Prescribing in Pregnancy

Treatment of rheumatic diseases

M A BYRON

Musculoskeletal disorders are common, and conditions such as low back pain and carpal tunnel syndrome may require treatment during pregnancy. Rheumatic conditions are more common in women, with the peak prevalence of rheumatoid arthritis and systemic lupus erythematosus occurring in women of childbearing age. Anti-rheumatic drugs are, therefore, often required for women of childbearing age.1

Table I lists the groups of drugs prescribed for rheumatic conditions. Analgesics and non-steroidal anti-inflammatory drugs are most commonly prescribed, some of which—for example, ibuprofen—are now available without prescription. Several reviews discuss the effects of these drugs during pregnancy and lactation.24

Table I—Antirheumatic drugs

<table>
<thead>
<tr>
<th>Drug group</th>
<th>Condition treated</th>
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<tbody>
<tr>
<td>Analgesics</td>
<td>Soft tissue lesions</td>
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<tr>
<td>Non-steroidal anti-inflammatory drugs</td>
<td>Inflammatory arthritides</td>
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<tr>
<td>Antimalarial drugs</td>
<td>Osteoarthritis</td>
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<tr>
<td>Sulphasalazine</td>
<td>Systemic lupus erythematosus</td>
</tr>
<tr>
<td>Gold salts</td>
<td>Rheumatoid arthritis</td>
</tr>
<tr>
<td>Penicillamine</td>
<td>Azathioprine</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>Rheumatoid arthritis (infrequently)</td>
</tr>
<tr>
<td>Cytotoxic agents</td>
<td>Other connective tissue diseases</td>
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Non-steroidal anti-inflammatory drugs

TERATOGENICITY

Studies in animals have linked a variety of skeletal and cranio-vertebral abnormalities with ingestion of large doses of salicylates during pregnancy. In humans several retrospective surveys have shown that significantly more mothers of malformed infants took salicylates regularly during pregnancy than mothers of normal infants.3 In these studies, however, factors such as the reason for taking salicylates, general health and nutrition of the mothers, and incidence of defects in the families of the malformed children were not always investigated. Three prospective studies have not shown a teratogenic effect of aspirin.6 The largest study, the Perinatal Collaborative Project of the United States of America, found that malformation rates were similar in the children of 35 418 women not exposed to aspirin, 9736 with intermediate exposure, and 5128 women heavily exposed during the first four months of pregnancy.8

Even in women identified as habitual aspirin users the prevalence of congenital malformation was not significantly increased.3 Overall, therefore, the evidence suggests that salicylates used in recommended doses are unlikely to produce fetal malformations.

Indomethacin is associated with teratogenicity in animals, but the link with human malformation is tenuous.26 Sulindac, diflunisal, and piroxicam have not been found to be teratogenic in animals, whereas azapropazone and diclofenac have, though at doses greater than those used in humans.20 No information is available for the fenamates and tometin. Studies in animals have found no evidence of teratogenicity with the commonly prescribed propionic acid derivatives such as ketoprofen, ibuprofen, flurbiprofen, and naproxen.9

EFFECTS ON FETAL GROWTH

A survey from Sydney showed that long term ingestion of aspirin was associated with an increased incidence of stillbirth and reduced birth weight compared with that of controls.7 Most of the aspirin preparations ingested, however, were compounds containing substances such as phenacetin and caffeine and were taken in large doses. Data from the United States showed no significant effect of aspirin ingestion on birth weight or perinatal mortality.11 There is no convincing evidence that indomethacin or other non-steroidal anti-inflammatory agents affect fetal growth.

EFFECTS MEDIATED THROUGH INHIBITION OF PROSTAGLANDIN SYNTHESIS

Table II summarises the conditions associated with the use of inhibitors of prostaglandin synthesis in pregnancy.

