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Accessibility
Objectives
Thalamotomy of the ventral intermediate nucleus (VIM) is effective in alleviating medication resistant tremor in patients with essential tremor (ET) and Parkinson’s disease (PD). MRI guided Focused Ultrasound (MRgFUS) is an innovative technology that enables non-invasive thalamotomy via thermal ablation.

Methods
Thirty eight ET and PD patients with severe medication resistant tremor underwent MRgFUS underwent unilateral VIM thalamotomy using MRgFUS. Effect was evaluated using clinical Rating Scale of Tremor (CRST) in ET patients and Unified PD Rating Scale motor part (UPDRS) in PD patients. Quality of life was assessed by Quality of life in ET Questionnaire (QUEST) and PD Questionnaire (PDQ-39).

Results
Tremor stopped in the treated hand in 37 patients immediately following the treatment. In one patients tremor was modified but not abolished. At one month post-treatment, the ET patients’ CRST score decreased from 38.6 ± 12.0 to 9.3 ± 7.7 (p < 0.001) and QUEST scores decreased from 44.8 ± 17.8 to 13.1 ± 15.9 (p < 0.001). In PD patients UPDRS-motor part decreased from 26.2 ± 8.7 to 16.3 ± 11.0 (p = 0.00087) and PDQ39 decreased from 40.8 ± 18.2 to 26.5 ± 15.1 (p = 0.027). During follow up of 1-24 months (mean 10.9 ± 8.1 months) tremor reappeared in seven of the patients, but in all but three, to a lesser degree than before the procedure.

Adverse events that transiently occurred during sonication included: Headache (n = 11), short lasting vertigo (n = 17) and dizziness (n = 4), nausea (n = 4), burning scalp sensation (n = 3), vomiting (n = 3) and lip paresthesia (n = 2). Adverse events that lasted after the procedure included gait ataxia (n = 5), unsteady feeling when walking (n = 4) unilateral taste disturbances (n = 3) and hand ataxia (n = 3). All adverse events were transient and none lasted beyond 3 months.

Conclusions
MRgFUS VIM thalamotomy to relieve medication resistant tremor was safe and effective in ET, and PD. Current results emphasize its low adverse events profile and high efficacy in treating tremor. Large randomized studies are needed to assess prolonged efficacy and safety.

Objectives
Essential Tremor (ET) is one of the most common neurologic conditions, and conservative measures are frequently suboptimal. Recent data from a multi-institution, randomized controlled clinical trial demonstrated that Magnetic Resonance-guided Focused Ultrasound (MRgFUS) thalamotomy improves upper limb tremor in medically refractory ET. This study assesses the cost-effectiveness of this novel therapy in comparison to existing procedural options.

Methods
PubMed and Cochrane Library searches were performed for studies of MRgFUS, Deep Brain Stimulation (DBS), and Stereotactic Radiosurgery (SRS) for ET. Pre- and post-operative tremor-related disability scores were collected from 32 studies involving 83 MRgFUS, 615 DBS, and 260 SRS cases. Utility (defined as percent change in functional disability) was calculated, and Medicare reimbursements were collected as a proxy for societal cost – costs of MRgFUS for ET were derived from a combination of available costs of approved indications and SRS costs where appropriate. A decision and cost-effectiveness analysis was then constructed, implementing meta-analytic techniques.

Results
MRgFUS thalamotomy resulted in significantly higher utility scores compared with DBS and SRS based on estimates of Medicare reimbursement (p < 0.001). MRgFUS was also the most inexpensive procedure out of the three (p < 0.001).

Conclusions
Preliminary experience with MRgFUS for ET suggests that this novel therapeutic may be more effective than available alternatives and potentially less costly for society. It thus will likely “dominate” DBS and SRS as a more cost-effective option for medically refractory ET. Our findings support further investigation of MRgFUS for ET and broad adoption.
Objectives

The ventral intermediate nucleus (VIM) is not visible on conventional Magnetic Resonance Imaging (MRI). A novel method for tractography-based VIM identification has recently been described. We report the short-term clinical results of prospective VIM targeting with tractography in a cohort of patients undergoing Focused Ultrasound thalamotomy.

Methods

All patients underwent structural and diffusion weighted imaging (60 diffusion directions, 2 mm isovoxel) with 3 Tesla MRI scanner (Philips Ingenia CX). The images were processed using streamline tractography (Stealth Viz, Medtronic Inc.). The lateral and posterior borders of VIM were defined by tracking the pyramidal tract and medial lemniscus respectively. A VIM region of interest (ROI) was placed 3 mm away from these borders (Figs. 1, 2 and 3). The structural connectivity of this VIM ROI was confirmed to the motor cortex (M1) and cerebellum. The coordinates of tractography-based VIM in relation to posterior commissure were noted for surgical targeting. The parameters analyzed include a clinical tremor scale (pre-, intraoperative, and post operative), operative time, and number of sonications.

Results

Tractography-based VIM targeting was successful in 7 out of 8 patients. The coordinates of tractography-based VIM were significantly different from the standard coordinates (3-D distance 3.9 ± 2.4 mm). Therapeutic sonication (>55 °C temperature, 10 seconds) at the tractography target resulted in >50 % tremor improvement with intraoperative objective tremor assessment without any motor or sensory side-effects. The mean operative time was 78 ± 3.3 minutes with 12.8 ± 3.9 average sonications. Overall the tremor scores significantly improved one month after surgery (preop CRST total 62.1 ± 15.5 versus 30.3 ± 14.1, two tailed t-test p = 0.006). None of the patients experienced sensory deficits or motor weakness during follow-up.

Conclusions

We report that prospective tractography-based VIM targeting is safe and feasible. The short-term clinical results are satisfactory. Long-term tremor efficacy outcomes are desirable to further assess the usefulness of this technique.

Targeted delivery of brain-penetrating non-viral GDNF gene vectors to the striatum with MRI-guided Focused Ultrasound reverses neurodegeneration in a Parkinson’s disease model

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Objectives

Parkinson’s disease (PD) is characterized by the degeneration of dopaminergic neurons in the motor control pathways of the brain. Gene therapy using glial cell derived neurotrophic factor (GDNF) has shown some limited promise for treating PD; however, we hypothesize that outcomes could be further improved by enhancing gene vector distribution. We previously developed a gene therapy approach that entails delivering systemically administered non-viral gene-bearing nanoparticles (BPN) across the Blood-Brain Barrier with MRI-guided Focused Ultrasound...
Focused ultrasound facilitated gene delivery for neuro-restoration in Parkinson’s disease mice

Shutao Wang, Ouyierni Olumolade, Tara Kugelman, Vernice Jackson-Lewis, Maria Eleni (Marilena) Karakatsani, Yang Han, Serge Przedborski, Elisa Konofagou

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**Objectives**
Not released for publication

**Methods**
Not released for publication

**Results**
Not released for publication

**Conclusions**
Not released for publication

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**A6**
MRI-g-FUS for the treatment of Alzheimer’s disease

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**Objectives**
Over the past ten years consistent effort has been put forward at the University of Toronto to develop Focused Ultrasound methods for the treatment of AD. This talk will review the progress made so far.

**Methods**
The first studies demonstrated safe antibody delivery in AD mouse model with significant reduction in the plaque load. A follow up studies with two-photon microscopy showed that blood vessels with plaque deposits showed a different type of opening than vessels in normal brain but large molecule delivery into the brain was still possible in these animals. Another study demonstrated that plaque reduction can be achieved by just opening the BBB with microbubbles.

**Results**
The histology revealed stimulation of neurogenesis. Multiple treatments of old mice resulted in memory rescue without any observable side-effects. A follow up study demonstrated that this neurogenesis was not induced with exposures that did not cause observable BBB opening even with the presence of the microbubbles. An ongoing study in large animals has shown that half-brain BBB opening can be safely and repeatable performed indicating the feasibility of clinical translation.

