Interaction of ketoprofen and frusemide in man


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Summary: The effects of ketoprofen on frusemide-induced diuresis, natriuresis and renin release were studied in 12 healthy male volunteers. Each received frusemide 40 mg once daily with either ketoprofen 100 mg twice daily or placebo for two periods of 5 days separated by a treatment-free period according to a randomized, double-blind, cross-over study design. Ketoprofen significantly reduced frusemide-induced diuresis on Day 1 but not on Day 5 of treatment. The natriuresis induced by frusemide on Day 1 or Day 5 of treatment did not differ significantly whether ketoprofen or placebo was administered, although the mean urinary sodium excretion values were consistently lower following ketoprofen. Ketoprofen did not affect the kaliuretic response to frusemide on Day 1 or Day 5 of treatment. The increase in plasma renin activity after frusemide was inhibited by ketoprofen on both Day 1 and Day 5. These results suggest that ketoprofen reduces the diuresis and renin release induced by frusemide, but that the reduction in diuretic response may become less important after their repeated coadministration.

Introduction

Conflicting results have been obtained with respect to the effects of nonsteroidal anti-inflammatory drugs (NSAIDs) on frusemide-induced diuresis and natriuresis, with a blunting1-4 or no change5-7 being reported. NSAIDs inhibit the early increase in plasma renin activity (PRA) which follows frusemide administration.1-3,6,7 Ketoprofen is a propionic acid derivative which has been shown to be safe and effective in the symptomatic treatment of rheumatoid and osteoarthritic diseases.8,9 There is no information available on the interaction of ketoprofen with frusemide. The absence of an interaction would make ketoprofen a NSAID of first choice when coadministration with frusemide is necessary.

This study was designed to investigate the effects of ketoprofen on frusemide-induced diuresis, natriuresis, and renin release following single and multiple doses of both drugs in healthy volunteers.

Methods

Twelve healthy male volunteers, aged 20–34 years (mean age 24) and within 10% of their ideal body weight, gave informed written consent to the study which was approved by the Ethical Committee of the City and Hackney District Health Authority. They were non-smokers and drank no alcohol during the trial periods. They were on no medication for at least one week prior to the start of the study and only the study medications were allowed during the study.

Each subject was randomly allocated to receive frusemide (40 mg once daily) together with either ketoprofen (100 mg capsule twice daily) or a matching placebo for 5 days according to a double-blind, cross-over design, the two treatment periods being separated by 9 days. Dietary advice was given in an attempt to moderate their intake of foods high in sodium and potassium for 5 days prior to and during the whole of each treatment period. Water intake was not limited but they were instructed to drink at least 2 litres of fluid daily. Twenty-four hour urine collections were made just before the start of each treatment period to assess sodium and potassium excretion and thus the approximate dietary intake of sodium and potassium of the volunteers.

The subjects attended on the mornings of Day 1 and Day 5 of each treatment period after fasting from 22.00 h the previous evening. A standard light breakfast without caffeine was taken. They
assumed a supine position for 30 minutes after
which a blood sample was drawn for measurement
of PRA. After emptying their bladders, they took
frusemide and either ketoprofen or placebo with
100 ml water. Following this, they drank 100 ml
water hourly for the next 5 hours. Another blood
sample for PRA was taken at 1 hour post-dosing,
after a 30 minute period of supine rest. Hourly total
urine collections were made during the first 6 hours
after dosing. The subjects were then allowed home
to complete their 24 hour urine collections.

Urine volumes were recorded and 10 ml aliquots
stored at −20°C until analysis for sodium and
potassium. Blood samples for measurement of
PRA were collected into tubes containing sodium
ethylenediamine tetra-acetate (EDTA) and kept on
ice. After centrifugation at 4°C, the plasma was

Figure 1  Cumulative urinary volume (ml), sodium and
potassium excretion (mmol) following administration of
frusemide 40 mg with either placebo (—O—) or keto-
profen 100 mg (—●—) on Day 1 of the treatment period.
Each point represents the mean value (± s.e.m.) of 12
subjects.

Figure 2  Cumulative urinary volume (ml), sodium and
potassium excretion (mmol) following administration of
frusemide 40 mg with either placebo (—O—) or ketopro-
fen 100 mg (—●—) on Day 5 of the treatment period.
Each point represents the mean value (± s.e.m.) of 12
subjects.

separated immediately and stored at −20°C until
analysis. PRA was measured by radioimmuno-
assay of angiotensin I (AI) generated under stan-
dard conditions and was expressed as pmol of
angiotensin I generated h⁻¹l⁻¹ of plasma at pH 7
and at 37°C. Urinary sodium and potassium concen-
trations were analysed by automated flame
photometry.

