significantly reduced by pseudopregnancy this therapy is valuable as a holding measure, preparing fertility for at least a few years. Pseudopregnancy is also valuable in cases with lesions inaccessible to surgery when castration might be the only alternative treatment. Even if relief is only temporary, hormonal treatment may enable surgical menopause to be postponed to an age when it is likely to be better tolerated. Precise diagnosis before treatment can be made without laparotomy. Formerly we used culdoscopes for this purpose, but often unsuccessfully because of adhesions in the pouch of Douglas. With the more versatile laparoscope this difficulty can almost always be overcome. Nausea and weight gain occurred in 25-30% of our cases, but the former usually ceased after about two months and normal weight was re-established a few weeks after the completion of treatment. Thromboembolic disease was not a problem.

In view of the foregoing I would urge gynaecologists not to discard the weapon of pseudopregnancy. It is often curative and even when further treatment, either hormonal or surgical, is required its contribution to the solution of what Andrews and Larsen call "a vexing clinical problem" is too valuable to be ignored.—I am, etc.,

J. A. CHALMERS
Worcester Royal Infirmary

Pitocin Bucal in Late Pregnancy

SIR,—In recent years we have pursued the policy, in certain cases, of prescribing Pitocin Bucal (oxytocin) tablets (Parke-Davis) in late pregnancy to "ripen" the cervix prior to inducing labour. Furthermore, from talking to colleagues, we believe we are not alone in advocating this practice. We have nevertheless been doubtful as to the efficacy of such therapy, though at first glance the concept of "priming the uterus" with repeated small doses of oxytocin would seem an attractive one.

We therefore undertook a small pilot study, conducted as a double-blind trial and in conjunction with Dr. David Evans of Parke-Davis. In this trial one Pitocin Bucal tablet (200 units oxytocin) or a placebo, was given three times a day to 44 primigravidae from 38 weeks onwards until delivery. Patients were fully informed and only those who were both certain of their dates and had had uncomplicated pregnancies were selected. All patients took a minimum of 21 tablets (seven days), the mean number being 57 for the 22 who took the active preparation and 55 for the 22 who took the placebo.

Cervical effacement (as judged by a cervix of more than 2 cm length at the beginning of therapy diminishing to less than 1 cm length at the onset of labour or induction) occurred in only 61% of those patients taking the active preparation, and when this figure is compared with 30% effacement in those taking placebo the difference is not significant. When we looked at the duration of the first stage of labour the results were likewise not significantly different in the two groups, though a mean of 7.6 hours for the Pitocin Bucal patients undoubtedly compared favourably with 10.2 hours for those taking placebo.

Our results lead us to conclude, therefore, that oxytocin as used in the form described here has no clinically detectable effect on uterine activity either in late pregnancy or labour. However, when we contrast our figures with those of Hayes and Kennedy, it counts as little as five years ago the mean duration of the first stage of labour at Guy's was 13.35 hours it is possible that any effect the oxytocin may be having in this respect is masked by other factors in the approach to management of labour.—We are, etc.,

T. H. COLTART
T. G. NASH
Department of Obstetrics,
Guy's Hospital Medical School,
London S.E.1

Eugynon 30 and Dermatitis

SIR,—Dr. S. J. Jachuck and Dr. H. L. Franks (3 August, p. 347) may be interested to hear of a woman who developed urticaria each time she took an oral contraceptive pill. In 1967, 1972, and 1974 she was prescribed Lyndiol-2.5, Norinyl-1, and Microgynon 30 respectively. On each occasion, after only a few tablets, she developed a mild diffuse urticarial rash, mainly on the limbs and chest. She continued with the first two drugs for only two weeks but was persuaded to complete the course of Microgynon 30. The rash faded during the pill-free week. Pirithon 4 mg was given with the second course of pills to see if it would modify the reaction. It had no effect and the woman stopped the pill of her own accord on the fifth day. During her first pregnancy, in 1957, she was found to be allergic to nickel but her subsequent pregnancies in 1961 and 1967 were uneventful. There is neither hormone, dye, nor a material common to these oral contraceptives, and no light has been shed on the problem so far.—I am, etc.,

DELLA F. MORRIS
Family Planning Association,
Wellington, Salop

Tetracycline-resistant Group A Streptococci

SIR,—Dr. R. J. Fallon (22 June, p. 663) expressed interest in the prevalence of tetracycline resistance among streptococci in different areas of Britain. Though the NHS has both social and commercial links with Glasgow the incidence of tetracycline-resistant Lancefield group A streptococci (Sr. pyogenes) observed here since 1970 differs markedly from that reported by Dr. Fallon for the Glasgow (north) area only some 30 miles (48 km) away.1 In 1970 3-5% of strains isolated from hospitals and general practice in Ayrshire were resistant, in 1971 14.2%, in 1972 36.8%, and in 1973 33.3%. Currently (the first half-year of 1974) the incidence is 19.4%. Quarterly figures reveal a much wider range of incidence, however, from 11% in the second quarter of 1970 to 55% in the same quarter of 1973. Invariably during this time respiratory tract isolates overall have exceeded numerically those of non-respiratory origin, with a fairly even distribution not only by year but also between hospitals and general practice. Interestingly, the mean peak incidences of tetracycline-resistance of 51.5% (fourth quarter of 1972) and 53.7% both coincided with unusually high but non-epidemic incidences of non-respiratory isolates, though not of Sr. pyogenes generally. Between these two quarter-years was one with an observed incidence of 26.8% for tetracycline-resistant Sr. pyogenes, and for that reason as much as any other I hesitate to predict what trend our last quarter's incidence of 22.2% is supposed to show. As reiterated in your leading article on tetracyclines (25 May, p. 400), resistance is now common enough to make a group of antibiotics unsuitable as a first choice for infections caused by Sr. pyogenes.—I am, etc.,

JOHN A. EMSLIE
Microbiology Department,
Ayrshire Central Hospital,
Irvine, Ayrshire


Antiemetics, Progynon, and Breast Cancer

SIR,—We agree with Dr. H. W. C. Ward (20 July, p. 169) that drugs which raise circulating prolactin levels should not be given to patients with breast cancer. Following our report on metoclopramide's potent prolactin-releasing properties (29 June, p. 729) we support his suggestion that it is best avoided in such patients.

It is well established that the phenothiazines raise circulating prolactin levels.1 We have shown that prochlorperazine, a phenothiazine which Ward suggests as an alternative to metoclopramide, also raises prolactin in man, thus making it, too, unsuitable. To find a suitable antiemetic which was without effect on prolactin levels we have screened a number of compounds of these cyclizine 50 mg intramuscularly did not alter circulating prolactin levels. We therefore recommend that cyclizine rather than prochlorperazine should be the antiemetic of choice for patients with breast cancer undergoing radiotherapy.—We are, etc.,

M. O. THORNER
G. VOLANS
G. M. BESSER
A. S. MCNEILLY
St. Bartholomew's Hospital, London E.C.1

1 Tuckington, R. W., American Journal of Medicine, 1972, 53, 389.

Blood Donors and the Transfusion Service

SIR,—While agreeing with much of your provocative leading article (27 July, p. 212) I must protest at your statement that the British blood transfusion service is clearly a lame duck. Foreigners reading your journal—and many do—might be forgiven for believing that our blood transfusion service,