Table II—Conditions associated with use of inhibitors of prostaglandin synthesis in pregnancy

<table>
<thead>
<tr>
<th>Effects on mother</th>
<th>Anti-inhibitors of prostaglandin synthesis</th>
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<tbody>
<tr>
<td>Prolongation of pregnancy</td>
<td>Anti-inhibitors of prostaglandin synthesis</td>
</tr>
<tr>
<td>Increased blood loss both before and after birth</td>
<td>Anti-inhibitors of prostaglandin synthesis</td>
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</table>

A retrospective study of women with musculoskeletal disorders showed that those who took more than 3.25 g of aspirin a day during the last six months of pregnancy had a significantly longer gestation, longer labour, and greater blood loss at delivery than women who had not taken aspirin.22 Collins and Turner also found an increased incidence of anaemia, antepartum haemorrhage, and pre-eclampsia in women who took aspirin for long periods.7 Haemostatic abnormalities and a higher incidence of intracranial haemorrhage...
have been found in neonates whose mothers ingested aspirin within a few days of delivery.\(^\text{15,14}\)

In the fetus prostaglandin E\(_1\) causes relaxation of systemic and pulmonary vessels as well as the ductus arteriosus, and 90% of blood ejected by the right ventricle passes through the ductus arteriosus to the descending aorta.\(^\text{15}\) Administration of single doses of an anti-inflammatory agent to a variety of animals results in reversible constriction of the ductus arteriosus and a substantial increase in pulmonary artery pressure in the fetus. Long term exposure to anti-inflammatory agents in animals and humans is associated with increased amounts of pulmonary artery smooth muscle which results in persistent pulmonary hypertension in the newborn infant, with or without premature closure of the ductus arteriosus.\(^\text{15}\)

Neonatal respiratory complications attributed to the use of non-steroidal anti-inflammatory drugs in the treatment of premature labour support this association.\(^\text{7,16}\) Since other studies, however, have found no increase in fetal mortality when premature labour was suppressed with non-steroidal anti-inflammatory drugs,\(^\text{17}\) it is likely that the dose and duration of administration of the drug, the gestational age of the fetus at the time of exposure, and the time between the last dose of the drug and the birth of the infant are important factors. Infants born to mothers receiving long term anti-inflammatory treatment are probably most at risk.

**BREAST FEEDING**

Because non-steroidal anti-inflammatory drugs are weak acids they do not achieve high concentrations in milk. All manufacturers state in their drug information that these drugs should not be used in lactating women. This caution is based on lack of specific information rather than known adverse reactions, and the benefit associated with breast feeding may outweigh the risks of a carefully chosen drug. The appropriate drugs should have a short elimination half life and metabolites which are inert or rapidly eliminated, or both. Hydroxy or methyl metabolites are relatively stable in the infant's stomach whereas glucuronide derivatives may be cleaved, releasing active metabolites.\(^\text{18}\) Table III shows the suitability of various drugs. Reported side effects are uncommon, but plasma salicylate concentrations of 24 mg/dl were found in a breast fed child with metabolic acidosis whose mother was taking 2.4 g aspirin a day, and a grand mal fit occurred in a child whose mother was taking indomethacin.\(^\text{19}\)

**Antimalarial drugs**

Chloroquine salts (4-aminoquinoline compounds) cross the placenta and rapidly accumulate in the fetal uveal tract of mice.\(^\text{20}\) Teratogenic effects of these substances are probably dose related, and experience from countries where malaria is endemic affirms the safety of weekly prophylactic doses in pregnancy.\(^\text{21}\) Exposure during the first trimester to the doses required to treat rheumatic diseases, however, has resulted in fetal sensorineural hearing loss.\(^\text{22}\)

**BREAST FEEDING**

Both chloroquine and hydroxychloroquine have been found in small quantities in human milk. Despite their widespread use by lactating women for malaria prophylaxis and the lack of reported adverse effects in breast fed infants, the daily doses required for treatment of chronic rheumatic diseases may cause retinal damage, which is difficult to monitor in children of this age. Their use in lactating women is, therefore, not recommended.

**Sulphasalazine**

Sulphasalazine is increasingly used as a second line treatment in rheumatoid arthritis. Experience with sulphasalazine in the treatment of inflammatory bowel disease has shown that it is safe to use throughout pregnancy and lactation.\(^\text{23}\) As it impairs absorption of folic acid, supplementation is recommended during pregnancy.

**Gold salts**

Both gold thiomolate and auranofin, an oral gold preparation, have proved teratogenic in animals. Gold has been found in the liver and kidneys of an aborted human fetus, and there are reports of possible teratogenic effects.\(^\text{24,25}\) Two studies, however, report the safe use of gold during pregnancy.\(^\text{26,27}\) and therapeutic gold concentrations have been detected in cord blood without evidence of congenital defects.\(^\text{28}\)

**BREAST FEEDING**

Trace amounts of aurothioglucose have been detected in the milk of lactating women, and gold has been found bound to the red blood cells of breast fed infants.\(^\text{29}\) The theoretical possibility of toxicity precludes its use during breast feeding.

