Focused liver ablation by cavitation in the rabbit: a potential new method of extracorporeal treatment

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Abstract
A new device was used to achieve focused tissue ablation by shockwave induced cavitation. The device produced a half cycle of negative pressure followed by a shock wave, thus enhancing cavitation. Twenty eight New Zealand rabbits were treated. Therapeutic ultrasound was targeted at the centre of the liver under ultrasound guidance. The focal volume was scanned with a computer operated x-y-z micropositioner. The number and frequency of bursts as well as the distance between two x-y-z displacements were pre-selected. The relation of tissue ablation seen to preselected parameters, effects on surrounding tissues, biological side effects, and mode of healing were studied. Macroscopy, planimetry, and quantitative microscopy were used. Focused and homogeneous tissue ablation was achieved within well defined limits. Maximal tissue ablation was seen in the centre of the target. Liver surrounding the target remained unaffected. Lesions were made of a-cellular spots surrounded by disorganised rims of necrotic hepatocytes; 24 hours after treatment, the changes (mean (SEM)) in alanine transaminase and haemoglobin were +225 (36)% and -2-4 (2)% respectively. Serum transaminases, haemoglobinemia, and packed cell volume were normal 21 days after treatment and the target area was covered by a fibrous scar. It is concluded that ultrasound cavitation may achieve extracorporeal intrahepatic tissue ablation inside a predetermined target. This technique should now be tested in an animal hepatic tumour model.

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Malignant tumours of the liver remain an important health problem throughout the world. Non-surgical treatments have shown essentially palliative effects and surgery is potentially curative in only a small percentage of patients. Surgery needs the resection of nontumoral liver tissue and does not prevent recurrence. Recurrences can now be detected at early stages with the help of modern diagnostic methods. Hence, it seems reasonable to contemplate a minimally invasive method providing radical and selective cancer destruction, which could be applied to such tumours with some benefit. It was our hypothesis that acoustic cavitation induced by shock waves could be such a method. Cavitation is defined as the non-linear oscillation and collapse of gas bubbles suspended in a liquid under the influence of acoustic (mechanical) pressure waves: the expansion of bubble nuclei results from a depression within the medium and their collapse may be induced by a sudden compression wave after expansion. It has been shown that cavitation, when occurring at an interface near a cell membrane for example, was deleterious to biological media through mechanical, non-thermal effects. We showed the feasibility of radical tissue destruction in the rabbit liver in vivo by a concomitant administration of focused extracorporeal electrohydraulic shock waves with intravascularily infused gas microbubbles. Our results showed improvement shortcomings, however, which made the method inapplicable to the human or even to animal models of tumours without an unacceptable toxicity: tissue ablation was not focused, non-homogeneous, and needed an invasive and potentially toxic intra-arterial infusion of gas microbubbles to be efficient. Our objectives were thus to perfect the method to enhance cavitation, eradicate bubble injection, and to improve the focusing and selectiveness of tissue destruction. In this study, qualitative and quantitative assessments of liver ablation in the rabbit show that improvements of the method have made these achievements possible.

Materials and methods

ANIMALS
Twenty eight New Zealand hybrid rabbits of both sexes (Elevage Scientifique des Dombes, Ain, France) weighing 2-0 to 2-5 kg were used. All the experiments were conducted with premedication by intramuscular acepromazine (Vetoquinol, Lure, France) 4 mg/kg and induction and maintenance of anaesthesia by a continuous venous infusion of a mixture of ketamine (Substancia Parke-Davis, Courbevoie, France) 125 µg/kg/min and 2% xylazine (Bayer, Leverkusen, Germany) 1 µg/kg/min. The abdomen was shaved thoroughly and the abdominal wall was covered with a coupling agent. The animals were positioned on a purpose built plexiglas set up.

