**New Drugs**

**New drugs in respiratory disorders: II**

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In this second article I will discuss the newer antimicrobial drugs and compare them with established antimicrobial drugs.

**New antimicrobial agents**

**TRIMETHOPRIM**

Trimethoprim, a selective antagonist of bacterial dihydrofolate reductase, is combined with sulphamethoxazole in co-trimoxazole, which is of proved value in lower respiratory tract infections, in particular the mixed infection of *Haemophilus influenzae* and *Streptococcus pneumoniae* that occurs so commonly in patients with chronic bronchitis and emphysema. A comparison of trimethoprim alone with co-trimoxazole showed no difference in outcome in adults with bronchitis, chronic bronchitis, or pneumonia, but side effects (rashes, nausea, etc) were more common with co-trimoxazole. Trimethoprim is marginally cheaper than co-trimoxazole and should therefore be considered when treatment with co-trimoxazole is indicated. Co-trimoxazole in high doses remains the treatment of choice, however, in infection with *Pneumocystis carinii* in immune compromised patients.

**AMOXICILLIN WITH CLAVULANIC ACID (AUGMENTIN)**

β-lactamases, which open the β-lactam ring of penicillins and cephalosporins, are a major source of bacterial resistance to these antibiotics. The combination of amoxycillin with a β-lactamase inhibitor, clavulanic acid, as in Augmentin (250 mg amoxycillin +125 mg clavulanic acid per tablet) is effective against many such respiratory pathogens, including most ampicillin resistant strains of *H influenzae*. Augmentin was therefore effective in 97% of patients with pneumonia, bronchiectasis, and bronchitis, but pathogens were identified in only 25% of these patients, even when transtracheal aspiration was used, and β-lactamase producing organisms were not important. Minor gastrointestinal disturbances or rashes occurred in 30% of patients. Only 6%, of strains of *H influenzae* currently isolated in the United Kingdom, however, are ampicillin resistant. Thus, first line treatment with Augmentin for an acute exacerbation of chronic bronchitis scarcely seems justifiable in preference to amoxycillin or ampicillin alone as Augmentin costs half as much again as the same dose of amoxycillin alone.

**CEPHALOSPORINS**

Of the 11 cephalosporins or cephamycins listed in the 1982 *British National Formulary*, only cephalaxin, cephadine, and cefaclor are active when given by mouth, whereas the others (cefotaxime, cefoxitin, cefuroxime, cefsludin, cefaloridine, cephalothin, cephamandole, and cephalolin) must be given parenterally. What is the role for this plethora of cephalosporin antibiotic drugs in today’s respiratory medicine? Unfortunately, the results that are available from trials rarely give a definitive answer, but the evidence will be briefly discussed for four major areas of bacterial infection of the lower respiratory tract—pneumonia acquired outside hospital; acute exacerbation of chronic bronchitis and emphysema; pneumonia in a previously compromised host; and chronic infections (cystic fibrosis, bronchiectasis, etc). The more general use of cephalosporins is considered elsewhere.

The new cephalosporins have few side effects. Pain after intramuscular injection is rare but may be reduced by adding lignocaine. The drugs are mostly excreted unchanged by both glomerular filtration and tubular secretion, so that renal failure may lead to toxic concentrations. Probenecid will prolong the half life. Hypersensitivity rashes may occur in 2-5% of all recipients and cross reactivity may occur in 5-16% of patients known to be sensitive to penicillin. The combination of a cephalosporin—for instance, cephaloridine—with an aminoglycoside may be nephrotoxic.

**Pneumonia acquired outside hospital**

Pneumonia acquired outside hospital is still most often due to *Strep pneumoniae*, which is nearly always as sensitive to penicillin as to any of the current cephalosporins. There is thus no justification for using a cephalosporin in preference to penicillin in treating proved *Strep pneumoniae* pneumonia except when the particular organism is known to be resistant to penicillin.

