Addition of dimethylsulphoxide to methyl-tert-butyl ether and ethyl propionate increases cholesterol dissolving capacity and cholesterol gall stone dissolution in vitro

J J G H M Bergman, A K Groen, K Huibregtse, G N J Tytgat

Abstract
There is a discrepancy between in vitro cholesterol dissolving efficacy of methyl-tert-butyl ether (MTBE) and ethyl propionate and cholesterol gall stone dissolution in vivo. This study investigated whether the presence of bile changes the cholesterol dissolving capacity of MTBE and ethyl propionate. The addition of dimethylsulphoxide to MTBE or ethyl propionate was also studied to discover if it improves the dissolving capacity for cholesterol gall stones. The presence of bile caused a 25% decrease in cholesterol dissolving capacity of both MTBE and ethyl propionate (p<0.0001). This inhibitory effect of bile could be overcome by the addition of dimethylsulphoxide: dimethylsulphoxide caused an increase in cholesterol dissolving capacity of MTBE and ethyl propionate, the increase depending on the dimethylsulphoxide/bile ratio in the mixture. Mean dissolution time of weight, size, and patient matched cholesterol gall stones was 220 minutes in MTBE and 130 minutes in MTBE/dimethylsulphoxide (p<0.0001). No stones dissolved completely in ethyl propionate or ethyl propionate/dimethylsulphoxide within 300 minutes. In conclusion, MTBE/dimethylsulphoxide is a more potent dissolving agent for cholesterol gall stones than MTBE, giving a 40% reduction in dissolution time. Addition of dimethylsulphoxide to ethyl propionate does not result in faster stone dissolution. MTBE and MTBE/dimethylsulphoxide are far superior to ethyl propionate as solvents for cholesterol gall stones.

Contact dissolution treatment of gall bladder stones, using methyl-tert-butyl ether (MTBE) or ethyl propionate, is successful in about 95% of selected cases.1-4 MTBE and to a lesser extent ethyl propionate have proved to be effective solvents for cholesterol in vitro.5 There is, however, a clear discrepancy between in vitro efficacy of MTBE and ethyl propionate as cholesterol solvents and time required for dissolution of gall stones in vivo. This is one of the important drawbacks of contact dissolution treatment: treatment is labour intensive and time consuming, negatively influencing cost efficacy. Although studies have been published describing an automated pump, such a device has not become commercially available since publication seven years ago.6-8 Treatment therefore still requires manual flushing of the gall bladder for 4–16 hours during several days.1,2 Changes to the treatment regimen resulting in faster stone dissolution can thus be of major importance.

Two factors are known to inhibit dissolution of stones in vivo. Smith et al showed that gallstone dissolution in vitro was inhibited by insoluble stone matrix, existing of calcium, bilirubin, and protein.9 10 They speculated that the matrix limits stone dissolution because of its spatial distribution within the stones as concentric rings that retard contact of the cholesterol solvent with underlying layers of crystalline cholesterol. Addition of a matrix dissolving agent has been shown to accelerate dissolution of gall stones in vitro.11-13

Less attention has been given to stone-solvent contact as a factor influencing contact dissolution in vivo: MTBE/bile and ethyl propionate/bile are biphasic mixtures with a lighter ether phase above and a heavier bile phase below. Because gall stones have a higher density than bile, they sink into the bile phase and do not come in close contact with the dissolving agents in the upper phase.14-16 Creating turbulence manually as during treatment in vivo can only partially overcome this effect.

So far the effect of bile components on cholesterol dissolution in MTBE and ethyl propionate has not been investigated. Almost all in vitro studies on solvents for gall stones have been performed in the absence of bile. Therefore such an effect cannot be excluded.

Addition of dimethylsulphoxide to MTBE or ethyl propionate could enhance stone dissolution in vivo. Dimethylsulphoxide is a potent solvent for bilirubin and easily mixes with MTBE and ethyl propionate in contrast with other stone matrix dissolving agents like EDTA or N-acetylcysteine.17 By its dissolving effect on bilirubin dimethylsulphoxide could overcome part of the shielding effect of the stone matrix on cholesterol dissolution in MTBE or ethyl propionate. Dimethylsulphoxide could also improve stone solvent contact.16 In the presence of bile, monophasic mixtures of MTBE/dimethylsulphoxide or ethyl propionate/dimethylsulphoxide become biphasic: MTBE or ethyl propionate in the upper phase, dimethylsulphoxide/bile in the...
lower phase. This probably reflects the high affinity of the bipolar dimethylsulphoxide for aqueous media and the poor solubility of bile in MTBE. Because dimethylsulphoxide has a higher density than bile, the density of the lower phase increases. Stones that sink into the bile phase will float at the interphase, coming in closer contact with the dissolving agents of the upper phase. Presence of dimethylsulphoxide could change the cholesterol solubilising capacity of MTBE and ethyl propionate in a similar way, however, as bile might have a negative effect on cholesterol dissolution in these solvents.

