A comparison of antrafenine and aspirin on platelet aggregation and frusemide-induced diuresis

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Summary
The effects of antrafenine were compared with aspirin and placebo on platelet aggregation and on the diuretic action of frusemide in normal volunteers. Aspirin significantly reduced platelet aggregation at 3 and 6 hr after administration, but antrafenine only at 3 hr. Only aspirin significantly reduced the increase in urine sodium and potassium produced by frusemide.

Antrafenine, an anthranilic acid ester derivative, is an analgesic agent which inhibits prostaglandin synthesis and has some anti-inflammatory properties (Wisanto et al., 1981). Other anti-inflammatory analgesics such as aspirin have been shown to inhibit, at least in part, the diuretic action of frusemide (Henry, 1980) and to inhibit platelet aggregation (Zucker and Peterson, 1968). The authors have compared the effects of antrafenine and aspirin on platelet aggregation and frusemide-induced diuresis in normal human volunteers.

Material and methods
Six healthy volunteers (5 male, 1 female) aged 20–22 years participated in the study. A history of peptic ulcer, analgesic-induced asthma and other chronic illness was excluded in all subjects, who abstained from use of any medication for at least one week before the study and from smoking on the days of the study. After a light breakfast of toast and fruit-juice they were given with 100 ml water the following treatments at weekly intervals in a double-blind randomized order based on 2 Latin squares, using a double-dummy technique:—(a) aspirin 600 mg in a capsule; (b) antrafenine 300 mg as two 150 mg tablets; (c) matching capsule and tablet placebos.

One hour after this treatment frusemide 20 mg as a tablet was given. The bladder was emptied before frusemide ingestion and a urine aliquot collected. All urine passed was then collected at hourly intervals for 7 hr, and urine volume, Na+ and K+ estimated.

Nine-millilitre blood samples were collected immediately before each analgesic treatment or placebo, and at 3 and 6 hr later into 1-ml 3-8% trisodium citrate solution.

Platelet-rich plasma was prepared as the supernatant after centrifugation at 120 g for 5 min. The remainder was centrifuged at 600 g for 5 min to prepare platelet-poor plasma. Platelet aggregation was measured in a Payton-Dual aggregometer using 1, 2 and 4 mol μmol adenosine diphosphate (ADP).

The effects on platelet aggregation are indicated in FIG. 1. The urinary excretion of sodium (unfilled symbols) and potassium (filled symbols) following administration of aspirin (—△—), antrafenine (—□—) and placebo (—○—) followed by frusemide. * indicates a significant difference of aspirin from placebo (P=0.05).

FIG. 1. Urinary excretion of sodium (unfilled symbols) and potassium (filled symbols) following administration of aspirin (—△—), antrafenine (—□—) and placebo (—○—) followed by frusemide. * indicates a significant difference of aspirin from placebo (P=0.05).
as the aggregating agent. Lateral shifts of the dose ratios were calculated for each blood sample.

Results

Urinary estimations

Aspirin significantly ($P=0.05$) reduced the frusemide-induced increase in sodium excretion at 2 and 3 hr, and potassium excretion at 2 hr (Fig. 1), but had no significant effect on urine volume. Antrafenine had no significant influence on either electrolyte excretion or urine volume.

Platelet aggregation

Both aspirin and antrafenine inhibited ADP-induced platelet aggregation at 3 hr ($P<0.05$), but only aspirin had a significant inhibitory effect at 6 hr ($P<0.05$) (Table 1).

Discussion

The mechanism underlying the inhibitory influence of non-steroidal anti-inflammatory drugs on the diuretic and anti-hypertensive effects of frusemide and thiazide diuretics is still uncertain. Although a pharmacokinetic mechanism may be present (Benet, 1979), it is also possible that the interaction involves inhibition of prostaglandin synthesis in the kidneys (Patak et al., 1975; Abe et al., 1978). Platelet aggregation is also thought to involve prostaglandin activity, and the anti-aggregatory effect of aspirin to involve inhibition of prostaglandin synthesis. This effect is permanent because aspirin irreversibly acetylates the platelet cyclo-oxygenase enzyme involved in prostaglandin synthesis (Roth et al., 1977). Platelet aggregation does not then return to normal until a new generation of platelets have entered the circulation (Marian, Packham and Mustard, 1980).

Antrafenine appears to differ from aspirin in certain important aspects. Firstly, although it has a marked inhibitory effect on platelet aggregation at 3 hr after administration, comparable to the effect of aspirin, this effect has disappeared at 6 hr while the effect of aspirin, as expected, is unchanged. Its effects in the platelet, therefore, appear to be reversible in comparison with the irreversible effects of aspirin. Secondly, while aspirin has a significant inhibitory effect on the sodium and potassium excretion induced by frusemide, antrafenine had no measurable effect under the conditions of this study, although such an effect in larger doses cannot be excluded. It can be concluded, however, that doses of aspirin and antrafenine with similar effects on platelet aggregation at 3 hr after ingestion had markedly different effects on frusemide-induced sodium and potassium excretion. The dose of antrafenine used, 300 mg, had been shown to be an effective analgesic in preliminary clinical testing (A. Hedges and P. Turner, personal communication) and in controlled double-blind clinical trials (Wisanto et al., 1981).

The clinical importance of these pharmacological differences has yet to be explored. Bowen et al. (1980) found that faecal blood loss after 7 days' treatment with antrafenine 450 and 900 mg daily was similar to that after placebo and significantly less than that after aspirin 1800 mg daily, but experience of its use is still too limited to conclude that it produces fewer episodes of major blood loss. There is little doubt that anti-inflammatory drugs such as aspirin and indomethacin can antagonize the effects of diuretics and precipitate heart failure (Turner and Warrington, 1980).

It will be interesting to see if the absence of interaction of antrafenine with frusemide in this single-dose volunteer study truly predicts a lack of this action in the clinical situation.

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