CAVITATION GENERATOR
The generator included an IBM/PC compatible computer for the overall control of the system, a pulse generator, an ultrasonic pulser (spark gap), a three axis stepper motor controller, a stepper motor driven micropositioner arrangement, an ultrasound scanner used to visualise the target, and a tank filled with degassed and deionized water (Fig 1A). The piezoelectric transducer array used as a shock wave source generator contained 18 single transducers mounted in a 28 cm diameter bowl and focused at 16 cm. The transducers were mechanically adjusted to pro-
The parallelepipedic target inside (2) the liver; (3) the plexiglas set up can be moved in the x, y, and z directions; the transducer and the animal are immersed into (4) degassed water; (5) imaging and targeting ultrasound (US) probe; (B) pressure tracing of the transducer and the computer.

METHODS

Short term end points

Macroscope – after fixation of the liver in acetic formalin, the whole organ was cut in 5 mm thick sagittal slices; slices including necrosis were compared with a video print of a sagittal ultrasound scan taken during the session. This methodology was used to accurately assess the x, y, and z dimensions of the macroscopically affected zone (as measured twice with a calliper square); calculations were corrected to take into account the volume reduction subsequent to formalin fixation (x 0.96) and correlated with the predetermined target volume.

Light microscopy – 4 μm thick tissue slices were prepared and stained with haemalum, cosin, and saffron (HES) for standard histological analysis.

Quantitative automated microscopy – this technique was used for the assessment of tissue lysis and viability: three 4 μm thick sections from the centre of the target area of each specimen were stained with periodic acid Schiff (PAS); each section was analysed three times under 10 magnification (nine fields/analysis) on a Quantimet 570 image analyser (Cambridge, UK); the analyser was requested to measure the surface of a section unstained with PAS (thus considered non-viable, even if morphologically unaffected) as a percentage of the whole surface included in the target area.

Planimetry was used to assess quality of tissue destruction as a function of the depth of the target. A video print was used as above to perform sectioning of the liver orthogonal to the z axis. Each 1 mm thick section was cut in 4 μm thick slices and stained with HES. Stained slices were photographed and magnified; planimetry (Digital polar planimeter 330, Germany) was repeated three times on each section to measure the surface of the lytic areas as a percentage of the whole surface.

Mid term end points

Clinical parameters – general condition of the animal during and after treatment, progress, and recovery from anaesthesia, weight variation between treatment and death were noted.

Biological parameters – haemoglobinemia, packed cell volume, and serum transaminases (alanine transaminase and aspartate transaminase) were assessed before treatment, 24 hours.
after, and 21 days after treatment in certain groups of animals.

Necropsy — thorough examination of the peritoneal cavity, diaphragm, lungs, heart, and kidneys was done. Particular attention was given to hepatic vessels damage.

TREATMENT PARAMETERS AND PROTOCOL
Twenty eight rabbits were assigned to the following groups of treatment:

Group 1 — eight rabbits were treated with the topographical parameters described above, at a burst frequency of 1 Hz, with a distance between points of 2 mm. They were killed 24 hours after treatment. Macroscopy of the liver, coeliac vessels, other intra-abdominal organs, kidneys, diaphragm, lungs, and heart was done, as well as a light microscopic examination of the target area, including quantitative microscopy.

Group 2 — eight rabbits treated with the same parameters were killed 21 days later; before death, all the animals were monitored clinically and biologically; on death, the animals had macroscopic and light microscopic examinations.

Group 3 — subgroups of three animals each were treated to assess the influence of the distance between two points (1, 2, and 4 mm) on the amount of tissue destruction, all other parameters remaining equal to group 1. Twenty four hours after treatment, the target area was submitted to a quantitative microscopic analysis as described above.

Group 4 — three rabbits were treated with the same parameters as group 1; after death at 24 hours, the target area was submitted to a planimetric analysis of tissue destruction as a function of the depth from the anterior pole of the target area.

Group 5 — three rabbits were submitted to two consecutive treatment sessions, 48 hours distant from one another; targeting for the second session was based upon the hyperechogenic remnant from the first one; 24 hours after the second session, the target area was examined by light microscopy.

Results

ULTRASOUND MONITORING OF THE TREATMENT
Progress of the treatment session was monitored through a sagittal view of the target zone, inside which tiny and fugitive hyperechogenic spots could be seen from the very first bursts; remnants of these spots gave the whole target area a whitish, brilliant aspect at the end of a session (Fig 2).

