## Excipients and additives: hidden hazards in drug products and in product substitution

E. Napke,\* MD, DPH D.G.H. Stevens,† MD, FRCP[C]

The excipients and additives in drug formulations have been described as inert because they do not have an active role in the prevention or treatment of particular ailments. This has led to the misconception among physicians, pharmacists, drug manufacturers and the public that excipients are harmless and unworthy of mention. In fact, pharmacists are allowed substitute drug formulations, without regard to the excipients, as long as they ensure that the active ingredients in the substitute are the same as those in the formulation prescribed. The inappropriateness of the term inert is becoming increasingly apparent as evidence of adverse reactions — some fatal — to excipients mounts. The likelihood that some "active" constituents, particularly erythromycin, have been blamed for such reactions deserves to be investigated. The public deserves to be better protected. For example, the United States has legislation requiring complete labelling of all food, drugs and cosmetics that incorporate

\*Chief, Product Related Disease Division, Bureau of Epidemiology, Health Protection Branch, Department of National Health and Welfare, Ottawa, and consultant, Adverse Drug Reaction Reporting Program of the Ontario Medical Association (OMA)

†Chief of pediatrics, York County Hospital, Newmarket, Ont., and long-standing member of the Drug and Pharmacotherapy Committee and the Adverse Drug Reaction Reporting Program of the OMA

This paper is the responsibility of the authors and should not be interpreted as a statement of policy of the Department of National Health and Welfare.

Reprint requests to: Dr. E. Napke, Product Related Disease Division, Health Protection Branch, Tunney's Pasture, Ottawa, Ont. K1A 0L2 more than one ingredient, no matter how innocuous the constituents are believed to be. In Canada, drug manufacturers are not even required to share this information with physicians or pharmacists when they introduce a new drug or reformulate a product already being marketed, nor are pharmacists required to disclose the contents of formulations that they prepare in the absence of commercially available products.

On dit souvent que l'excipient d'une préparation pharmaceutique est inerte parce qu'il n'exerce aucune action préventive ou curatrice contre une maladie donnée. D'où, parmi les médecins, pharmaciens, industriels pharmaceutiques et dans le grand public, l'illusion que l'excipient est inoffensif et indigne de mention. Le pharmacien a même le droit de substituer une autre préparation à celle que le médecin a prescrite, pourvu que les ingrédients soi-disant actifs soient les mêmes, sans considération l'excipient. Pourtant, de vu la fréquence apparemment en hausse des réactions indésirables, voire mortelles, à des excipients, il faut mettre en doute la justesse de l'adjectif "inerte" qu'on leur applique. Il est probable que certaines de ces réactions ont été à tort imputées à des principes actifs, telle l'érythromycine: la chose demande à être étudiée de plus près. Car la protection du public est en cause. Alors qu'aux Etats-Unis, par exemple, la loi exige l'étiquetage complet des ingrédients de tout aliment, médicament ou produit de beauté qui en compte plus d'un, quelle qu'en soit l'innocuité présumée, ce n'est pas le cas pour les médicaments au Canada: le fabricant n'est pas tenu de passer ces renseignements au médecin ou au pharmacien lorsqu'il lance une nouvelle préparation ou qu'il modifie la formule d'une préparation qui existe

déjà, et le pharmacien n'est pas tenu non plus de dévoiler le contenu des médicaments qu'il prépare secundum artem.

Most pharmaceutical products are a combination of constituents. In addition to the active or therapeutic ingredients, product formulations contain a number of "inert" materials known as additives or excipients. Classified according to the part they play in the finished product, the excipients include diluents, binders, lubricants, disintegrators, colours, flavours, sweetening agents etc.