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**A7**
Scanning Focused Ultrasound disruption of the blood-brain barrier as an Alzheimer’s disease therapy

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**Objectives**
Alzheimer’s disease is the most common form of dementia. Pathological abnormalities in the Alzheimer’s disease brain includes the presence of amyloid-beta plaques, hyperphosphorylation and intracellular aggregation of tau and synaptic degeneration. Focused ultrasound combined with intravenous injection of microbubbles has been shown to reversibly open the blood-brain-barrier (BBB). By moving the focus in scanning mode we are able to open the BBB throughout the brain of a mouse. Here we tested the effects of repeated scanning ultrasound (SUS) in APP23 amyloid plaque-bearing mice, pRS tau mice and wild-type mice to determine the effects of SUS on amyloid, tau and dendritic spines.

**Methods**
The device used was the Therapy Imaging Probe System (TIPS, Philips Research), which has an eight-element annular array transducer with a focal length of 80 mm a radius of curvature of 80 mm, a 33 mm central opening, and a motorized 3D positioning system. The focus 6 dB size was 1.5 mm x 1.5 mm x 12 mm at 1 MHz. Settings that were applied were 1 MHz centre frequency, 0.7 MPa peak rarefractional pressure applied outside the skull, 10 Hz pulse repetition frequency, 10 ms pulse length and 10 % duty cycle immediately after retroorbital injection of i-house made microbubbles. APP23...
mice that accumulate amyloid beta, pRS mice that overexpress FTD-mutant tau, and wild-type C57Bl/6 mice were treated weekly by scanning ultrasound (SUS) for periods of 4 to 7 weeks.

Results
In APP23 mice we used repeated scanning ultrasound (SUS) treatments of the mouse brain to remove amyloid-beta. Spinning disk confocal microscopy revealed extensive internalization of Abeta into the lysosomes of activated microglia in mouse brains subjected to SUS. Plaque burden was reduced in SUS-treated AD mice compared to sham-treated animals. Treated AD mice also displayed improved performance on three memory tasks.

In pRS mice we investigated the efficacy of a novel tau isoform-specific single chain antibody fragment, RNX, delivered by passive immunization in the P301L human tau transgenic pRS mouse model. When administration of RNX was combined with scanning ultrasound (SUS), RNX delivery into the brain and uptake by neurons was markedly increased, as were reductions in tau phosphorylation and anxiety-like behavior.

In wild-type mice we investigated the effects of SUS on neuronal excitability and morphology. We performed patch-clamp recordings from hippocampal CA1 pyramidal neurons in wild-type mice 2 and 24 hours after a single SUS treatment, and one-week and three months after six weekly SUS treatments. No change in CA1 neuronal excitability was observed compared to sham-treated neurons at any time-point. Multiple SUS treatments had the effect of preventing the loss of CA1 synapses that occurred in sham-treated neurons.

Conclusions
We show that scanning Focused Ultrasound disruption of the BBB has multiple biological effects in the brain which make it an attractive candidate for an Alzheimer disease therapy. SUS reduced plaque burden and amyloid-beta levels in APP23 amyloid mice, through activation of microglia, and improved performance on tests of memory function. In pRS tau mice SUS alone reduced hyperphosphorylation of tau, and enhanced the delivery of anti-tau antibodies resulting in improved reductions in pathology and behavioral abnormalities. In wild-type mice SUS was shown to have no effect on the firing of hippocampal neurons or their morphology, but prevented spine loss at 3 months after six weekly SUS treatments. If these effects on Abeta and tau pathology, and dendritic morphology are recapitulated in human patients SUS may emerge as a promising AD therapy.

A8 Scanning ultrasound as a treatment tool of proteinopathies including Alzheimer's disease
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Objectives
Neurological disorders constitute a substantial social and economic burden, as they cause considerable ill health but few direct deaths. Treatment strategies for neurodegenerative diseases are hampered by the fact that the Blood-Brain Barrier (BBB) establishes an efficient barrier for therapeutic agents (Leinenga et al., Nature Reviews Neurology 2016). We have recently shown that scanning ultrasound (SUS) allows microglial-mediated clearance of extracellularly deposited amyloid-beta in APP mutant APP23 mice and restores memory functions in three cognitive tests to wild-type levels, in the absence of overt damage to the brain (Leinenga and Götz, Science Translational Medicine 2015) However, it had not been determined whether SUS treatment reduces the intracellular tau pathology that together with amyloid deposition characterizes Alzheimer's disease.

Methods
We investigated the efficacy of a novel tau-specific single chain antibody fragment, delivered by passive immunization in the human tau transgenic pRS mouse model, a model of the tau pathology of Alzheimer's disease (Götz et al., Science 2001). To further assess the efficacy and drug-delivering ability of SUS, we established four experimental groups, using the novel anti-tau antibody that was injected weekly over four weeks, either on its own, or together with SUS. A third group used SUS only, and a fourth was the anaesthesia control group. The mice were analysed on the elevated plus maze, histologically and biochemically. Furthermore, uptake of the antibody by the brain was determined using fluorescently labelled single-chain antibody fragments.

Results
A histological and biochemical analysis of the pRS tau transgenic mice revealed that SUS as well as the employed antibody ameliorated the tau pathology that characterizes the pRS mice. In addition, the anxiety-like behaviour that characterizes pRS mice was significantly reduced. We furthermore found enhanced delivery of the antibody using SUS yielding a synergistic therapeutic effect as determined by histology and using the elevated plus maze.

Conclusions
Our study suggests that SUS is a method that benefits diseases with protein aggregates more generally, whether they are intra- or extracellular. The therapeutic delivery combined with SUS could offer significant clinical benefits for the treatment of patients with Alzheimer's disease and related tauopathies. Considering that the yearly costs of passive immunotherapy for AD is expected to exceed $25,000 per patient, combining SUS with antibody delivery could drastically reduce these costs.

A9 Enhancement of FUS mediated delivery of stem cells to the brain
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Objectives
FUS mediated Blood-Brain Barrier disruption (BBBD) can enable even large therapeutics such as stem cells to enter brain from the bloodstream and could be a major advance in cell delivery over current invasive methods of brain injection. The efficiency of cellular entry after FUS mediated BBBD alone however is low. We hypothesized that this process could be enhanced by combining it with a complementary strategy termed magnetic targeting. Stem cells can be safely loaded with super-paramagnetic iron oxide nanoparticles (SPION) in culture, allowing cells to be attracted by an external magnet. Our previous study showed SPION loaded stem cells to have enhanced brain retention near a magnet on the skull in a rat model of traumatic brain injury, where BBBD also occurs. The goal of our current project was to determine if magnetic attraction of SPION loaded stem cells would also enhance their delivery to brain after FUS mediated BBBD.

Methods
With a small animal MRI guided FUS device (Image Guided Therapy, IGT and 7 T Bruker MRI), we sonicated young adult rats (~120 g) with both radiologic (enhancement of the target region with gadolinium contrast) and histologic (staining with Evans' blue dye) evidence of BBBD, without tissue damage or hemorrhage. Confirmation of the cells within brain as those injected was performed by staining with Perl's reagent for iron and by immuno-histochemistry with a human specific antigen. The procedure was then combined with the application of a powerful magnet to the head directly after IV injection of hNPCs.