CLINICAL RESULTS
No animal died from anaesthesia during this
study; no animal died or presented any clinical complication of any sort during the follow up period, in any of the groups studied. Urine and faeces were normal in all cases. Food intake was unchanged after treatment and weight presented no abnormal (mean (SEM)) variation (2.17 (0.5) kg after treatment compared with 2.11 (0.3) before treatment). All animals presented petechiae (0.5 mm in diameter) over the abdominal wall, on a surface consistent with the cross section of the ultrasound beam at the skin level (diameter 2.5 cm, section 4.9 cm²). Petechiae disappeared after a few days.

BIOLGy
All animals from group 2 showed an increase in serum transaminase activities 24 hours after the session, between 1.5 and 6 times the normal value. Mean (SEM) variation of alanine transaminase was +225 (36)%; in all animals, the recovery was complete 21 days after treatment; haemoglobin records showed non-significant variations (−2.4 (2)%).

HISTOLOGICAL EXAMINATION
The target zone was always constituted of individual lytic spots surrounded by a rim of necrotic hepatocytes of disorganised architecture. The hepatic tissue around the affected area remained morphologically strictly normal (Fig 3A and B). Quantitative microscopy with PAS staining showed that tissue ablation was surrounded by a morphologically normal but unstained (non-viable) zone. In group 2, the livers examined 21 days after treatment presented always a fibrous whithis scar at the surface of the capsule; the scar was usually rectangular, 0.5 cm in width and less than 1 cm in length, retractile, with a regenerative aspect of the surrounding liver. The healing process was a collagenic fibrosis, poorly inflammatory, and sometimes nodular surrounded by proliferative hepatocytes presenting images of mitoses (Fig 3C).

Figure 4: (A) Percentage of tissue ablation/section of the affected volume as a function of the distance between points as estimated by quantitative microscopy and colorimetry (PAS); (B) percentage of tissue ablation as a function of the depth from the anterior pole of the target, as estimated by planimetry. Vertical bars represent standard error of the mean.

ANALYSIS OF PARAMETERS

Macroscopy – the distribution of affected tissue volumes from group 1 showed a dispersion around the predetermined (desired) volume; the mean (SEM) volume seen was 3.59 (1.23) cm³ for a predetermined volume of 2.56 cm³. The affected zone presented grossly more spheric than parallelepipedic, within sharp, well defined limits. Its position in the liver was always consistent with the desired coordinates – that is, in the centre of the organ and its top polar plane 25 mm deep from the skin. Within these limits, individual lesions were homogeneously distributed, although not falling into straight lines. Overall dimensions of the target zone of animals having two consecutive sessions (group 5) presented no difference with other groups, but tissue ablation presented was more complete, with almost no remaining viable hepatocytes. The percentage of tissue affected (as assessed by quantitative microscopy) decreased significantly when the distance between points was increased from 2 to 4 mm (p<0.01, Student's t test); however, a distance between points shorter than the focal spot diameter – that is, 1 mm – did not improve tissue ablation (Fig 4A).

The planimetric study of in depth tissue lysis (group 4) showed that the maximum effect was obtained 3 mm from the anterior (bottom) pole of the lesion (70% tissue lysis) and decreased anteriorly and posteriorly (Fig 4B).