**Penicillamine**

The chelating agent penicillamine (dimethylcysteine) is used for treating Wilson’s disease, cystinuria, and rheumatoid arthritis. Its use in pregnancy has been associated with the development of a generalised connective tissue defect, similar to that of Ehlers-Danlos syndrome, in three babies.\(^\text{30,31}\) Two died, but in the third the cutis laxa was reversible. Many normal children, however, have been born to mothers taking penicillamine for Wilson’s disease,\(^\text{32,33}\) and it was proposed that in this disease the fetus was protected from the effects of penicillamine by the excessive maternal pool of copper. In another survey, however, one ventricular septal defect was the only abnormality reported in 27 pregnancies in patients with rheumatoid arthritis and cystinuria.\(^\text{34}\)

**Corticosteroids**

The pharmacology of corticosteroids and potential effects on the fetus are discussed in the article on treatment of asthma. In
two studies of pregnancy associated with corticosteroid treat-
ment 37 pregnancies in 24 patients with rheumatic diseases
were evaluated. Although five pregnancies resulted in abortion and
four in fetal death, this was in a group of patients at high risk of fetal
loss. Length of gestation and birth weight were within normal limits and
a few minor fetal abnormalities were considered unrelated to
steroid treatment in the mother. Both studies emphasised the
importance of giving the mother additional corticosteroids during
delivery and the rarity of fetal adrenocortical insufficiency.

BREAST FEEDING

In the doses most commonly used for treating rheumatic diseases
(15 mg of prednisolone a day or less) there is little chance of an infant
receiving appreciable amounts of prednisolone in breast milk.\(^3\)

Cytotoxic drugs

Alkylating agents and antimetabolites may be teratogenic and
mutagenic, and even if used after the first trimester of pregnancy the
fetus is susceptible to bone marrow depression, infection, and
haemorrhage.\(^3\) Azathioprine is the cytotoxic agent most commonly used
in rheumatic disorders. In a study of 125 pregnancies in renal
transplant recipients taking both azathioprine and prednisolone only
one infant showed a congenital abnormality, though a number had
lymphopenia, growth retardation, and an increase in chromosomal
breakage.\(^3\) Results of long term follow up studies in these children
would not be available. The risk of lymphoproliferative and gonadal
 disorders would be likely to cause concern.

BREAST FEEDING

Many cytotoxic drugs are found in appreciable amounts in human
milk, and the risk to the infant would outweigh any benefits of
breast feeding.

Guidelines for antirheumatic treatment in pregnancy

Adequate explanations of the possible risks of any proposed
treatment, with appropriate advice on contraception, are essential
when treating women of childbearing age. The use of drugs that
pose the least threat to the fetus will minimise anxiety should
pregnancy occur. In women with established rheumatic diseases it is
important to appreciate that without the use of agents which suppress
the disease pregnancy may not have occurred or may not
have been carried to term. Adequate control of the disease may also
enable a woman to feel capable of bearing and raising children.
There is a good chance of remission of rheumatoid arthritis during
pregnancy, though aggressive treatment may continue to be neces-
sary in systemic lupus erythematosus.\(^7\)

NON-STEROIDAL ANTI-INFLAMMATORY AGENTS

Drug action is a function of concentration and time. To minimise
the effects on the fetus drugs with a short elimination half life and
inactive metabolites—for example, ibuprofen, flurbiprofen, and
ketoprofen—should be used at the maximum tolerated dosage
interval. The most potent inhibitors of prostaglandin synthesis,
such as salicylates and indomethacin, should be avoided throughout
pregnancy, if possible, and certainly during the last trimester. Well
motivated women with moderate symptoms may be managed with
simple analgesics, paracetamol being the drug of choice.