Results
With BBBD alone human neuro-progenitor cells (hNPCs) loaded with SPION were observed in rat brain after intravenous (IV) injection directly after sonication only within the treated regions. To demonstrate the effect of magnetic attraction, we injected equal numbers of SPION and non-SPION labeled cells, where each cell type was labeled with a different fluorophore. In animals that had FUS mediated BBBD...
followed by a magnet applied to the head, significantly greater numbers of SPION labeled cells were observed compared to the non-labeled cells. This result was most pronounced in regions of the brain close to the skull (cerebral cortex) and magnet surface. More powerful magnets including magnetic arrays resulted in more effective retention of SPION labeled cells in even deeper brain regions such as the striatum. There, 90% of hNPCs observed contained SPIONs compared to 60-70% with a less powerful magnet.

**Conclusions**

These results demonstrate that the use of magnetic attraction can substantially enhance delivery of stem cells after BBBD. In prior published work, stem cells were delivered to brain after FUS mediated BBBD using cells injected directly into the carotid artery. In an effort to accomplish this goal in a safer and less invasive manner, our study utilized IV cell injection (tail vein), supporting the view that the combination of FUS mediated BBBD and magnetic attraction can allow stem cells to enter brain with a minimally invasive strategy.

**A10**

Fluorescent lipid microbubbles for targeted brain drug delivery through the Focused Ultrasound-induced blood-brain barrier opening in vivo

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**Objectives**

Focused ultrasound (FUS) in the presence of lipid microbubbles can induce non-invasive, transient and reversible Blood-Brain Barrier (BBB) opening. This study entailed assessment of the feasibility of fluorescently loaded microbubbles, labeled with the fluorophore 5-dodecanoylaminfluorescein (C-12), as a vector for targeted brain drug delivery. Compared to prior studies by our group, where fluorescently-labeled dextran were co-administered with microbubbles, this new methodology improves the safety and allows a more targeted drug delivery with potentially lower toxicity, avoiding systemic exposure. The main objective was thus to determine feasibility and safety of using the loaded microbubbles as carriers towards targeted brain drug delivery with simultaneous cavitation monitoring.

**Methods**

A spherical, single-element, FUS transducer (center frequency 1.5 MHz) was used. A pulse-echo transducer (center frequency 10 MHz), confocally mounted at the center of the FUS transducer, was utilized for passive cavitation detection (PCD). FUS (pulse length 10,000 cycles; pulse repetition frequency 5 Hz; duration 5 minutes; acoustic pressure 450-750 kPa) targeted mouse brains in vivo, in combination with fluorescent microbubbles for C-12 delivery, which was evaluated by in vivo transcranial PCD, through the quantification of inertial (ICD) and stable harmonic (SCDh) and ultraharmonic (SCDu) cavitation doses at 30, 60 and 300 s; together with ex vivo fluorescence imaging. The BBB opening was verified using in vivo T1-w MRI showing BBB opening at pressures (a) 450, (b) 600 and (c) 750 kPa. Fluorescence delivery (green) and DAPI (blue) in two horizontal sections of mouse brains sonicated at (d) 600 and (e) 750 kPa.

**Fig. 5 (abstract A10).** BBB opening and fluorescence delivery: T1-w MRI showing BBB opening at pressures (a) 450, (b) 600 and (c) 750 kPa. Fluorescence delivery (green) and DAPI (blue) in two horizontal sections of mouse brains sonicated at (d) 600 and (e) 750 kPa.

**A11**

Ultrasound-mediated delivery of gadolinium and fluorescent-labelled liposomes through the Blood-Brain Barrier

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**Objectives**

The main objectives of this study were: 1) to examine whether gadolinium and fluorescent labelled liposomes can extravasate into the brain parenchyma after ultrasound mediated Blood-Brain Barrier disruptions; and 2) to test whether extravasated liposomes were size dependent or not. The liposomes were labelled with gadolinium (Gd) and fluorophore, thus enabling detection of extravasated liposomes via MRI in vivo and fluorescence methods in tissue, respectively.

**Methods**

Liposomes labelled with gadolinium and fluorophore were prepared using lipid film hydration and extrusion to two different sizes: ~70-85 nm and ~130-150 nm. Animals were divided into two different groups based on the use of particle sizes; group A (~70-85 nm) and group B (~130-150 nm). Focused ultrasound mediated Blood-Brain Barrier disruption (BBBD) was produced in one hemisphere in 15 mice survived up to one week after opening. Iba-1 immunohistochemistry also showed microglial activation.

**Conclusions**

FUS was applied in conjunction with fluorescent microbubbles and, for the first time, the existence of CD thresholds for assessing successful drug delivery was defined. For CD above these thresholds, significant fluorescent enhancement was observed, demonstrating C-12 targeted delivery. One week after sonication, edema was cleared out but microglial activation was observed in certain cases. Therefore, this study indicates the feasibility and safety of a new methodology of FUS-induced BBB opening for targeted albeit potentially riskier brain drug delivery and provides a platform for predicting successful delivery via PCD.
mice. Particles were injected before sonication. Sonications (0.69 MHz at 0.42 MPa) were performed in two locations combine with Definity (10 μl/kg). Acoustic emissions were recorded during FUS. T1-weighted contrast enhanced and T2*-weighted MRI were used to confirm Gd leakage and damage detection respectively. Mice were euthanized 5-24 hours after FUS and post-process for fluorescence measurement.

Results
In T1-weighted contrast enhanced MRI, gadolinium-leakage was able to detect on sonicated area at 5-24 after FUS but not on non-sonicated area (control). Detection of fluorescence signal from brain tissue homogenates confirm the liposomal particles extravasation on sonicated locations. On group A, gadolinium and fluorescence signal intensities on sonicated locations were increased by 26 % and 62 % respectively as compared with control and signal enhancement were statistically significant compared with control (p = 0.017 and p = 0.02 respectively, two-tailed, paired ttest). On group B, gadolinium and fluorescence signal intensities on sonicated locations were increased by 24 % and 40 % respectively as compared with control. Comparison of fluorescence signal intensities between two groups on sonicated location was statically significant whereas it was not significant on controls (p < 0.05 and p = 0.07 respectively, one-tailed unpaired ttest).

Conclusions
Overall, this work demonstrates that ultrasound can deliver up to ~150 nm liposomes that labelled with gadolinium and fluorophore through the Blood-Brain Barrier. The results indicate that the extravasation of liposomes were size dependent.

A12
Sterile inflammatory response (SIR) in the brain following exposure to low intensity pulsed Focused Ultrasound and microbubble infusion
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Objectives
Magnetic Resonance Imaging (MRI)-guided pulsed Focused Ultrasound (pFUS) in combination with systemic injection of microbubbles (MB) is being advocated to increase drug or gene delivery by causing localized Blood-Brain Barrier (BBB) disruption (D). The objective of this study is to investigate the molecular and cellular responses following pFUS + MB associated with BBBD in the rat brain.

Methods
Female Sprague Dawley rats (<200 g) were sonicated at 0.3 MPa acoustic pressures with 10 ms burst length and 1 % duty cycle (9 focal points, 120 sec/9 focal points) using a single-element spherical FUS transducer (SB9,636 kHz; FUS Instruments). 100 μl Optison™ MB (GE Healthcare) was administered intravenously. Gadofosveset-enhanced T1w images were obtained with a 3.0 T MRI (Phillips). Proteomic and mRNA expression in the brain following pFUS + MB were analyzed with ELISA, Western blot, quantitative real-time PCR or immuno-fluorescent staining (Fig. 6a). Proteomics were normalized to sham and statistical analysis was performed by one-way ANOVA corrected for multiple comparisons. Rats were also injected with 8 mg/kg Rhodamine encapsulated magnetic polymers (MicroTRACK™; Bio-Pal) 3 days prior to sonication to label splenic macrophages (CD68) to monitor tropism to the brain.

