desire to take drugs. With easier access to the General Practitioner and the explosion of scientific advance in the field of clinical pharmacology with its attendant publicity via the mass media, this difference is very much more marked now than it was even in Osler's day. The doctor himself feels an almost innate desire to prescribe, and Parish noted in 1971 that 'It is easier to prescribe than to give advice'.

These two pressures contribute to the fact that between 60% and 70% of consultations in general practice result in a prescription being given.

Having reached the stage where the decision to give a prescription has been made by the General Practitioner, the conscious question of whether this should be written in trade name or approved name form arises only infrequently. We are creatures of habit and a number of factors have been involved in the evolution of the method of writing a prescription for a particular drug.

Undergraduate learning

During our undergraduate training most of us will have been impressed by our teachers to use approved names for drugs at all times, and to eschew brand names at all costs. I wonder what the reaction would have been if in our final MB examination papers we had suggested Imperacn or Terramycin as a suitable antibiotic for the treatment of bronchitis, instead of oxytetracycline.

Postgraduate learning

During vocational training, General Practitioners become more aware of brand names as by this time they find themselves reading the journals as well as the text books, and who can fail to be influenced by the mass of advertising that many journals now necessarily contain. Most General Practitioners read something out of most issues of Update which is The Postgraduate Journal of General Practice. In the 1971 volume of Update Plus, out of 966 pages there were 388 full pages of pharmaceutical advertising and in one monthly issue there were only three double pages with no full page advertising material at all. The brand names of drugs are thrust at us, and the proper names are usually only to be found in small print at the bottom of the page.

Having become principals in General Practice, there is less exposure to formal courses of instruction in therapeutics, and practitioners are more easily influenced (not always adversely) by visits from representatives of pharmaceutical companies whose brand of a particular drug is always better than that of its competitors. Reading the Prescribers' Journal or the Drug and Therapeutics Bulletin may make one at least aware of the approved name of the newer compound being used, but the total exposure to this name is only just above the subliminal level.

Ease of writing

Most brand names of drugs are shorter, more easily associated with the condition or symptom they treat, and more euphonious than the approved name. In a busy surgery, when the doctor is working at full pressure, these can become very important factors. Table 1 is taken from a recent paper on drug usage in general practice (Berkeley and Richardson, 1973). It shows the frequency of prescription of the most commonly used drugs amongst twelve General Practitioners in Scotland. Table 2 shows the preparations most commonly prescribed when a single-handed doctor looked at his prescribing habits over a period of 4 separate months (Parish, 1971). You will notice that in both these
It is sometimes necessary to take into account the cost of drugs to the patient, particularly to those people who may require a number of different compounds. If the doctor is aware that they are not eligible for free prescriptions and that they are financially vulnerable he may feel it desirable to use combined preparations where feasible. In almost every case these are prescribed by the brand name. He may also use combined preparations where he feels the patient’s reliability in taking drugs is suspect.

**Effectiveness**

Our prime consideration in prescribing should be whether or not the drug is going to be effective, reliable and consistent in its action. Most concern in this area is directed at drugs which are to be used over the long term in conditions such as diabetes mellitus, epilepsy, hypertension, arthritis and cardiovascular disease. Many of the newer and effective preparations which have been introduced are, or will be shortly, available from sources different from those with which we have always associated them. What we really want to know is whether the methyl-dopa which we have been prescribing with consistent effectiveness for the last 5 years for our 45-year-old hypertensive coal-miner will continue to give consistent control when it comes from another source. The fact that each tablet will contain 250 mg of the active ingredient is of little significance if the particle size changes, or its rate of absorption is altered because the machine which manufactures the tablet is pressing those particles more firmly together, resulting either in a rise of the patient’s blood pressure to uncontrolled levels, or excessive lowering leading to hypotensive episodes. Oral anti-diabetic agents are even more important in this respect, particularly those which can cause hypoglycaemic coma. We have all felt the effects of ‘the digoxin affair’, and unless action is taken now, a repeat performance in another field must eventually come, perhaps with disastrous results.

The study of general practice prescribing mentioned above (Berkeley and Richardson, 1973) covered 26 randomly selected days over a 1-year period, and the twelve General Practitioners involved prescribed a total of 401 different drugs; the average for each doctor being 116. Wilson (1971) in an analysis of his own practice found that he used 148 different compounds in 1 year and he was confident that he used only a small number of well tried preparations. Patterson (1972) found that he used 403 different preparations when he audited his prescribing habits retrospectively in 4 different months over a 3-year period.

It is, therefore, apparent that the General Practitioner has to carry a lot of prescribing information.
about with him. It is unreasonable to expect him to remember several names for the same drug, let alone the possible differences between varying preparations of the same compound. For the time being, the safest course would seem to be to use the cheapest, most effective brand name until the regulations on the manufacture of drugs are tightened up to include the method of manufacture, particle size, solubility and bio-availability for every drug marketed. We shall then not be confronted with the potential hazard which now faces us every time we write a prescription.

References