"Dorland's Illustrated Medical Dictionary", 26th edition, defines an excipient as "any more or less inert substance added to a prescription in order to confer a suitable consistency or form to the drug; a vehicle" and an additive as "a substance . . . preservative, or vitamin, added to another substance to improve its appearance, increase its nutritional value, etc.".2 The inappropriateness of the term inert has become increasingly apparent: excipients may be active ingredients, though not in the pharmacologically accepted sense of components intended to affect the structure or function of the body in a way that contributes to the diagnosis, cure, mitigation or prevention of disease.3

In general the potential for adverse reactions to excipients has not been well recognized, probably because of both lack of knowledge about the excipients in products and, more importantly, the misconception that these substances are harmless and, therefore, unimportant. This misconception has contributed to the general acceptance that drug substitution is safe, practical and economical as long as the active ingredients remain constant.

"Epidemics" of adverse reactions due to toxic "inert" substances or changes in drug formulations were reported as early as the 1930s.<sup>4,5</sup> Although recent advances in biochemistry and in diagnostic procedures have expanded our understanding of the role that various constituents of drug preparations play in both the treatment of disease and the production of complications, adverse reactions to inert substances have not been eliminated.

Practising physicians encounter patients who prefer one product to another having identical active ingredients. Frequently the preference results from the patient's having had an adverse reaction to one of the products. This phenomenon was largely unrecognized until 1971, when Lockey6 reported a case of severe headaches and protracted gastrointestinal disturbance associated with the ingestion of Premarin. At that time Premarin's formulation contained 28 ingredients, of which only one, the conjugated estrogen, was considered active. Included in the formulation was the yellow dye FD&C (Food, Drug, and Cosmetic [Act] — USA) no. 5, commonly known as tartrazine. Lockey confirmed that the patient was reacting solely to the tartrazine and could, by washing the colouring agent from each tablet, continue taking the medication without adverse effects. Had Lockey not tested the patient against all components of the formulation, the adverse effects would almost certainly have been attributed to the active ingredient. More importantly, the patient would have been denied useful treatment.

#### Tincture of orange

Similarly, one of us (D.G.H.S.), in a consulting pediatric practice, found tincture of orange to be hazardous. A patient was referred for advice and help in the management of recurrent upper respiratory tract infections. The infections tended to be refractory because the patient was hypersensitive to penicillin and its derivatives and had also demonstrated gastrointestinal intolerance to other antimicrobial medications. Pneumococcal infection was confirmed, and oral administration of 5 mL of liquid erythromycin — Erythrocin 125, each millilitre of which contains 25 mg of erythromycin — was prescribed. Immediately after ingesting the first dose of the suspension, the patient suffered severe abdominal pain, nausea and vomiting. Because of scepticism about an allergy to erythromycin, Erythrocin 250, each millilitre of which contains 50 mg of erythromycin, was substituted at the same dosage. This formulation produced none of the previous symptoms.

Several months later a local pharmacist disregarded the "No Substitution" order on the prescription for the patient and substituted the offending formulation. The patient's pain, nausea and vomiting recurred. Told of the consequences of the substitution, the pharmacist responded as have many practitioners in the past: "The active ingredient is the same; the colouring and flavouring don't matter."

During an outbreak of pertussis on an Indian reserve, Georgina Island, Ontario, the regional officer of health prescribed Erythrocin 125 as prophylaxis and had complaints of similar gastrointestinal symptoms from several of the children. Of the two children who required hospital admission for pertussis, one also experienced intolerance to the one formulation of erythromycin (Erythrocin 125) but not the other (Erythrocin 250) (D.G.H.S.: unpublished data, 1982).

A review of the records of the same consulting pediatric practice disclosed that during the previous 15 years 16 patients with confirmed hypersensitivity to penicillin had manifested the same type of intolerance to Erythrocin 125. Communication with the manufacturer, Abbott Laboratories, Limited, revealed that the sole difference between the two erythromycin formulations was the colouring and flavouring agent, tincture of orange. This alcohol extraction of orange peel, available commercially from several sources, was present in the offending formulation but not in the other, which is formulated with cherry syrup. There is no tartrazine in either preparation.