The aims of this study were twofold. Firstly, to test if the presence of bile changes the cholesterol dissolving capacity of MTBE and ethyl propionate, and secondly to investigate if addition of dimethylsulphoxide improves the dissolution capacity of MTBE and ethyl propionate for cholesterol gall stones.

Methods

SOLVENTS AND BILE
MTBE, ethyl propionate, and dimethylsulphoxide (all synthetical grade) were purchased from Merck (Darmstadt, Germany). Cholesterol (monohydrate, 99% pure) was provided by Sigma Chemical Company (St Louis, MO). Hepatic bile was collected from one single patient by a T drain after cholecystectomy. To eliminate changes in bile composition because of surgery, bile collection was started three days postoperatively.

GALL STONES
Human cholesterol gall stones were obtained at laparoscopic cholecystectomy. Gall stones were selected from patients whose gall bladders contained more than 10 stones similar in size and morphology. Black pigment stones were identified visually and excluded from this study. A total of 46 gall stones obtained from three patients were selected. Stones from each patient were kept separate from those of other patients and stored in bile at −20°C. To verify if stones were actually cholesterol gall stones one stone from each patient was subjected to compositional analysis. All stones were incubated in bile and x rayed using the same settings as for plain abdominal x ray. Only stones that were completely radioluent were used for dissolution experiments. After washing in deionised distilled water to remove bile and debris, stones were air dried for 30 minutes. Using a Mettler HL52 balance, individual gall stone weight was determined. Maximum and minimum diameter were measured using calipers and rulers. For paired dissolution experiments 13 stone pairs, matched for weight, size, and patient, were selected. A difference of more than 10% between two stones of the same stone pair was not permitted in respect to weight or sum of maximal and minimal diameter.

INFLUENCE OF BILE AND WATER ON CHOLESTEROL DISSOLUTION CAPACITY OF MTBE AND ETHYL PROPIONATE
Mixtures of MTBE/bile, MTBE/water, ethyl propionate/bile, and ethyl propionate/water were made in volume ratios of 90/10, 80/20, 70/30, and 50/50 such that total volume of each mixture was 5 ml. All mixtures were biphasic: the lighter MTBE or ethyl propionate phase above, the heavier bile or water phase below. To each biphasic mixture an excess of pure cholesterol crystals was added (ratio cholesterol/solvent: 200 mg/ml MTBE, 100 mg/ml ethyl propionate). After addition of cholesterol, mixtures were incubated at room temperature in a REAX 2 circulator (60 rpm) for 30 minutes. The mixtures were then centrifuged at 2000 rpm for two minutes and cholesterol concentration in the upper phase (MTBE or ethyl propionate) was determined using the cholesterol oxidase method (Boeringer, Mannheim). The cholesterol concentration measured was considered to represent the maximum cholesterol dissolving capacity of the solvent at room temperature.

EFFECT OF ADDITION OF DIMETHYLSULPHOXIDE TO MTBE/BILE AND ETHYL PROPIONATE/BILE MIXTURES ON CHOLESTEROL DISSOLUTION CAPACITY OF MTBE AND ETHYL PROPIONATE
Monophasic MTBE/dimethylsulphoxide and ethyl propionate/dimethylsulphoxide mixtures were prepared in volume ratios of 90/10, 80/20, 70/30, and 50/50. These mixtures were then added to different amounts of bile to give final percentages of 10, 20, 30, and 50% of bile. The total volume of each mixture was 5 ml. Addition of bile caused an immediate phase separation leading to biphasic mixtures with an upper MTBE or ethyl propionate phase and a lower phase consisting of dimethylsulphoxide and bile. This phase separation was an exothermic process causing a rise in temperature of about 8°C in every mixture. To ensure that dissolution tests were performed under isothermic conditions all MTBE/dimethylsulphoxide/bile and ethyl propionate/dimethylsulphoxide/bile mixtures were circulated in a REAX 2 circulator (60 rpm) for 30 minutes before addition to cholesterol. To each mixture an excess of cholesterol was added (ratio cholesterol/solvent: 200 mg/ml MTBE, 100 mg/ml ethyl propionate). Mixtures were subsequently treated as described above. During the dissolution experiments no change of temperature was seen.