Discussion
In a previous study, we have shown the feasibility of tissue destruction in the liver of rabbits by cavitation. Several shortcomings appeared, however, because of the technology used to produce cavitation: (1) the lesions obtained were poorly focused; (2) the need for an intravascular infusion of gas bubbles to enhance cavitation was invasive and difficult; (3) bubbles were not small enough to diffuse into liver capillaries and subsequently, bubbles trapped in the portal spaces generated mostly portal and periporal lesions, while the lobule was generally not affected. In this study we describe reproducible and well focused lesions in the liver by an extracorporeal approach, without the need to inject bubbles in the circulation. Toxicity was not serious; there were no deaths; biological side effects were constantly noted, but always transitory and with complete recovery after three weeks. There were no perforations of surrounding structures, intra-abdominal haemorrhages or thromboses of large vessels. These improvements were permitted by the small size and constant positioning of the focal spot (as tested in vitro in our laboratory by hydrophone pressure recordings (data not provided), and mostly by the effect of the initial negative half cycle of the pressure tracing (expansion wave). High intensity ultrasound produces cytotoxicity through essentially hyperthermia and cavitation; heating is certainly the dominant mechanism of tissue destruction at intensities below 1000 W/cm² and exposure times longer than one second; several research groups, as well as ours, are trying to develop devices for extracorporeal tumour treatment based on focused, high intensity necrosis induc-
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ing hyperthermia.20–23 In this study, it can be assumed that hyperthermia plays a very marginal part, if any, and that almost exclusively caviation participates in the creation of tissue necrosis.

Cytotoxicity such as seen here is probably thus both direct, by changes of cell and nuclear membranes, cytoskeleton, and mitochondria24 and indirect through microvascular disruption4 and production of free radicals.25 It is very difficult to assess accurately the occurrence of caviation in vivo25; however, the hyperechogenic spot induced by each burst at the focal point in the liver, as seen by ultrasound monitoring, is the landmark of transitory bubble production – that is, of caviation.

Another advantage of this technology is the reproductiveness of the lesion in a given model. A perfect alignment of individual lesions, however, was not seen; this is partly because of physiology (respiration movements) and also because of acoustic phenomena of diffraction, refraction, and interactions with the blood flow, which are not well understood in living media. A discrepancy was noted between the desired volume of tissue necrosis and the lesion seen; this is partly explained by respiratory movements, because shots were not breath monitored; caviation can also be created early ahead of the focus, thus extending the lesion beyond the desired volume.26 Testing the role of the distance between points in the target clearly showed that this parameter influenced the density of tissue ablation; the absence of increase in tissue ablation for a distance shorter than the focal spot diameter may be a result of caviation bubbles occurring when a lesion was created very close to the next one (1 mm distant), thus preventing the creation of the next lesion. The actual amount of tissue ablation, however, included a rim of hepatocytes morphologically unchanged but functionally affected, as shown by quantitative microscopy with PAS staining.

Several other parameters might influence the quality of tissue ablation, which could not be investigated with this experimental model: in the liver for instance, the quality and quantity of lesions will be mostly influenced by the acoustic attenuation of the tissue as a function of depth.27 To study easily the conditions of tissue ablation deeper than 30 mm from the skin would need bigger animal models. Modulations of the amplitude of the acoustic wave (its negative as well as its positive components), of the peak positive and negative pressures of the burst frequency might prove important for this factor. The histological lesion induced in the liver by caviation is a necrosis entailing both microvascular lesions and direct parenchymal cell lysis. In contrast with our initial study, tissue ablation here affected the lobule as well as the portal space. The lesions were sharply defined, and the surrounding tissue seemed unaffected. The tissue effect was immediate and radical, and seemed to generate a repair process stimulating both collagenic fibrosis and hepatic regeneration. Indeed, tissue destruction is temporary and not all cells are killed by caviation, suggesting that once applied to tumours, the remaining living cancer cells would continue growing. This

important point can be considered in several ways: firstly, microvascular damage results in ischaemia and fibrosis; it can thus be expected that cells in the focal area continue to die after shockwave is given, and that tumour growth rate would seriously be affected, as has already been shown in vitro28 and in vivo.29 Secondly, the repetition of treatment sessions will probably complete and reinforce therapeutic effects. Thirdly, caviation, as a local treatment, should probably be part of combined therapeutic schedules including chemotherapy or immunomodulators for instance. Several recent studies conducted by various laboratories have shown encouraging results.30–34

The development of a caviation based device for the local treatment of liver cancer in humans is readily conceivable: it could be an alternative to surgery in certain cases of liver cancers. We are now investigating the effects of caviation in a liver cancer model in the rabbit.

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6 Miller DL. A review of the ultrasonic bioeffects of micronsona.


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