Tincture of orange is sometimes used by pharmacists to formulate suspensions for infants when a commercially prepared suspension is not available. For example, a 6-week-old boy with congenital heart disease

and heart failure who had been receiving digoxin and Aldactazide (spironolactone – hydrochlorothiazide) in a suspension formulated by the hospital began vomiting after being given the medication provided by a local pharmacy. The local pharmacist had substituted tincture of orange for cherry syrup. Reversion to the cherry formulation resolved the problem immediately and completely (D.G.H.S.: unpublished data, 1981).

In short, gastrointestinal disturbance attributed to erythromycin and other medications formulated with tincture of orange should be reassessed.

#### Metabisulfite

Two papers presented at the annual meeting of the American Society of Allergy8,9 described adverse reactions to metabisulfite, an antioxidant widely used in the food and drug industry. The first paper described four patients with chronic asthma who suffered acute asthma attacks following the ingestion of certain foods and wines. When challenged with each ingredient separately, the patients reacted only to capsules of sodium metabisulfite. The second paper discussed two cases, one of which was similar to the first four. The second involved a patient who had respiratory arrest following intravenous administration of dexamethasone sodium phosphate (Decadron) combined with aminophylline (on four occasions) and, later, metoclopramide hydrochloride (Maxeran) (on one occasion). Double-blind challenge with 500-mg capsules of sodium metabisulfite, which is present in the dexamethasone and metoclopramide preparations but not in the aminophylline formulation, resulted in identical clinical episodes.

#### Polyethylene glycol

Another excipient to which adverse reactions have been reported is polyethylene glycol, an "inert" ingredient in drugs, shaving lotions, milk shakes etc. Kwee and Dolovich<sup>10</sup> reported their experience with a 36-year-old man who in 6 years had five documented episodes of anaphylaxis characterized by hypo-

tension, loss of consciousness and major generalized seizures. The patient reported a past history of hives following the topical use of shaving lotions and perfumes and remembered having taken a multivitamin tablet before the most recent episode. The patient was challenged with the individual constituents of the tablet, and polyethylene glycol was identified as the offending agent.

#### Benzyl alcohol

In 1982 the director of drugs and biologics of the US Food and Drug Administration notified hospital pharmacists of reported problems with benzyl alcohol,11,12 a preservative found not only in multidose vials of sterile water and sodium chloride for parenteral use but also in many parenteral drug formulations, such as Solu-Cortef (hydrocortisone sodium succinate), morphine, heparin and Valium (diazepam). Deaths had occurred in premature infants weighing less than 1250 g; the infants died after the development of a condition named the gasping syndrome, which was eventually attributed to benzyl alcohol. Benzyl alcohol had also been implicated in exacerbation of asthma following the intravenous injection of Solu-Cortef.13 When preparations containing benzyl alcohol were removed from nurseries for premature infants, reports of the gasping syndrome and related deaths stopped.

Questions subsequently arose concerning complications and complaints associated with the epidural administration of morphine for intractable pain. Studies to investigate whether these problems also stem from adverse reactions to benzyl alcohol are under way in Canada.

#### Propylene glycol

Propylene glycol, a polyalcohol, is widely used as a solvent in cosmetics, lotions, ointments and drugs, including many injectable formulations (e.g., of benzodiazepines, digoxin, dimenhydrinate, pentobarbital, phenobarbital and phenytoin), as well as co-trimoxazole preparations for oral use, erythromycin ethylsuccinate and a number of

multivitamin products. This agent is generally considered to be a stable, pharmacologically inert substance with low systemic toxicity. However, it was reported to be associated with systemic toxic effects in 1970 and, more recently, with a protracted seizure unresponsive to anticonvulsant medication. In the latter case the patient's electroencephalogram, grossly abnormal during the seizure, reverted to normal when the exposure to propylene glycol was stopped.