PAIRED STONE DISSOLUTION EXPERIMENTS
Of the 13 stone pairs selected, seven were used to compare stone dissolution in MTBE with a dissolution regimen using MTBE/dimethylsulphoxide (70/30). The other six stone pairs were subjected to dissolution in ethyl propionate and ethyl propionate/dimethylsulphoxide (70/30). Stones were put in separate glass test tubes in which the tip of a polyethylene
nasobiliary pigtail catheter (diameter 7F, length 120 cm), was placed. This same type of catheter is used in transpapillary contact dissolution treatment in vivo in our unit. Hepatic bile (2 ml) was added and gall stones were subsequently dissolved in 8 ml of either MTBE, MTBE/dimethylsulphoxide (70/30), ethyl propionate or ethyl propionate/dimethylsulphoxide (70/30). The in vivo situation was mimicked by creating turbulence manually using glass syringes (4–6 cycles a minute) and by exchanging solvents and bile every 15 minutes. Test tubes were incubated in the dark at 37°C. Every 15 minutes gall stones were air dried for three minutes and weighed on a Mettler HL52 balance. Dissolution was expressed as the percentage of initial dry stone weight remaining at each time point. If during the dissolution process stones disintegrated no further weight determination was performed until the end of the experiment to avoid artificial fragmentation caused by handling of the fragments. The experiment was stopped whenever gall stones had completely dissolved or a time limit of 300 minutes had passed.

STATISTICS
Results are expressed as percentages or mean

Figure 1: Cholesterol concentration in MTBE phases of MTBE/dimethylsulphoxide (80/20)/bile and MTBE/bile mixtures after addition of an excess of cholesterol.

Figure 2: Cholesterol concentration in MTBE phases of MTBE/dimethylsulphoxide (70/30)/bile and MTBE/bile mixtures after addition of an excess of cholesterol.

In MTBE/bile mixtures cholesterol concentration was 25% (95% confidence intervals: 22 to 27%, p<0.0001). In ethyl propionate mean decrease in cholesterol concentration was 28% (95% confidence intervals: 23 to 34%, p<0.0001). Data are given as mean of four separate experiments. Standard deviation were less than 5%. For this negative effect on cholesterol dissolution only small amounts (5%) of water or bile were necessary. Increasing the volume ratio of water or bile in the mixtures beyond this 5% did not cause any further decrease in cholesterol dissolution capacity of the solvents. There was no difference in effect on cholesterol dissolution between addition of water or bile.

RESULTS

INFLUENCE OF WATER AND BILE ON CHOLESTEROL DISSOLUTION CAPACITY OF MTBE AND ETHYL PROPIONATE

Compared with pure MTBE or ethyl propionate, the presence of water or bile caused a 25% decrease in cholesterol dissolution capacity of both solvents. In MTBE decrease in cholesterol dissolution capacity was 25% (95% confidence intervals: 22 to 27%, p<0.0001).

In ethyl propionate mean decrease in cholesterol concentration was 28% (95% confidence intervals: 23 to 34%, p<0.0001). Data are given as mean of four separate experiments. Standard deviation were less than 5%. For this negative effect on cholesterol dissolution only small amounts (5%) of water or bile were necessary. Increasing the volume ratio of water or bile in the mixtures beyond this 5% did not cause any further decrease in cholesterol dissolution capacity of the solvents. There was no difference in effect on cholesterol dissolution between addition of water or bile.

EFFECT OF ADDITION OF DIMETHYL SULPHOXIDE TO MTBE/BILE AND ETHYL PROPIONATE/BILE MIXTURES ON CHOLESTEROL DISSOLUTION CAPACITY OF MTBE AND ETHYL PROPIONATE

Figures 1 and 2 show the mean cholesterol concentration in MTBE phases of MTBE/dimethylsulphoxide/bile and MTBE/bile mixtures after addition of an excess of cholesterol. Figures 3 and 4 show this information for ethyl propionate/dimethylsulphoxide/bile and ethyl propionate/bile mixtures. Data are given as mean of four separate experiments. Standard deviation were less than 5%. In mixtures containing dimethylsulphoxide, the MTBE and ethyl propionate phases had a significantly higher mean cholesterol concentration than in the mixtures without any dimethylsulphoxide, p<0.001. The increase in cholesterol dissolution capacity depended on the percentage of bile in the mixtures, as shown in Figures 1–4 separately.