Results
Post contrast T1w MRI and histology showed open BBB without evidence of microhemorrhage. Within 5 minutes following sonication, increased expression of pro-inflammatory and anti-inflammatory cytokines, chemokines and trophic factors (CCTF) was detected in the parenchyma lasting up to 24 hours (Fig. 6b), increases in heat shock protein 70 (HSP70), tumor necrosis alpha (TNFα), and interleukin (IL) 1a, 1b and 18 consistent with damage associated molecular patterns (DAMP1) and activation of nuclear factor kappa-light-chain-enhancer of activated B cells (NFκB) inflammatory pathways were observed with SIR to injury2 (Fig. 6b). NFκB pathway-related gene activation along with anti-apoptotic genes, immune cell chemotractants, selectins, and cell adhesion molecules showed significant (>2fold) increases in mRNA expression (Fig. 6c). Histological analysis showed significant increases (p < 0.05) in the following: number of TUNEL positive cells within 6 hrs, GFAP and Iba1 staining for activated astrocytes and microglia (1-24 hrs) and increase of ICAM up to 24 hrs post sonication. We also detected a >4 fold greater (p < 0.05) CD68 positive cells on day 6 post sonication containing intracellular fluorescent beads within the pFUS + MB treated hemisphere compared to contralateral hemisphere.

Conclusions
The temporal molecular response to pFUS + MB is indicative of SIR2 originating from the parenchyma. The pattern of cytokines immediately after pFUS + MB is initiated by cellular release of DAMPs and TNFα observed with mild cerebral trauma or ischemia [1,2]. Increases in monocyte chemotactrant protein (MCP-1), vascular endothelial growth factors (VEGF), stromal derived factor 1 (SDF-1), erythropoietin (EPO) and brain derived neurotrophic factor (BDNF) are associated with BBBD stimulating angiogenesis, neurogenesis and stem cell migration observed with ischemia and trauma. These results indicate that pFUS + MB rapidly affects the cerebral vasculature as evident by BBBD in addition to the shockwave from MB collapse that induces mild stress within various cellular elements in the parenchyma inducing a SIR.

References

A13
Volumetric MR thermometry in a clinical transcranial MR guided Focused Ultrasound system
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Objectives
Transcranial MR-guided Focused Ultrasound (tcMRgFUS) applied within a small central brain volume has achieved excellent outcomes in treatment of movement disorders (Elia NEJM 2013). Although
transducers used in tcMRgFUS have been designed with large apertures to spread the beam energy over as much skull as possible to reduce skull and cortex heating, currently used MR temperature imaging (MRTI) methods cannot monitor the temperature increase over the entire insonified brain volume. Instead, temperatures are typically measured in one 2D slice, leaving the majority of the insonified brain volume unmonitored. We have previously developed and published methods to achieve fully 3D MRTI covering the entire insonified brain with good spatial and temporal resolution (Todd MRM 2009/2010) but the techniques have not been evaluated on clinical tcMRgFUS systems. In this work-in-progress study we demonstrate the value of volumetric MRTI with two different pulse sequences applied during heating on a clinical tcMRgFUS system.

**Methods**

PRF MRTI was performed with a product 3D gradient recalled echo (GRE) pulse sequence and a custom-implemented 3D GRE segmented echo planar imaging (EPI) pulse sequence on a 3 T MRI (Discovery 750 T, GEMS). tcFUS heatings were performed in an ex vivo human skull filled with tissue-mimicking gel (ATS Laboratories) in a clinical tcMRgFUS system (ExAblate Neuro, Insightec).

Pulse sequences parameters are listed in Table 1. All data were zero filled interpolated (ZFI) to 1-mm isotropic spacing (GRE data additionally ZFI to 0.5-mm spacing). The skull was physically positioned in the FUS system, and the focus electronically steered, to target deep brain-structures located outside the normal treatment envelope. FUS sonications were applied at 940 W for 30/60 s while imaging with the GRE/EPI sequences, respectively. The EPI sequence used an echotrain length of 16 with bi-polar readout gradient, sampling 192 phase encodings in each direction, so that full “positive only” and “negative only” images could be reconstructed.

K-space data were retrospectively down sampled by a factor of $R = 4$ and $8$ (split into multiple time-frames without throwing any data away) for the GRE and EPI experiments, respectively, giving acquisition times of 3.6 and 7.8 s, and reconstructed with a temporally constrained reconstruction algorithm (Todd MRM 2009).

**Results**

Figure 7 shows three orthogonal views of temperature maps overlaid on magnitude images in the GRE experiment. Attempting to focus this far outside the normal treatment envelope results in severe near- and far-field heating. Heating on both the cortex and in the far-field along the petrous bone is visible. In Fig. 8 the temperature evolution at the focal spot is compared to that near the petrous bone. The gel near the bone shows a delayed and greater maximum temperature compared to the focus.

Three orthogonal views of the larger FOV in the segmented EPI experiment are shown in Fig. 9. Heating along large parts of the cortex can be seen. In the underlying magnitude images it can be seen that the EPI sequence experiences more artifacts than the GRE sequence. The focal spot position (evaluated as the temperature center-of-mass) was tracked as a function of time during the heating in the GRE data ZFI to 0.5 mm (data not shown). The focal spot did not experience any shift to within the finer ZFI spacing.

**Conclusions**

This study shows that volumetric thermometry over a FOV covering the focal spot and the skull base can be achieved with readily available pulse sequences. With custom implemented pulse sequences the fully insonified FOV (from skull cap to skull base) can be covered. Even though the reconstruction is done retrospectively, the described methods are valuable as research tools in e.g. treatment envelope evaluations. By utilizing 3D imaging ZFI can be performed in all directions to minimize partial volume effects, and very accurate dynamic focal spot localization can be performed.

Future studies will compare the accuracy and precision of the described methods to standard 2D MR thermometry. Experiments comparing 3D MRTI with fiber optic probe measurements at potential target positions outside the currently available treatment envelope will also be performed (Monteith JNS 2016).

**Table 1 (abstract A13).** MR scan parameters. TR – Repetition time, TE – Echo time, FA – Flip angle, BW – Bandwidth (readout), FOV – Field of view, Res – Resolution, Tacq – Acquisition time (before subsampling)

<table>
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<th></th>
<th>TR (ms)</th>
<th>TE (ms)</th>
<th>FA (deg)</th>
<th>BW (kHz)</th>
<th>FOV (mm)</th>
<th>Matrix</th>
<th>Res. (mm)</th>
<th>Tacq (s)</th>
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<td>1.25x2.50x3.00</td>
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<td>30</td>
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**Fig. 7 (abstract A13).** Three orthogonal views of GRE temperature maps overlaid on magnitude image

**Fig. 8 (abstract A13).** Temporal evolution of heating comparing focal spot to far-field next to petrous bone
Multi-echo MR thermometry compared to single echo MR thermometry in the treatment of essential tremor
Kim Butts Pauly, Mike Marx, Pejman Ghanouni
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Objectives
The choice of receive bandwidth in MR thermometry acquisitions needs to balance two competing choices. Low bandwidths improve SNR, while high bandwidths reduce spatial shift of the hotspot due to the temperature off-resonance. The low bandwidth MR thermometry sequence in use in essential tremor MRgFUS treatments required repeated swapping of phase and frequency directions since the location of the hotspot could only be trusted in the phase encode direction.

A solution is to use multiple high bandwidth acquisitions, which when averaged, regain much of the SNR of the lower bandwidth image. Such a multiecho sequence has recently become available for clinical use. The purpose of this work was to compare the performance of the multi-echo (ME) MR thermometry to the single echo (SE) MR thermometry in clinical treatments of essential tremor.