#### Azo dyes

Although in the United States preparations including tartrazine must, by law, be labelled accordingly, no such requirement exists in Canada. In 1982, 450 Canadian pharmaceuticals contained tartrazine,17 though the number was declining as the recognition of adverse reactions spread. However, other aniline or azo dyes, including amaranth, sunset-yellow, FD&C no. 6 and new coccine, have also been implicated in adverse reactions.18 These dyes are present not only in pharmaceuticals but also in dairy products, juices, candies, foodcolouring kits, cosmetics and toile-

When a generic rifampicin-isoniazid preparation caused a reaction and the patient showed no reaction to either rifampicin or isoniazid administered as separate preparations, the excipients were investigated. The combination product contained only a minute amount of sunset-yellow, 0.76 mg per tablet, but challenge with 1 mg of the dye resulted in identical gastrointestinal tract signs and symptoms, which began 6 hours after ingestion and persisted for 12 hours.18 This azo compound is closely related chemically to tartrazine and has now been substituted for it in many formulations.

#### **Discussion**

The lack of understanding of the importance of excipients<sup>19</sup> is demonstrated with a relatively new antihistamine-decongestant. The 1984 edition of the "Compendium of Pharmaceuticals and Specialties" lists the contents of both the liquid and tablet preparations as two — the

### Now available with a pleasant cherry flavour

#### Septra\* Pediatric Suspension b.i.d.

Each teaspoonful (5 mL) contains 40 mg trimethoprim and 200 mg sulfamethoxazole

■ SEPTRA\* (Trimethoprim + Sulfamethoxazole)

B. Summary

**INDICATIONS:** Indicated for the following infections when caused by susceptible organisms.

caused by susceptible organisms.

UPPER AND LOWER RESPIRATORY TRACT INFECTIONS —
particularly chronic bronchitis and acute and chronic otitis
media.

URINARY TRACT INFECTIONS — acute, recurrent and

GENITAL TRACT INFECTIONS — uncomplicated gonococcal urethritis.
GASTROINTESTINAL TRACT INFECTIONS.

SKIN AND SOFT TISSUE INFECTIONS.

SEPTRA is also indicated in the treatment of infants and children with a diagnosis of *Pneumocystis carinii* pneumonitis, especially if they are immunosuppressed. SEPTRA is not indicated in infections caused by *Pseudomonas, Mycoplasma* or viruses. This drug has not yet been fully evaluated in streptococcal infections.

CONTRAINDICATIONS: Patients with evidence of marked liver parenchymal damage, blood dyscrasias, known hypersensitivity to trimethoprim or sulfonamides, marked renal impairment where repeated serum assays cannot be carried out, premature or newborn babies during the first few weeks of life. For the time being SEPTRA is contraindicated during pregnancy.

ADVERSE REACTIONS: Most frequent: nausea; vomiting; gastric intolerance; and rash. Less frequent: diarrhea; constipation; flatulence; anorexia; pyrosis; gastritis; gastroenteritis; urticaria; headache; and liver changes (abnormal elevations in alkaline phosphatase and serum transaminase).

Occasionally reported: glossitis; oliguria; hematuria; tremor; vertigo; alopecia; and elevated BUN, NPN, and serum creatinine. Hematological changes occurring particularly in the elderly, are mostly transient and reversible (primarily, neutropenia and thrombocytopenia; less frequently, leukopenia, aplastic or hemolytic anemia, agranulocytosis, and bone marrow depression).

PRECAUTIONS: As with other sulfonamide preparations, critical appraisal of benefit versus risk should be made in patients with liver damage, renal damage, urinary obstruction, blood dyscrasias, allergies or bronchial asthma.

The possibility of a superinfection with a non-sensitive organism should be borne in mind.

DOSAGE AND ADMINISTRATION: Adults and children over 12 years. Standard dosage: 2 SEPTRA tablets or 1 SEPTRA DS tablet twice daily. Minimum dosage and dosage for long-term treatment: 1 SEPTRA tablet or 1/2 SEPTRA DS tablet twice daily.