The differences between Figures 1 and 2 and Figures 3 and 4 show, however, that changing the MTBE/dimethylsulphoxide or ethyl propionate/dimethylsulphoxide ratio, also changed the cholesterol dissolution capacity of the solvents. Therefore the dimethylsulphoxide/bile ratio was calculated for all mixtures and related to the cholesterol concentration in the MTBE or ethyl propionate phases (Fig 5). Increase in cholesterol dissolution capacity clearly depended on the dimethylsulphoxide/bile ratio in the mixture, a rise of about 100% being reached at a ratio of about 2. Both MTBE and ethyl propionate showed the same effect, however cholesterol dissolution capacity of ethyl propionate was about 50% of that of MTBE.
PAIRED STONE DISSOLUTION TESTS
All stones used in paired dissolution experiments were radiolucent in x-ray and had a cholesterol content higher than 95%. Stones added to MTBE or ethyl propionate all sank into the bile phase whereas stones treated with MTBE/dimethylsulphoxide or ethyl propionate/dimethylsulphoxide all floated at the interphase. This difference in stone solvent contact, however, was probably minimised by the turbulence created manually with glass syringes. In MTBE and MTBE/dimethylsulphoxide mixtures all cholesterol gall stones dissolved completely. Mean time required for complete stone dissolution was 220 minutes in MTBE (SD 21 minutes, range 180–240 minutes) and 130 minutes in MTBE/dimethylsulphoxide (SD 28 minutes, range 105–180 minutes). Figure 6 shows the results of these paired dissolution tests. Compared with MTBE, MTBE/dimethylsulphoxide mixtures gave a faster stone dissolution in every stone pair. Mean time reduction was 90 minutes (95% confidence intervals: 65 to 115 minutes, p<0.0001).

In ethyl propionate and ethyl propionate/dimethylsulphoxide no cholesterol gall stone dissolved completely within the time frame of 300 minutes, leaving a mean stone residue of 35% in ethyl propionate and 28% in ethyl propionate/dimethylsulphoxide mixtures (p=0.13). In one stone pair we continued the dissolution process up to 480 minutes. Stone dissolution was still incomplete with a stone residue of 16–7% in the ethyl propionate treated stone and 15–8% in the ethyl propionate/dimethylsulphoxide stone. The difference in dissolution of cholesterol gall stones between mixtures with ethyl propionate and mixtures with MTBE was highly significant both for dissolution time (p<0.0001) and for successful stone dissolution (p<0.05).

Discussion
We showed that addition of bile has a negative effect on cholesterol dissolution in MTBE and ethyl propionate. The presence of bile caused a 25% decrease in the cholesterol dissolving capacity of both MTBE and ethyl propionate. There was no difference between addition of water or bile, suggesting that the inhibitory effect of bile on cholesterol dissolution in MTBE and ethyl propionate is probably caused by its water component. The fact that only small amounts of water or bile were necessary to cause this effect reflects in our opinion the poor solubility of water in MTBE and ethyl propionate. Only a small amount of water is sufficient to completely saturate the solvent with water and to cause the decrease in cholesterol dissolving capacity. Addition of more water does not result in a higher concentration of water in the ether phase and thereby has no further effect on cholesterol dissolution. Because of the negative effect of water and bile on cholesterol dissolution in MTBE and ethyl propionate, we feel that all in vitro testing on solvents for contact dissolution of gall stones should be performed in the presence of water or bile.

The negative effect of bile can be compensated by the addition of dimethylsulphoxide: in our experiments we showed that addition of dimethylsulphoxide to MTBE/bile or ethyl propionate/bile mixtures increases the cholesterol dissolving capacity of MTBE and ethyl propionate compared with mixtures without dimethylsulphoxide. This increase depends on the dimethylsulphoxide/water ratio of the mixture, a maximum increase of 100% being reached at a ratio of 2. We hypothesise that the positive effect of dimethylsulphoxide on cholesterol dissolution in MTBE and ethyl propionate in the presence of bile is caused by extraction of water from the MTBE (ethyl propionate) phase into the bile/dimethylsulphoxide phase, thereby overcoming the negative effect of water on cholesterol dissolution in MTBE and ethyl propionate.