**Chronic bronchitis and emphysema**

In chronic bronchitis and emphysema *Strep pneumoniae* and *H influenzae* are the commonest pathogens in the sputum. Of the oral cephalosporins, only cefaclor is active against both. A 10 day course of cefaclor (250 mg three times a day), however, is three times more expensive than an equivalent course of ampicillin. There was no difference in clinical outcome in a comparative trial of these two drugs in 50 patients aged 65-85, but the causative pathogens were not determined. Furthermore, although 250 mg cefaclor three times daily “cured” 93% of patients with pneumonia or acute exacerbations of chronic bronchitis (97% with 500 mg twice daily), the results of another trial showed 250 mg of ampicillin three times a day for seven days to be as effective in an acute exacerbation of chronic...
bronchitis as either 500 mg of ampicillin or amoxycillin in either 250 or 500 mg doses, all given four times daily. Therefore in treating purulent exacerbations of chronic bronchitis and emphysema there seems little to justify oral cefaclor in preference to oral ampicillin (or amoxycillin), and clearly cephalaxin and cephadrine (the other drugs active when given by mouth) are not indicated as they are not active against both Strep pneumoniae and H influenzae. If, however, the patient has a proved infection with an ampicillin resistant strain of H influenzae cefuroxime, cefotaxime, or ceftazidime are all active, at least in the laboratory. None the less, superior clinical efficacy, as compared with co-trimoxazole, or possibly ampicillin, remains to be established in a properly controlled clinical trial where the invading pathogens are identified before treatment. Although purulent exacerbations in patients with relatively mild chronic bronchitis may not be appreciably influenced by antibiotic treatment (tetracycline), severe exacerbations with CO₂ retention (type 11 respiratory failure) still carry a mortality rate of 9-24%. Bacterial pneumonia in the compromised host Whether bacterial pneumonia in the compromised host results from a congenital abnormality—for example, hypogammaglobulinaemia—or is today more commonly induced therapeutically (as in organ transplantation, treatment of autoimmune disease, or malignancy) it is often due to Strep pneumoniae infection but may also arise from H influenzae, Klebsiella spp, Pseudomonas aeruginosa, or anaerobic organisms. Klebsiella pneumoniae carries a mortality of over 40%, and although gentamicin is often an agent of first choice, in Britain, where chloramphenicol is rarely used in treating other conditions, this drug in combination with streptomycin and tetracycline has often been effective. Cefotaxime is very active against Klebsiella spp in vitro, as is cephamandole and other parenteral cephalosporins. Cefotaxime and ceftazidime have the greatest in vitro activity against Klebsiella spp. With cefotaxime this has been confirmed clinically, klebsiellae being eliminated in 91% of 189 adult patients, but the cases of pneumonia were not separately identified. Cefotaxime may be the only drug that is active against gentamicin resistant strains of Klebsiella spp. Mycobacterium tuberculosis, Legionella pneumophila, viral or fungal agents, and Pneumocystis carinii are other infective agents recognised as causing pneumonia in the compromised patient, but none will respond to any of the cephalosporin antibiotics. Antiviral and antifungal chemotherapy are considered elsewhere in this series.

Chronic infections

Chronic infections include bronchiectasis, where again Strep pneumoniae and H influenzae are common infective agents, with, more rarely, pseudomonas or anaerobes, or both, and cystic fibrosis, where in childhood staphylococcal infections and in the adult pseudomonas infections are common, particularly with the miliary strain of Ps aeruginosa, which carries a particularly grave prognosis. Carbencillin and gentamycin by aerosol over many months can slow the progress of cystic fibrosis in adults with pseudomonas. Cef sulodin and ceftazidime (not yet available in the United Kingdom) also seem very promising as they may be more active than any other cephalosporin against Ps aeruginosa. Cefazidime is also active against pseudomonas strains resistant to carbenicillin. The cephalospors are the most expensive of any group of antibiotics in use. Good patient care may be maintained despite a voluntary restriction of prescribing cephalospors in hospital. Treatment with 2 g of cefotaxime a day for seven days costs £90-120 as compared with £1-2 for 250 mg of ampicillin three times daily for seven days. Thus if all new episodes of an exacerbation of chronic bronchitis in the United Kingdom were treated with cefotaxime, which is certainly not recommended, the bill would be around £100-200m a year for this alone compared with £1-2m a year if ampicillin were used; these figures were calculated from estimates of the probable numbers of such acute exacerbations each year in the United Kingdom. Ampicillin, amoxycillin, and co-trimoxazole (or possibly trimethoprim alone) are rational choices for first line treatment of this very common condition, so allowing excellent patient care to be combined with sensible thrift.