Maximum dosage. Overwhelming infections: 3 SEPTRA tablets or 1½ SEPTRA DS tablets twice daily. Uncomplicated gonorrhea: 2 SEPTRA tablets or 1 SEPTRA DS tablet four times daily for 2 days.

Pneumocystis carinii pneumonitis: 20 mg/kg/day trimethoprim and 100 mg/kg/day sulfamethoxazole in four divided doses for 14 days.

Children 12 years and under. t

Young children should receive a dose according to biological age:

Children under 2 years: 2.5 mL of suspension twice daily. Children 2 to 5 years: 2.5-5 mL of suspension twice daily. Children 6 to 12 years: 5-10 mL of suspension twice daily.

the children this corresponds to an approximate dose of 6 mg trimethoprim/kg body weight/day, plus 30 mg sulfamethoxazole/kg body weight/day, divided into two equal doses.

DOSAGE FORMS:

SEPTRA DS TABLETS, each containing 160 mg trimethoprim and 800 mg sulfamethoxazole, and coded SEPTRA DS 02C. Bottles of 50 and 250. SEPTRA TABLETS, each containing 80 mg trimethoprim and

SEPTRA TABLETS, each containing 80 mg trimethoprim and 400 mg sulfamethoxazole, and coded SEPTRA Y2B. Bottles of 100 and 500.

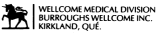
SEPTRA PEDIATRIC TABLETS, each containing 20 mg trimethoprim and 100 mg sulfamethoxazole, and coded WELLCOME H4B. Bottles of 100.

SEPTRA PEDIATRIC SUSPENSION, each teaspoonful (5 mL) containing 40 mg trimethoprim and 200 mg sulfamethoxazole. Bottles of 100 and 400 mL. SEPTRA FOR INFUSION, each ampoule (5 mL) containing

SEPTRA FOR INFUSION, each ampoule (5 mL) containin 80 mg trimethoprim and 400 mg sulfamethoxazole.

#### Product Monograph available on request.

References: 1. Krause PJ, Owens NJ, Nightingale CH, et al: Penetration of amoxicillin, cefaclor, erythromycinsulfisoxazole, and trimethoprim-sulfamethoxazole into the middle ear fluid of patients with chronic serous otitis media. *J Infect Dis* 1982, 145:815-821. 2. Zack BG: Otitis media: Diagnosis and treatment. *Drug Therapy* 1982; 7 (February): 209-217. 3. Data on file, Burroughs Wellcome Inc.



\*Trade Mark

W-2026

PAAB CCPP antihistamine and pseudoephedrine. In fact, the syrup contains, in addition to the two active ingredients, 11 excipients, 6 of which have been implicated in adverse reactions. The tablet has four excipients, two of which are known to cause reactions (D.G.H.S.: unpublished data, 1983–84).

The hazards of excipients are not limited to new drug products and drug substitutions: they extend to undisclosed changes in drug formulations. Within the past 15 months two preparations — an aqueous aminophylline formulation and an anticonvulsant suspension — widely used in the practice of pediatrics were reformulated and caused adverse reactions that were reported to the Adverse Drug Reaction Program of the Ontario Medical Association. In neither case were the pertinent professions, medicine and pharmacy, notified. In fact, it was difficult to obtain specific information about the formulation changes.

Physically any drug product is the sum of its individual components. Pharmacologically, however, a drug does not include its excipients. The fundamental dictum of therapeutics is that any substance producing an effect has the potential for side effects; this dictum must be applied to the so-called inert excipients. The increasing numbers of adverse side effects should lead to purer drug formulations in the future.

Although Lockey21 had reported "allergic" reactions to tartrazine in 1959, it was not until 1980 that labelling regulations in the United States required documentation of the presence of this dye; such regulations are not in effect in Canada. Yet the known adverse reactions to this agent and to the other excipients we have cited — a list that is certainly not all-inclusive — clearly indicate that the time has come for drug manufacturers to disclose to physicians and pharmacists all ingredients of all formulations, both new and revamped. The present information vacuum has contributed to unnecessary illness and even fatalities and is no longer acceptable.