We found during contact dissolution treatment in our patients that about 10–20% in the volume extracted from the gall bladder after 15 minutes of flushing, existed of bile. Therefore we speculated that mixtures of MTBE/dimethylsulphoxide and ethyl propionate/dimethylsulphoxide in volume ratios of 70/30 would be most suited for stone dissolution experiments. In this way the increase in cholesterol dissolution capacity of the solvents could compensate for the loss of absolute volume of solvent entered in the gall
bladder at every cycle. More importantly, the volume of dimethylsulphoxide entered in the gall bladder would be enough to raise the density of the bile phase and to exert its dissolving effect on the stone matrix.

Our paired stone dissolution tests show that addition of dimethylsulphoxide to MTBE results in a 40% time reduction compared with a treatment regimen with only MTBE. This time reduction achieved by addition of dimethylsulphoxide is impressive, a mean reduction of 90 minutes on a total dissolution time of 220 minutes. The time required for complete stone dissolution in MTBE is in agreement with in vivo data from other studies in which solitary stones were dissolved after a mean dissolution time of four hours. We feel that this supports the adequacy of our in vitro model for gall stone dissolution.

In ethyl propionate mixtures addition of dimethylsulphoxide does not result in a significant change in dissolution time. We speculate that the more polar molecular structure of ethyl propionate causes ethyl propionate to compete with bilirubin for dissolution in dimethylsulphoxide.

Stone dissolution experiments were performed in pairs. The main analysis was aimed at comparing MTBE with MTBE/dimethylsulphoxide and ethyl propionate with ethyl propionate/dimethylsulphoxide. Comparing mean dissolution time of MTBE mixtures and ethyl propionate mixtures, however, provides important information. Stone pairs used in dissolution tests with ethyl propionate were significantly lower in weight than stone pairs obtained from the same patient dissolved in mixtures with MTBE. Any bias resulting from stone size or weight was therefore directed towards a faster stone dissolution in ethyl propionate.

MTBE mixtures were far superior to ethyl propionate mixtures both for dissolution time and ultimate successful dissolution. We think that these results reflect the higher cholesterol dissolving capacity of MTBE (about twice that of ethyl propionate) and its lower viscosity. This contrasts with the results of Zakko et al who found ethyl propionate more effective in dissolving gall stones in vitro than MTBE. In their study they used a more artificial model with an automated high flow system. Because such a pump is not available commercially and has not been approved by the Food and Drug Administration we feel that our system represents the in vivo situation in a more proper way.

After MTBE dissolution insoluble debris may remain. This might act as a nidus for gall stone recurrence. Nelson et al found this debris to exist predominantly of calcium bilirubin and calcium carbonate. They showed that a mixture of EDTA with either cholate or polysorbate was most efficient in dissolving these components. Dimethylsulphoxide also has dissolving capacity for these substances and can, in contrast with hydrophilic stone matrix solvents like EDTA or N-acetylcysteine, be mixed with MTBE, making application in vivo more favourable. Addition of dimethylsulphoxide to MTBE could thus not only result in faster stone dissolution but could also lead to less residual debris after dissolution. In our studies residual debris after successful dissolution with MTBE or MTBE/dimethylsulphoxide was too small to show such a difference. Others have, however, shown this effect using different matrix solvents.

Dai et al studied the cytotoxicity of MTBE and MTBE/dimethylsulphoxide (70:30) infused in gall bladders of rabbits. Their histological study of several rabbit tissues showed that chemical irritation by MTBE/dimethylsulphoxide (70:30) was identical to that caused by MTBE (100%) and localised to the level of the gall bladder wall. Normal morphology was rapidly recovered in two weeks and no changes were seen in liver cells, duodenum, kidney or medulla. Somnolence occurred more frequently in the MTBE group suggesting that systemic absorption of MTBE was not increased by the presence of dimethylsulphoxide.

We conclude that using MTBE/dimethylsulphoxide (70:30) instead of MTBE may result in a considerable gain of time in contact dissolution treatment in vivo. Compared with ethyl propionate, MTBE is a superior solvent for contact dissolution in vitro. Addition of dimethylsulphoxide to MTBE may also result
in a more complete stone dissolution, leaving less residual debris. Toxicity of MTBE/ dimethylsulfoxide seems to be comparable with MTBE. Further testing, however, especially of toxicity in humans is mandatory before application in vivo.

The results of these studies were presented in part at the annual meeting of the American Gastroenterological Association in May 1993 in Boston and were published in abstract form in Gastroenterology 1993; 104: A350.

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