Erythromycin

Erythromycin may have a new role as possibly the most effective drug for treating legionnaires’ disease, in addition to its traditional role as an alternative to penicillin for use in penicillin sensitive patients and also for treating chlamydial pneumonia in infancy. Although evidence from a controlled trial is not available, in infection with L pneumophila (legionnaires’ disease), pneumonia, and baceraemia, Pontiac fever, and febrile illness without pneumonia) erythromycin, possibly at first given intravenously in the sick patient followed by oral maintenance treatment in a dose of 2-4 g/day for three weeks, is recommended. Relapse has been noted with shorter courses of treatment. Rifampicin is also effective in vitro but should not be used as the sole antimicrobial drug. L pneumophila produces a β-lactamase, which may explain the lack of effect of penicillin and cephalosporins. Erythromycin estolate is well absorbed by mouth but may cause liver damage, whereas erythromycin stearate and propionate both yield lower serum concentrations, but this does not seem to affect the results of treatment, and neither of these preparations is hepatotoxic. Intravenous erythromycin is given as the lactobionate, 0.5 g every six hours for adults. In severe infections this may be increased to 6-8 g daily, but intramuscular injection is not advisable.

Anaerobic pulmonary infections

METRONIDAZOLE

Metronidazole is active against nearly all Gram negative anaerobic bacteria—that is, the anaerobes that contribute to the mixed flora of lung abscess and aspiration pneumonia and most strains of Bacteroides fragilis. Metronidazole by mouth (500 mg four times daily in an adult) yields adequate serum concentrations at much less cost than the very expensive intravenous form. Serious neurological side effects (vertigo, ataxia, confusion, and fits) or reversible peripheral neuropathy are rare at these doses, except with prolonged courses or in patients with liver disease. Penicillin, ampicillin, and amoxycillin are all also active against many anaerobes (except for β-lactamase producing Bacteroides spp) as are most cephalosporins.

CLINDAMYCIN

Clindamycin is active against many anaerobes, including penicillin resistant strains, and is thus superior to penicillin in treating lung abscess, being actively transported to the infection site in leucocytes. It causes diarrhoea in 10-20% of patients which usually improves when the drug is stopped, but may cause Clostridium difficile induced colitis in 0-1-10% of patients.

Influenza

AMANTADINE

Amantadine, which has an established use in the management of Parkinsonism, has recently been shown to be effective in
Chemotherapy for tuberculosis

RIFAMPICIN

Rifampicin has radically changed the drug treatment of tuberculosis. The former 18-24 month regimes using streptomycin, isoniazid, and aminosalicylic acid (now designated classic chemotherapy) have been replaced in economically developed countries by standard chemotherapy—rifampicin (600 mg if the patient is over 50 kg, 450 mg (8-12 mg/kg) for those under 50 kg), with isoniazid 300 mg (5 mg/kg) once daily in a combination tablet of both drugs taken one hour before breakfast on an empty stomach for nine months. In addition, ethambutol (15 mg/kg) is included for the first two months. This standard chemotherapy has been shown to be effective, with insignificant failure rates, in controlled trials in Britain and elsewhere.

Rifampicin is fully absorbed, with therapeutic concentrations maintained for 12 hours due to an enterohepatic recirculation after an oral dose. Most of the drug is excreted in the bile and with daily treatment minor skin reactions, gastric upsets, and a rise in liver transaminases are all recognised but uncommon side effects, particularly during the early weeks of treatment, but these resolve despite continuing treatment. Thrombocytopenia, probably due to a circulating antibody, is a rare but important side effect as it is a contraindication to further use of the drug, but the platelet count rises when treatment is stopped. Clinical hepatitis is uncommon with today’s standard chemotherapy, but pre-existing severe liver disease remains as a possible contraindication to treatment with rifampicin.

Rifampicin induces liver enzymes, so increasing the elimination of common drugs such as warfarin, corticosteroids, and sulphonyleureas, whose dosage will thus need to be increased during treatment with rifampicin. Oral contraceptives are also metabolised more rapidly, and an alternative contraceptive method should be used during treatment with rifampicin. Intermittent rifampicin treatment, which is not included in the standard regimen, can induce a flu-like syndrome, with fever and chills, after three months’ treatment because of a circulating antibody. If the drug is restarted after a long interval the dose should be built up gradually. Rifampicin is contraindicated in pregnancy. An intravenous rifampicin preparation is available for those too ill to take tablets.

ETHAMBUTOL

Ethambutol is included in the standard regimen to prevent development of rifampicin resistance if the patient’s organisms are initially resistant to isoniazid. This occurred in under 2% of all cases in the United Kingdom in 1978-9 but in 8% of cases originating in the Indian subcontinent. It is only a weak antituberculosis drug, 80% of the drug being absorbed after an oral dose, with a plasma half life of some eight hours. Optic neuritis, indicated by blurred vision, loss of visual acuity, and inability to see colours, is a recognised hazard in doses over 25 mg/kg but is rare if the dose is not above 15 mg/kg, as in the standard chemotherapy regimen. Vision usually returns after stopping the drug, but patients should be warned to report any visual disturbance during treatment with ethambutol.