#### References

- Dorland's Illustrated Medical Dictionary, 26th ed, Saunders, Philadelphia, 1981: 473
- 2. Ibid: 30
- Ansel HC: Introduction to Pharmaceutical Dosage Forms, 3rd ed, Lea & Febiger, Philadelphia, 1981: 104

 Elixir of sulfanilamide-Massengill: chemical, pharmacologic, pathologic and necropsy reports; preliminary toxicity reports on diethylene glycol and sulfanilamide; special article from American Medical Association Chemical Laboratory. JAMA 1937; 109: 1531-1539

- Geiling EMK, Cannon PR: Pathologic effects of elixir sulfanilamide (diethylene glycol) poisoning; clinical and experimental correlations — final report. JAMA 1938; 111: 919-926
- Lockey SD Sr: Reactions to hidden agents in foods, beverages and drugs. Ann Allergy 1971; 29: 461-466
- Napke E: The Canadian Drug Adverse Reaction Reporting Program. Drug Inf J 1975; Sept: 224-231
- Stevenson DD, Simon RA: Sensitivity to ingested metabisulfites in asthmatic subjects. J Allergy Clin Immunol 1981; 68: 26-32
- Allen DH, Collett P: Life threatening asthmatic reactions to food and drug preservative, sodium metabisulfite. Ibid: 67-71
- Kwee YN, Dolovich J: Anaphylaxis to polyethylene glycol (PEG) in a multivitamin tablet: part 2. J Allergy Clin Immunol 1982; 69: 138
- Gershanik J, Boecler B, Ensley H et al: The gasping syndrome and benzyl alcohol poisoning. N Engl J Med 1982; 307: 1384-1388
- Brown WJ, Buist NRM, Cory Gipson HT et al: Fatal benzyl alcohol poisoning in a neonatal intensive care unit [C]. Lancet 1982; 1: 1250
- Grant JA, Bilodeau PA, Guernsey BG et al: Unsuspected benzyl alcohol hypersensitivity [C]. N Engl J Med 1982; 306: 108
- Seidenfeld MA, Hanzlik PJ: General properties, actions and toxicity of propylene glycol. J Pharmacol Exp Ther 1932; 44: 109-121
- Martin G, Finberg L: Propylene glycol: a potentially toxic vehicle in liquid dosage form. J Pediatr 1970; 77: 877-878
- Arulanantham K, Genel M: Central nervous system toxicity associated with ingestion of propylene glycol. *J Pediatr* 1978; 93: 515-516
- MacCara ME: Tartrazine: a potentially hazardous dye in Canadian drugs. Can Med Assoc J 1982; 126: 910-914
- Jenkins P, Michelson R, Emerson PA: Adverse drug reaction to sunset-yellow in rifampicin/isoniazid tablet [C]. Lancet 1982; 2: 385
- Napke E: Adverse Reactions: Some Pitfalls and Postulated Side Effects of Drugs, Annual 7, Excerpta Medica, Amsterdam, 1983: xv-xxvi
- Compendium of Pharmaceuticals and Specialties, 19th ed, Can Pharm Assoc, Ottawa, 1984
- 21. Lockey SD Sr: Allergic reactions due to FD&C #5, tartrazine. Ann Allergy 1959; 17: 719-721

Doctors...businessmen...academic groups... trekkers...wildlife and archeology buffs...

# The group that (tours together, saves together.

You'll get more out of your Indian vacation when you travel with people who want to see and do the same things as you.

You may also qualify for greatly reduced prices when you travel as a group with a common interest.

Your group can join one of the many exciting India tour packages, or we can customize a tour especially for you.

Plan now by contacting your travel agent, or your nearest Air-India general sales agent.



Montréal: 842-1805 or toll-free 1(800) 361-2813

**Toronto**: 865-1033 or toll-free: 1(800) 268-9582

Vancouver: 879-0271 or toll-free 1(800) 663-3433