Methods
Fifteen patients were treated for essential tremor, 12 with only single echo thermometry (TE and BW/pix = 12.8 and 44), 2 with only multi-echo thermometry (TE and BW/pix = 3, 8, 13, 18, 22 and 278), and 1 with both sequences. All other image parameters remained the same, including TR = 100. All thermometry images were processed offline in Matlab with a single baseline subtraction (α = -0.00909), followed by referenceless processing for constant and linear terms. The multi-echo thermometry was further processed with a phase unwrapping algorithm in the TE dimension. The multiple echoes were then combined with a weighting by the square of the temperature SNR of each image.

Results
The decrease in sampling time for the ME thermometry dictates a theoretical decrease in temperature SNR of only 11 % due to the decrease in sampling time alone. In the one patient that had both sequences in the same scan planes for two sonications, there was a 9 % decrease in SNR in the ME thermometry as measured in the frame used for the referenceless processing, comparing well with theory.

A review of all sonications with ME thermometry when phase encoding was S/I revealed much reduced artifacts over SE thermometry with phase encoding in the S/I direction. Example images are provided in Fig. 10. The region of interest measurement indicated a 30 % improvement in temperature SNR for the ME over the single echo, due to this artifact reduction.

The spatial shifts should theoretically be reduced by a factor of 6.3 in the ME thermometry, as compared to the SE thermometry, due to the increase in receive BW. For example, a 23 °C temperature rise would result in a 0.6 mm shift with the single echo, and only a 0.1 mm shift with the multi-echo.

Conclusions
When phase encoding is S/I, multi-echo thermometry is superior to single echo thermometry due to a reduction in ghosting artifacts and spatial shifts. Alignment of the focal spot should not require swapping phase and frequency directions. Future work will include a prospective study to verify this. In the other directions, ME thermometry is comparable to SE thermometry with the reduction of spatial shifts coming with a loss of temperature SNR of about 10 %.

High-resolution whole-brain MR thermometry with a 3D EPI stack-of-stars pulse sequence
Sumeeth Jonathan, William Grissom
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Objectives
Real-time whole-brain MR thermometry is needed for transcranial MR-guided Focused Ultrasound to accurately track rapid heating at the focus and to monitor for unsafe heating in the near- and far-field. Previous efforts to meet this need have been based on 3D segmented echo planar imaging (EPI) and spiral MR readouts (Fielden et al., 2014). Here, we propose a 3D EPI stack-of-stars temperature mapping pulse sequence that enables greater volume coverage than has been reported with previous approaches. The sequence allows flexible adjustment of the scan acceleration factor and does not require a high density of receive coils for high frame rate thermometry.

Methods
Readout Trajectory: Fig. 11 shows the proposed readout trajectory. Each TR comprises a 2D EPI plane that is frequency-encoded in the axial (x-y) and phase-encoded in the slice (z) dimensions. The plane is rotated between TRs by 111.25 degrees to sample a golden angle...
stack-of-stars k-space. The sequence was implemented on a Philips Achieva 3 Tesla scanner with parameters: field-of-view = 28.0 x 28.0 x 11.9 cm, 43 slices, in-plane resolution = 1.50 x 1.50 x 2.75 mm, 17 ms TE/45 ms TR, 14.5° flip angle, spectrally-selective fat suppression.

**In vivo** Experiment: To confirm the achievable brain volume coverage and image quality, with local IRB approval, brain images in a healthy volunteer were acquired without heating using the pulse sequence with one, two, and three interleaved shots and with an 8-channel receive coil array. Fully-sampled images were reconstructed using the density-compensated conjugate gradient method.

**Phantom Heating Experiment:** A tissue-mimicking gel phantom was sonicated using a clinical MR-HIFU system (Philips Sonalleve) operated at 1.2 MHz and 80 W for 12 s, while scanning with the proposed sequence with one shot and with 5 abdominal receive coils. Temperature maps were reconstructed in one second increments using the k-space hybrid method at three scan acceleration factors.

**Results**

**In vivo Results:** Distortions and signal dropouts caused by off resonance are visible in the sagittal and lower axial single-shot brain images in Fig. 12 (white arrows). The distortions can be reduced by increasing the number of EPI shots per angle, which increases the pixel bandwidth in z. An axial slice positioned for monitoring Focused Ultrasound thalamotomy (red arrow) is also shown, which contains no visible distortions at any multishot factor.

**Phantom Heating Results:** Fig. 13a shows reconstructed sagittal and axial temperature maps through the middle of the phantom at peak heat. Significant temperature aliasing does not appear until the scan is accelerated by a factor of 14.1 or a 1 second window width. The hotspot temperature curve plots in Fig. 13b show that the reconstructed temperatures all coincide, though the 14.1x curve appears noisier. Importantly, at all acceleration factors, there is agreement with a hot spot measurement obtained by a 2D multislice Cartesian EPI temperature mapping pulse sequence (7 slices, same in-plane resolution) which was used as a reference standard.

A video of the entire sonication monitored using this pulse sequence can be viewed at https://www.youtube.com/watch?v=z3cVXjkgyLg.

**Conclusions**

A 3D EPI stack-of-stars temperature mapping pulse sequence was proposed and validated against a 2D multislice Cartesian EPI temperature mapping pulse sequence. The sequence enables fine spatiotemporal resolution with large volume coverage, without requiring temporally-regularized reconstruction or a large number of receive coils.

**Objectives**
Not released for publication

**Methods**
Not released for publication

**Results**
Not released for publication

**Conclusions**
Not released for publication

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**A17**

**Tissue stiffness imaging with interleaved multiple-point MR-ARFI**

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*Journal of Therapeutic Ultrasound* 2016, 4(Suppl 1):1A17

**Objectives**
Although Focused Ultrasound (FUS) has the potential to treat a number of pathologies the methods used to guide treatment are still...
limited in the ability to identify effective treatment endpoints. Changes in MRI parameters such as proton density and T1 and T2 relaxation times may result from reversible edema instead of tissue death. While dynamic contrast enhanced and late Gd-enhanced MRI are more specific, Gd contrast cannot be administered multiple times during a procedure to monitor treatment progression. MRI acoustic radiation force impulse imaging (MR-ARFI) can monitor tissue stiffness, but the acquisition is relatively slow, and must be repeated to measure tissue displacement at more than a single point [1]. The purpose of this work was to develop an efficient method to use MR-ARFI to measure tissue displacement volumetrically with an array of points as a first step towards monitoring tissue stiffness changes during MRgFUS procedures.

Methods
A 3D segmented gradient echo (GRE) echo-planar pulse sequence, designed to simultaneously measure tissue displacement with MR-ARFI and the corresponding tissue heating [2–5], was further modified to allow interleaved acquisition of multiple point MR-ARFI (mpMR-ARFI) volumes. Phase change due to displacement was separated from that due to temperature by complex subtraction of an interleaved volume acquired with no FUS applied. Temperature was obtained using the proton resonance frequency method with a referenceless background phase subtraction [6].

Experiments were performed in a gelatin phantom and excised pig brain on 3 T MRI scanners (Siemens Tim Trio and PrismaFit). Pulse sequence parameters were: TR/TE = 73/43 ms, FA = 30°, readout bandwidth = 752 Hz/pixel, echotrain length = 7, FOV = 160x114x55 mm, matrix = 128x91x22 giving voxel dimensions of 1.25x1.25x2.5 mm before zero filled-interpolation. Bipolar motion-encoding gradients (MEG) of 15 ms duration (each bipolar lobe) were applied prior to signal readout. Ultrasound pulses of 50 acoustic watts were applied during the second 15 ms bipolar MEG lobe. Navigator echoes were used to measure and compensate for B0 field drift due to gradient heating caused by high MEG strength and duty cycle. The same navigators echoes can be used for respiratory correction in vivo [7].