PYRAZINAMIDE

Pyrazinamide, introduced in 1954, was previously considered too hepatotoxic for wide use. Interest has been reawakened for use in lower doses as a valuable part in short courses of antituberculosis regimens, possibly by killing mycobacteria within macrophages at a low intracellular pH and so reducing the relapse rate in six month regimens. Thus an oral daily dose of 1-5 g (for those weighing under 50 kg) to 2-5 g (weight over 75 kg) in two combinations with rifampicin and isoniazid, showed no excess hepatotoxicity (as detected by liver function tests, gastrointestinal symptoms, or jaundice) that could be attributed to the pyrazinamide alone. Pyrazinamide has a role in the management of tuberculous meningitis as it rapidly diffuses into the cerebrospinal fluid. Rifampicin, isoniazid (10-12 mg/kg intramuscularly or intravenously), pyrazinamide 30 mg/kg to a maximum of 2-5 g daily, and streptomycin 1-0 g daily (or 0.75 g daily if the patient is over 35) are combined, along with pyridoxine 50 mg daily by mouth to prevent the risk of peripheral neuropathy from the high dose of isoniazid. Corticosteroids are also used to prevent formation of adhesions and hydrocephalus in treating this life threatening condition.

SHORT COURSE CHEMOTHERAPY

In patients whose sputum is positive on smear there are now several effective regimens lasting for six months, based on rifampicin, isoniazid, and pyrazinamide for the first two months (some including thrice weekly streptomycin in addition) followed by isoniazid and rifampicin for the remaining four months. Even shorter regimens are being evolved.

COMPLIANCE WITH TREATMENT

Compliance with treatment is now the main problem remaining in managing tuberculosis. Both the patients and all their doctors must realise that cure can be guaranteed only if all doses are taken for the whole course of treatment. This advice must be repeated frequently. The reddish colour of the urine after taking rifampicin can be intensified by extraction with butanol, so proving that the drug has been taken within 12-24 hours. Supervised chemotherapy may be needed in feckless patients, which includes alcoholics and drug addicts. Partially treated patients may excrete drug resistant organisms and so be a menace to themselves and to others.

CYTOXIC CHEMOTHERAPY

Chemotherapy in treating lung cancer has attracted much recent interest. Cytotoxic agents in combination are now being widely used in small cell lung cancer, where they may offer a substantial improvement in survival over that to be expected without treatment, although cure or prolonged survival is possible in very few. These drugs are thus used only in inoperable cases, which in small cell cancer unfortunately includes most cases, as the disease is usually disseminated at the time of diagnosis. The most active agents include cyclophosphamide, doxorubicin, vincristine, methotrexate, lomustine, and etoposide. Although all are active when used alone, combining these drugs in various regimens gives the best results. The drugs are usually given parenterally either together or only a day or two apart, at intervals of three to six weeks. All act by interfering with cell division, but of course they not only attack malignant cells but also depress the bone marrow, hair growth, and rapidly dividing cells in the gut. Side effects therefore occur frequently and may threaten life. Thrombocytopenia, with resultant spontaneous bleeding, is an uncommon but serious side effect, but nausea, vomiting, and alopecia are so common as to
be expected in all patients. The peripheral blood count should thus be checked before the first dose of the drugs and a week to 10 days after these bolus doses (or three to four weeks after 5-lomustine), when the white cell count will reach its nadir, but it recovers thereafter. This pancytopenia is also associated with varying degrees of immunosuppression, so that infections may also complicate treatment.23 None the less, many patients with small cell lung cancer can have a pain-free life of reasonable quality considerably extended by these regimens at the cost of three or four days of anorexia and nausea every three or more weeks. Loss of hair is disguised by a wig. Most physicians think that the patient should know why such treatment is being recommended so that he may make an informed decision as to whether to accept this advice. Further information on new cytotoxic drugs is given elsewhere in this series.