DRUGS THAT MODIFY RHEUMATIC DISEASE

Only a small proportion of women require disease modifying
drugs. Antimalarial drugs are contraindicated in pregnancy or
lactation, whereas sulphasalazine appears to be safe, though foetal
supplements should be given. Treatment with gold and penicillina-
should not be started during pregnancy, and these drugs
should not be used by breast feeding women. If a woman becomes
pregnant while taking these drugs gold can be continued at the
longer possible dosage interval and penicillamine should be slowly
reduced or withdrawn. Pyroxidine supplements are recommended
as penicillamine may deplete maternal stores.\(^6\) Careful considera-
tion should be given to the use of cytotoxic agents in pregnancy.
Although azathioprine appears to be relatively safe, long term
follow up data are not yet available. Corticosteroids in the doses
required to treat rheumatic conditions seem to be safe in pregnancy
and lactation. Additional doses are needed to cover delivery. Fetal
adrenal insufficiency is rare.

References

3 Needleman BK. Antithrombotic medication during lactation. Br J Rheumatol 1985;24:
291-7.
5 Collins E. Maternal and fetal effects of acetoaminophen and salicylates in pregnancy. Obstet
7 Turner G, Collins E. Fetal effects of regular salicylate ingestion in pregnancy. Lancet 1975;i:
338-40.
8 Shrimpton OR, Heineisson O, Kaufman DW, Sinkvist V, Monson RR, Shapiro S. Aspirin and congenital
Sauunders, 1979:5-2.
11 Shapiro S, Monson RR, Kaufman DW, Sinkvist V, Heineisson O, Stone D. Perinatal mortality and
12 Lewis RR, Shulman JD. Influence of acetylsalicylic acid on the hypothalamic-pituitary system,
13 Rumack CM, Guggenheim MA, Rumack BH, Peterson RG, Johnson ML, Braultaite WY.
52-60.
14 Stuart MJ, Gross SJ, Eldred H, Greaves J. Effects of acetylsalicylic-acid ingestion on maternal
15 Rudolph AM. The effects of non-steroidal anti-inflammatory compounds on fetal circulation
16 Levin DL, Mills LJ, Weinberg AG. Haemodynamic pulmonary vasculature and myocardial
abnormalities secondary to pharmacological constriction of fetus ductus arteriosus: a possible mechanism
for persistent pulmonary hypertension and transient tricuspid insufficiency in the new born infant.
Circulation 1979;60:360-4.
17 Zuckerman A, Renz V, Robinovitch S. Inhibition of human premature labour by indomethacin.
18 Wilkinson AR, Aynsley-Green AJ, Mitchell M. Persistent pulmonary hypertension and abnormal
prostaglandin E levels in preterm infants after maternal treatment with aspirin. Arch Dis Child
1979;54:942-5.
19 Wilkinson AR, Landstroem V, Green K. Premature labour and indomethacin. Prenat Diagn
21 Lewis R, Larsen NH, Birnbom S. Malaria associated with pregnancy. Obstet Gynecol
1978;51:1006-700.
22 Hart CW, Naunton RF. The ototoxicity of chloromphenicol. Arch Otolaryngol 1964;80:
407-12.
24 Rogers JG, Anderson RMcD, Cow C. Possible teratogenic effects of gold. Aust Paediatr J
25 Stern E. Drug therapy in the perinatal period. In: Morselli PL, Garattini S, Sereni S, eds. Basic and
26 Zuckerman A. Gold and anti-malaria therapy. In: Mccarthy DJ, ed. Arthritis and allied conditions.
28 Mjölnernor OK, Dommerud SA, Rasmussen K, Gerdslien ST. Congenital connective tissue
defect probably due to d-penicillamine treatment in pregnancy. Lancet 1972;i:673-5.
29 Solomon L, Abrams G, Dinsor M, Berens L. Neonatal abnormalities associated with
30 Linacre A, Graeme J, Rodriguez-Alcoro J, Diaz-Perez JL. Reversible cutis laxa due to maternal
34 Popert AJ. Pregnancy and adrenal cortical hormones, some aspects of their interactions in
35 Yackel DB, Kemper RD, McCoulskey WM. Adrenocorticotetrioid therapy in pregnancy. Am J
36 Macnamara SA, Selley JA, Agnew JE. Secretion of prednisolone into breast milk. Arch Dis Child
1975;50:894-5.
37 Barber HRK. Fetal and neonatal effects of cytotoxic agents. Obstet Gynecol 1981;58(suppl 5):
705-11.
38 Nolan GH, Sweet RL, Laros RR. Renal cadaver transplantation followed by successful