Results
Results from a13-point mpMR-ARFI acquisition in a gelatin phantom using the Siemens Tim Trio and MEG = 20 mT/m are given in Fig. 14. An example cropped single slice from 10 of the 13 acquired displacement volumes is shown in Fig. 14a with a cropped slice of the mpMR-ARFI composite of all 13 displacement volumes shown in Fig. 14b. The corresponding temperature increase measured during the mpMR-ARFI acquisition is shown in Fig. 14c and was relatively low (<3 °C). (For all mpMR-ARFI images, point separation is 5 mm).

Figure 15 shows the improved image quality achieved by the phase-navigator correction in mpMR-ARFI measurements obtained using a PrismaFit 3 T MRI scanner with MEG = 40 mT/m. The ghosting artifact in Fig. 15a due to B0 field drift caused by high amplitude (40 mT/m) and duty cycle (40 %) MEG gradients during mpMR-ARFI acquisition. After correction (Fig. 15b) mpMR-ARFI displacement images have negligible artefact.

Results of the displacement measured in excised pig brain using the Siemens PrismaFit 3 T MRI scanner and MEG = 40 mT/m and duty cycle (40 %) MEG gradients during mpMR-ARFI acquisition. After correction (Fig. 15b) mpMR-ARFI displacement images have negligible artefact.

Results of the displacement measured in excised pig brain using the Siemens PrismaFit 3 T MRI scanner and MEG = 40 mT/m and duty cycle (40 %) MEG gradients during mpMR-ARFI acquisition. After correction (Fig. 15b) mpMR-ARFI displacement images have negligible artefact.

Figures 14 (abstract A17). a Individual images, b Composite displacement, c simultaneous heating

Figures 15 (abstract A17). Ghosting (a) is corrected by navigator echoes (b) allowing clean mpMR-ARFI displacement measurement (c,d).

Figures 16 (abstract A17). mpMR-ARFI displacement measurements in an excised pig brain before (a) and after (c) MRgFUS ablation with estimated cumulative dose in CEM (b)

Conclusions
Measuring tissue stiffness (displacement as a function of ultrasound intensity) and changes in displacement before, during, and after the procedure provides the unique potential to remotely palpate the ablated volume to monitor the formation of lesions created with MRgFUS. It will also provide a crucially needed assessment of tissue change in regions of fat where conventional MR thermometry fails.

References


A18
Binary localization of cavitation activity based on harmonic content for transcranial brain therapy
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1Institut Langevin Ondes et Images, ESPCI ParisTech, CNRS UMR 7587, INSERM U979, Paris, France; 2CENIR, ICM, CNRS U7225, INSERM U975, Paris, France
Objectives
Cavitation activity may occur during BBB disruption (due to UCA injections) and also during HIFU treatment due to nucleation under high negative pressure. The corresponding microbubble activity has to be monitored to assess the safety and efficiency of brain treatments by ultrasound. The purpose of this study is to binary discriminate the position of microbubbles, inside or outside the skull, in order to know whether cavitation occurs in the brain of the patient or not.

Binary localization is achieved here by taking advantage of the attenuation properties of the skull. The skull acts indeed as a low pass filter for acoustic signals. Thus we hypothesize that the harmonic content of signals from cavitation events could be used as a binary indicator of their localization: inside or outside the brain case.

Methods
A wideband Passive Cavitation Detector (PCD) recorded the acoustic signals from microbubbles activity.

An in vitro setup (Fig. 17) mimics a BBB opening configuration (contrast agent flow with 0.6 MPa sonication during 10 ms at focus) or a thermal ablation (calf brain sample at focus with 3–4 MPa sonication during 0.3 s at focus). Experiments were performed either with no skull in place, or with human (6 samples) or monkey (1 sample) skull in front of the PCD.

In vivo BBB opening were conducted on macaque (900 kHz, 0.6 MPa at focus) with the same PCD mounted on the monkey head. The spectra are computed from data recorded by the PCD and reveal harmonics, subharmonic and ultraharmonics of the excitation frequency (900 kHz in all cases).

The ratio of a high frequency harmonics over a lower frequency harmonic was then calculated. This ratio is expected to decrease when the acoustic signal from cavitation activity crosses through the skull. In order to achieve binary localization, two types of ratio have been computed: ultraharmonic ratios (e.g. 5/2 over 1/2) and harmonic ratios (e.g. 4 over 2).

Results
The ultraharmonics ratio obtained during in vitro thermal-like ablation is plotted in Fig. 18. This ratio significantly decreases when acoustic signal crosses through the skull. Thus a threshold can be introduced to binary localize cavitation inside/outside the skull.

Table 2 summarises the results for harmonic ratio 4 over 2 and ultraharmonic ratio 5/2 over 1/2 obtained in vitro and in vivo during BBB opening and thermal necrosis. For BBB opening, neither the sub- nor the ultraharmonics appears in vivo, probably due to the confinement of the largest microbubbles by the vasculature. Thankfully in vitro experiments point out that the harmonic ratio remains relevant to binary localize microbubbles in the human case. Indeed the harmonic ratio 4 over 2 exhibits a -30 ± 3 dB decrease when crossing through human skull.

In order to ensure the repeatability of this method, each configuration was statistically investigated by plotting receiver operating characteristics (ROC) curves. These plots (Fig. 19) illustrate the performance of this approach. Except for in vivo BBB opening on monkey, ROC curves demonstrate that a sensitivity and a specificity which are simultaneously nearly 100% can be obtained for the binary localization of cavitation activity.

Conclusions
This preliminary study, mainly done in vitro, shows that a low-cost and easy-to-use PCD can binary localize cavitation activity inside/outside the skull using the filtering effect of the skull on the harmonic content of the spectra. The statistical study shows that the method is able to localize microbubble activity inside or outside the skull with high sensitivity and high specificity. We look forward to testing extensively this technique in vivo for both BBB opening and thermal necrosis.

<table>
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<tr>
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<th>Harmonic 4</th>
<th>Ultraharmonic 5/2 over 1/2</th>
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<tr>
<td></td>
<td>in vitro</td>
<td>Reference: no skull</td>
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<tr>
<td>BBB opening</td>
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<tr>
<td>(UCA + US)</td>
<td>-5</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Monkey skull</td>
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</tr>
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Fig. 17 (abstract A18). In vitro experimental setup for mimicking BBB opening

Fig. 18 (abstract A18). Ultraharmonics ratio (3/2 to 15/2 over 1/2) recorded during in vitro thermal ablation

Table 2 (abstract A18), ultraharmonic (5/2 over 1/2) and harmonic (4 over 2) ratio during thermal necrosis and BBB opening in dB scale
Inflection of temperature vs. power curve in tcMRgFUS: correlation with lesion location

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1Stanford University, Stanford, California, USA; 2University Children’s Hospital Zurich, Zurich, Switzerland

Objectives
During tcMRgFUS, power levels are progressively increased to align, verify treatment location, and create a durable ablation. It is predicted that the measured temperature at the focus for a consistent timepoint will increase monotonically with power. However, the temperature rise is often observed to actually decrease with increasing power, creating an inflection in the peak temperature vs. power graph (Fig. 20). The purpose of this work was to show this behavior in the temperature rise is correlated with a lack of alignment between the thermal lesion and the monitored scan plane.

Methods
Fifteen ET datasets were included in this study. Post-treatment 3D T2-weighted FSE images (FOV 24 cm, matrix 320x320, slice thickness 1 mm) were thresholded at the signal intensity of zone 1 (Fig. 20). Measurements from the targeted ACPC plane to the top and bottom edges of zone 2 were made in the sagittal plane.

MR thermometry (TE/TR = 12.8/100 (n = 13); TE/TR = 3.8, 13, 18, 22/100, (n = 2)) was processed with a single baseline subtraction (α = 0.00909), followed by referenceless processing for constant and linear terms. The maximum temperature of the third time point (k-space center 8.8 s after sonication initiation) in the axial scan plane was plotted vs. power. This time point was chosen because all sonications were at least 10s in duration. The number of sonications after any inflection was noted.