CYCLOPHOSPHAMIDE

Cyclophosphamide, the well known alkylating agent, damages deoxyribonucleic acid (DNA) and so interferes with cell replication after hepatic metabolism that renders it active. Nearly all effective regimens for small cell cancer include cyclophosphamide, and in addition to the expected alopecia and pancytopenia (seen above) this drug may also cause a distressing haemorrhagic cystitis from metabolites that accumulate in the bladder. This is common with regimens using an intravenous bolus of 2 g or more of cyclophosphamide, but symptoms of cystitis may develop in any patient with a disorder of the lower urinary tract. A high intake of fluid can minimise this but is not always effective. A new drug, mesna (mercaptopropane sulphonate), prevents cystitis by binding to the metabolites of cyclophosphamide in the urine, and so enables high doses (10-15 g) of cyclophosphamide to be given, without causing cystitis, in combination with autologous marrow transplantation.31 Mesna is not yet available on prescription in the United Kingdom.

DOXORUBICIN

Doxorubicin also interferes with DNA and RNA (ribonucleic acid) replication and produces complete alopecia as well as pancytopenia. It also increases sensitivity to therapeutic irradiation, given either before or afterwards. It is given intravenously in a fast running intravenous drip, as disastrous local reactions may follow extravasation into tissues. Prolonged administration may cause cardiomyopathy, but arrhythmias are common at even lower dosage. The total dose should not exceed 550 mg/m2 and should be less in patients with previous myocardial disease.21

VINCRISTINE

Vincristine is a plant alkaloid that is a useful adjunct to other chemotherapeutic regimens as it causes minimal marrow depression. A major side effect is neuropathy (sensory, motor, or autonomic), which is seen after long term administration. Extravasation into tissues from an intravenous infusion can cause local cellulitis.

METHOTREXATE

Methotrexate is an antimetabolite that inhibits dehydrofolate reductase and thus DNA synthesis. It has recently been given in high doses (20-100 mg/m2) intravenously followed after 12-24 hours by rescue with folic acid, which overcomes the enzymatic inhibition. Treatment, although effective, is potentially dangerous in patients with large pleural or peritoneal effusions, as the drug may later be released from these sites with subse-

quent profound marrow depression. These high dose regimens, however, are thought to penetrate the central nervous system to treat micrometastases at that site.32

LOMUSTINE

Lomustine is a nitrosourea and alkylating agent (like cyclophosphamide) that can also cross the blood brain barrier and may be active in disease disseminated into the nervous system. Delayed suppression of the bone marrow may occur.

ETOPOSIDE

Etoposide is one of the most effective new agents against small cell lung cancer.4 It is usually given over three to four days every three to four weeks either by mouth or intravenously, and the major side effects are alopecia, marrow depression, and gastrointestinal disturbances.

As with antituberculosis chemotherapy, such chemotherapy will usually be started by the respiratory physician or oncologist who has established the diagnosis of small cell cancer. As treatment may be prolonged, however, other doctors concerned with the patient’s care during this time should have some knowledge of the potential side effects of these chemotherapeutic regimens for these may well affect at present to the patient’s family doctor.

References

MATERNA NON MEDICA

Amateur naturalists and their occupation

A serious interest in natural history is unusual even among countrymen, but its incidence has a noteworthy association with the professions of medicine and law. Is the Victorian amateur naturalist, whose standards have seldom been equalled, dominated by Anglican clergy and the medical profession?

No doubt because of the decreasing likelihood of discovering new species and the inhibiting effect of more professional naturalists, there was, at least in this county, a marked decline in the excellence of amateur naturalists during the first half of this century; but in the second half there has been a revival, stimulated partly by television, the profusion of superbly illustrated books, and the remarkable sophistication of the camera, leading to widespread rejuvenation of natural history societies by enabling lecturers to show slides which so beautifully reveal the intimate details of plant and animal life.

What, now, are the occupations of amateur naturalists? Certainly they come from more diverse backgrounds, but the medical profession remains very much to the fore. Now that the profession is more specialised it might even be possible to correlate a love of natural history with different disciplines. My impression is that naturalists are to be found, at present, mainly among general practitioners and physicians, but seldom surgeons.

But how about the current neglect by the clergy? Not just, I think, that they are dispirited by Darwinism. And why the lack of interest by lawyers—do they wrongly suppose that nature is chaotic?

8 8 MILES, retired physician, Hereford.

The age of the pak pai?

In Hong Kong, with its overcrowded and inadequate roads, the “pak pai” is the answer to many businessmen’s perennial problem of how to travel and work efficiently each day. The pak pai (the minicab or licensed taxi) is a hired vehicle whose legality is doubtful (due to lack of insurance cover for the occupants), whose mechanical condition would rarely pass an MOT inspection, and is driven by a young Chinese. A cab as ecumenically eccentric as the cabby, leading to widespread rejuvenation of natural history societies by enabling lecturers to show slides which so beautifully reveal the intimate details of plant and animal life.

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