The hourly total urine volumes, sodium and
potassium excretion in the first 6 hours after drug
administration were analysed using multiple linear
regression analysis with treatment, time or
measurement and subjects included as independent
variables, employing the dummy variable techni-
que. Student’s paired t test was used to examine the
treatment effect on PRA, 24 hour urine volumes,
sodium and potassium excretion.
Results

Twenty-four hour sodium excretion values on the day prior to each treatment period were similar: 113 ± 93 mmol (mean ± s.d.) prior to placebo and 117 ± 79 mmol prior to ketoprofen. The potassium excretion values were 38 ± 14 and 50 ± 26 mmol respectively. These differences were not significant.

Ketoprofen, when compared with placebo, significantly reduced frusemide-induced diuresis over the 6 hours after drug administration (mean difference = -67 ml, \( P < 0.05 \)) and the 24 hour urine output (mean difference = -651 ml, \( P < 0.05 \)) on Day 1 (Figure 1). On Day 5 of treatment (Figure 2), the changes did not reach significance.

The natriuresis induced by frusemide on Day 1 or Day 5 of treatment did not differ significantly whether ketoprofen or placebo was administered, although the mean urinary sodium excretion values were consistently lower following ketoprofen (Figures 1 and 2).

Similar changes were seen for frusemide-induced kaliuresis although the magnitude of the mean differences was much smaller (Figures 1 and 2).

Frusemide, when administered with placebo, significantly increased PRA over basal values on both Day 1 (mean difference = 183 pmol/l/h, \( P < 0.05 \)) and Day 5 (mean difference = 355 pmol/l/h, \( P < 0.01 \)) of the treatment period. Ketoprofen abolished this frusemide-induced rise in PRA on both days (Figure 3).

Discussion

The results of this study demonstrated that ketoprofen suppressed frusemide-induced diuresis and increased in PRA following concomitant administration of single doses of both drugs. On Day 5 of treatment, however, ketoprofen had no significant effect on diuresis while PRA remained suppressed. Ketoprofen did not have a significant effect on the natriuretic response although the mean urinary excretion values were consistently lower when ketoprofen was administered. These results were obtained with doses and route of administration of frusemide and ketoprofen which are commonly used in every day clinical practice. We chose to study for any interaction after both drugs were administered simultaneously because patients commonly take their medication together and the times to peak concentration of the drugs were comparable.\(^\text{10,11}\)

The interference by NSAIDs with the diuresis and natriuresis induced by frusemide has previously been demonstrated by others\(^\text{1-4}\) and is believed to be due to inhibition of renal prostaglandins by NSAIDs. However, other investigators
have demonstrated no interference with these actions of frusemide.\textsuperscript{5-7} The reasons for the disparity in results between these studies are unclear and may be partly due to difference in trial design, route of frusemide administration, specific NSAID employed, salt intake of the subjects, duration of treatment and dosage used. It is also possible that the natriuresis following frusemide is not entirely prostaglandin-dependent and that other prostaglandin-independent mechanisms may be responsible for the interaction of NSAIDs on frusemide-induced diuresis and natriuresis.\textsuperscript{7} Our finding of a greater effect following acute than chronic administration of frusemide and ketoprofen also suggests that inhibition of prostaglandin inhibition may not be the only mechanism involved in this interaction, since one would expect a greater effect on prostaglandin inhibition with repeated administration of ketoprofen. Whatever the mechanisms involved, the results obtained in this study, using doses which represent commonly prescribed treatment regimens for both drugs, indicate that this interaction may be less important on longer term dosing.

A biphasic renin response is described with loop diuretics. There is an early stimulation, mostly observed after intravenous administration, which is thought to be a prostaglandin-mediated event, followed by a late rise which is due to salt and water depletion. The early increase in PRA after frusemide is consistently blocked by NSAIDs\textsuperscript{1-3,6,7} and our results are in accordance with these studies. The suppression of frusemide-stimulated renin release by ketoprofen in the absence of a significant effect on natriuresis is in keeping with the findings of other investigators that sodium retention and presumed consequent volume expansion is not the mechanism for renin suppression of NSAIDs\textsuperscript{1,2,6,7}.

We conclude that ketoprofen interferes with the diuresis and renin release induced by frusemide but the interference with the diuretic response of frusemide may become less important after repeated coadministration of both drugs. The absence of such an effect by a useful analgesic drug may be a potentially valuable property in clinical situations requiring both drugs. From these data, it cannot, however, be predicted whether similar results will be obtained in patients.

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References