In 12 cases, temperature-power curves from early sonications in the axial scan plane were extrapolated to higher power levels. The amount that the measured temperature was below the estimated temperature was measured.

Results
The results are shown in Fig. 21. While the distance from the treatment plane to the inferior aspect of the lesion remains essentially constant, the distance from the treatment plane to the superior aspect of the lesion increases with number of sonications after inflection. While it is expected that multiple sonications at the same location may increase lesion size, this data demonstrates that lesion size increases preferentially in the direction of the transducer as we increase sonications after inflection. The movement of the lesion superiorly from the AC-PC plane is correlated with the difference between the actual and estimated temperature (ΔT). Although this was quantitated only in the axial scan plane, the inflection was seen in all three scan planes. Prior work demonstrated that the lesion is double oblique (the superior aspect of the lesion is posterior and medial); therefore, movement of the focal spot towards the effective transducer aperture moves it also out of the monitored sagittal and coronal scan planes.

Conclusions
One interpretation of this data is that an increase in the acoustic absorption effectively shields the focal spot, with subsequent hotspots located closer to the transducer. A second explanation is that the acoustic properties of the skull may be changing during treatment. An additional implication of this work is for comparison of simulation with thermometry. To simplify this comparison without the complication of the alignment of the thermometry plane, a sonication early in the treatment should be used for comparison.
target brain tissue. After tuning of a closed loop control system in a phantom, one acute and seven chronic pig experiments with three-day survival were conducted to investigate the correlation of lesions between the MR images of pig brain tissue with the corresponding histology. Cranectomy was performed to create an acoustic window, and sonication was applied on 4 spots in the thalamus of each pig. Absolute temperature in pig brain tissue was computed based on the MR thermometry using the rectal temperature as a baseline. TD varied from 7 to 195 CEM with target temperature between 46 and 52 °C at appropriate acoustic powers. This pig study was approved by the University of Alberta.

**Results**

From the acute pig experiment, we observed one large and 2 small lesions on MR images 1 hour after a sonication and subsequent histology showed 4 lesions of target. For the chronic pig experiments, 22 sonication spots in 6 pigs were analyzed through MR images and histology. One pig was excluded due to air bubbles introduced between the dura and scalp during the surgery procedure, and 2 sonication spots were failed to generate due to technical problems. Results show that large brain tissue damage was observed in MR images in all 7 spots with doses larger than 100 CEM and the corresponding histology results confirmed infarction with necrotic center for all except one with a dose of 101 CEM. The diameter of those lesions on T2-weighted axial MR images was measured to 2.9 ± 0.4 mm (mean ± SD) with a mean volume of 30.7 ± 12.9 mm³. All with TD lower than 17 CEM produced no visible lesions in either MR images or histology. There was a discrepancy in generating lesions with TD between 18 and 100 CEM, so that six smaller lesions (3 in volume) were shown except one large change in MR images at

**Conclusions**

In conclusion, large tissue damages were observed on MR images and histology for all TD above 104 CEM, but no change was shown for all TD below 17 CEM. There is a variability in tissue changes between these TD levels. These results may contribute toward prescription of thermal dose rather than peak temperature or acoustic power for brain treatments, and expand the treatment envelope beyond the current limitations in selecting targets and patients.

**A21 Visualization tools for transcranial Focused Ultrasound procedure planning, simulation and analysis**

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**Objectives**

The interactions between particular transcranial transducer and skull geometries are challenging to understand without interactive visual computing tools. Such a tool has been created to allow the visual exploration of the estimated treatment envelope of a transcranial transducer given a patient specific imaging dataset. The impetus for developing such a system is to aid in understanding treatment envelope constraints and to serve as a tool for specifying the input to various acoustic simulation systems.

**Methods**

A visualization application was created which makes heavy use of a modern GPU to interactively display a transcranial Focused Ultrasound transducer, patient specific brain and skull anatomy, and the interaction of each transducer element beam path with the skull. The incident angle of each transducer element beam axis with the outer table of the skull is calculated and displayed in an interactive fashion as the natural focus of the transducer is moved within the intracranial volume. Skull geometry is derived from a treatment compatible CT dataset. An MR dataset is registered with the CT for targeting. Due to the interactive nature of the tool, prescriptive targets can be rapidly and intuitively explored and understood in terms of geometric constraints and estimated treatment efficiency.

**Results**

The visualization system functions will be demonstrated including targeting, transducer positioning and assessment of transducer efficiency in terms of effective transducer element count.

**Conclusions**

An interactive visualization system has proven valuable for facilitating understanding of the complex interaction of transducer and skull geometries. Future applications of this system may include patient screening, post-treatment analysis, indication feasibility screening and acoustic simulation initialization.

**A22 HIFU for Pediatric Operations (HOPE) – a pediatric neurosurgical treatment system**

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**Objectives**

Pediatric patients have distinctive neuroanatomic features and specific disorders that make them unique candidates for transcranial MR-guided Focused Ultrasound treatment. Children have thinner skulls, and neonates in particular, possess a natural acoustic window through their fontanelle. This results in lower phase aberration and decreases the need for the larger hemispherical dome transducers used for current adult transcranial procedures. These systems also require fixation of the head in a stereotactic frame, which is dangerous for their fragile skull. In addition, neonates often require MR-compatible incubators and dedicated neuro-interventional coils for imaging. Focal brain ablation/disconnection for medical refractory epilepsy and lysis of intra-ventricular hemorrhage are some of the proposed noninvasive treatment options for this population. To provide such treatment, we have developed HOPE – HIFU for Pediatric Operations – an integrated neonatal HIFU treatment system.

**Methods**

HOPE is composed of multiple elements: a 5 degree of freedom MR-compatible robot positioning device, a 256 element phased-array transducer, an 8 channel neuro-interventional coil, and a real-time Python-based treatment planning system for treatment delivery and software. The specifications for HOPE were determined by a team of clinicians and researchers as follows: compatible with clinical neonatal incubators and MRI techniques, imaging coil integrated into the incubator, positioning system to couple and deliver the HIFU treatment and real-time control/monitoring of the treatment (Fig. 22). The hardware elements are designed and tested with a Philips Achieva 3.0TX MRI with an Imasonic 256 element transducer. For coupling, a custom designed water bag system was created to interface with the patients’ head. The software platform includes robot kinematics, robot visualization, registration, transducer control and MRI communication. A calibration procedure was developed using a series of Vitamin E markers visible in a 3D gradient echo sequence without fat suppression; the markers were attached to an extension of the robot that recreates the acoustic cone of the HIFU beam. A user-interface module was developed to calculate the center of mass of each marker and to co-register the coordinate systems of the MRI and the robot.

**Results**

The hardware system of HOPE has been designed and tested to show it can perform T1, T2 and DTI imaging that is comparable to clinical coils while delivering HIFU treatment in a phantom model. The neuro-interventional coils provide high resolution images for treatment planning (Fig. 23). The MR-conditional robotic system positions the transducer to an accuracy of 0.59 +/- 0.25 mm and delivers thermal ablation treatment to targets in a Philips HIFU quality assurance phantom. The custom water coupler bag provided a transmission path for the HIFU with minimal energy loss. The HOPE software platform controls each of the robotic positioning axes with hardware safety switches in a real-time Python interface (Fig. 24). The software registration of the patient, MRI and robot frame allows the user to select specific brain targets. During treatment, real-time thermometry is displayed.
Conclusions
HOPE is an MR-guided Focused Ultrasound system aimed at delivering HIFU therapy (both thermal ablation and cavitation-based treatment) for neonatal and pediatric patients. The system has been designed to operate within an incubator and clinical MRI system which minimizes the impact to the patient. Future work involves characterizing the treatment accuracy and performing in vivo animal studies to test the overall system feasibility and usability for treatment of epilepsy and IVH clots. Other treatments that will be investigated are thermal ablation of brain tumors, hyperthermia and targeted drug delivery.

Fig. 22 (abstract A22). HOPE Concept (1 – incubator, 2 – coil, 3 – neonatal patient, 4 – transducer, 5 – robot and 6 – MR bore)

Fig. 23 (abstract A22). Coil comparison in a T1-TFE image of a porcine brain in vivo (Left: Prototype 8 channel neuro-interventional coil, Right: Philips clinical 32 channel head coil)

Fig. 24 (abstract A22). HOPE software interface with robot, treatment planning and real-time monitoring of HIFU exposure in phantom material

A23
Simultaneous stimulation of the human primary and secondary somatosensory cortices using transcranial Focused Ultrasound
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Objectives
Low-intensity transcranial Focused Ultrasound (FUS) is making progress as a new mode of non-invasive brain stimulation, having potential for superior spatial selectivity and depth penetration compared to transcranial direct current stimulation (tDCS) or transcranial magnetic stimulation (TMS). With accumulating evidences of the FUS-mediated neuromodulatory effects in small and large animal models, the sonication given to the primary somatosensory cortex (SI) has recently shown to be capable of eliciting tactile sensations in humans. Here, we further investigated the creation of tactile sensations induced by simultaneous FUS stimulation of the secondary somatosensory cortex (SII) and SI of the hand.

Methods
Ten healthy volunteers (two females, ages = 23–34, average of 27.8 ± 4.1 yrs) participated, and all procedures were conducted under the approval of the local Institutional Review Board of the Catholic University of Korea. For targeting of FUS focus to the individual functional neuroanatomy, each participant’s brain image was acquired using a 3-T MR scanner with anatomical (T1-weighted) and functional MRI (fMRI, T2*-weighted) protocols. The SI and SII areas were mapped while using four types of external tactile stimuli to the palm of the right hand—vibrotactile, pressure, warmth, and coolness. CT scan of the head was also acquired for the planning of transcranial sonication path. As guided by these individual-specific neuroimage data, FUS sonication (210 kHz, single-element FUS transducer with focal length of 38 mm) was administered to the SI and SII simultaneously (or separately), with an incident acoustic intensity of 35 W/cm2 Isppa, tone-burst-duration of 1 ms, pulse-repetition frequency of 500 Hz (yielding a 50 % duty cycle), and a sonication duration of 500 ms. The SII was targeted as divided into four sub-regions that are specifically activated by the external tactile stimuli.

Results
Across the differential selective stimulations (i.e., SI only, SII only, or SI and SII simultaneously), participants felt various types of elicited tactile sensations (while ‘tingling’ was dominant) from the hand areas contralateral to the sonication, such as the palmar/dorsal side of the hand or as single/multiple adjacent fingers. These results were similar to our previous study on FUS stimulation of the SI, while the elicitations of ‘vibrotactile’ and ‘warmth’ sensations were newly reported in the present study. The types of tactile sensations did not match to the sensations that are associated with the specific sub-regions in the SII. However, two individuals reported matching types of sensations (‘vibrotactile’, ‘pressure’, and ‘warmth’) during stimulation of the SI/SII simultaneously or the SII only. The stimulatory effects of the FUS were transient and reversible, and the sonication procedures did not induce any discomforts or adverse changes in the subjects’ physical/mental status.

Conclusions
Simultaneous stimulation of the SI/SII in the same hemisphere was achieved by using multiple FUS transducers, which elicited various types of tactile sensations. Stimulation of the SII only also induced the creation of tactile sensations. The ability to stimulate multiple region-specific brain areas may shed light on examining the causal relationships between regional brain activities and subsequent behavioral/cognitive outcomes.
A24
Transcranial Focused Ultrasound stimulation of the primary visual cortex in humans
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Objectives
Transcranial Focused Ultrasound (FUS) has been suggested as a new non-invasive modality of regional brain stimulation, with potential to be more spatially-selective and to reach deep cortical/subcortical areas compared to the conventional methods of transcranial magnetic stimulations (TMS) or transcranial direct current stimulation (tDCS). In humans, low-intensity FUS sonication has been demonstrated to temporarily change the neural activities in the primary somatosensory cortex (SI), based on the observations of subjective sensory manifestations and electrophysiological responses. However, functional neuroimaging evidence of increased neural activity in the stimulated region, as well as the associated network-wide brain responses, has not yet been shown in humans. Here, we administered stimulatory FUS to the primary visual cortex (V1) as guided by the individual-specific neuroanatomy. Concurrent functional MRI was acquired to assess the brain regions that were activated due to the stimulation.

Methods
19 healthy volunteers (five females, ages 20–45, average 26.1 ± 5.4 yrs) participated, and all procedures were conducted under the approval of the Institutional Review Boards of both the Catholic University of Korea and Korea University. Functional MRI (fMRI); for mapping of the visual areas) and cranial CT were obtained from each participant to provide the individual-specific V1 location for sonication planning/targeting. Then, in a separate session, an MR-compatible sonication setup (270 kHz, single-element FUS transducer) was used to deliver FUS to the V1 under a clinical 3-T MR scanner for the image-guidance and the simultaneous acquisition of fMRI data. Separate from the FUS-fMRI session, electroencephalographic (EEG) potentials elicited by the FUS stimulation were also measured. We used a pulsing scheme having a sonication duration of 300 ms with a tone-burst-duration of 1 ms repeated at a pulse repetition frequency of 500 Hz (yielding a 50 % duty cycle). The incident acoustic intensity at the FUS focus was 16.6 W/cm² Isppa. Retrospective numerical simulation of the transcranial acoustic wave propagation was performed proximal to the sonicated area to estimate the in situ acoustic intensity and spatial accuracy of sonication.

Results
Simultaneous acquisition of fMRI during FUS sonication to the V1 revealed the elicited activation not only from the sonicated brain area, but also from the network of regions involved in visual and higher-order cognitive processes. Accompanying phosphene perception was also reported. The EEG responses showed distinct peaks associated with the sonication, having similarities with the classical visual evoked potentials (VEP) generated by photic stimulation. The procedures did not induce any discomforts or adverse effects from the participants, based on the subjective reporting and neuroradiological/neurological examinations. Retrospective numerical simulation of the transcranial FUS suggested the variability in individual responsiveness to the stimulation.

Conclusions
Simultaneous fMRI acquisition during FUS application to the V1 revealed the functional neuroimaging-based evidence in humans that the FUS stimulation activates the sonicated brain area and concurrently elicits the associated phosphene perception. Successful stimulation of the V1 was also supported by the presence of the evoked EEG potentials associated with FUS. The individual variability in responsiveness to the stimulation suggested needs for an elaborate image-guidance.

A25
Focused Ultrasound modulation of visual search performance and associated EEG in monkeys
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Objectives
Focused ultrasound (FUS) is a promising tool for neuromodulation because of its noninvasiveness and better spatial precision compared to other noninvasive methods, such as transcranial magnetic stimulation (TMS) or transcranial direct current stimulation (tDCS). FUS neuromodulation has been demonstrated in multiple animal models, including a prior study where ultrasound was applied transcranially over macaque frontal eye field (FEF) to influence saccade response time in an anti-saccade task. In this work, we applied FUS through a craniotomy over the macaque FEF while measuring saccade response times and EEG signals associated with selective attention.

Methods
A single element focused transducer was positioned through a craniotomy over FEF, a cortical area that plays a key role in the eye movement and attention systems. FUS stimulation was applied during a complex visual search task where the monkey was required to shift its gaze to a target among distractors. We alternated blocks of trials with or without FUS stimulation (300 ms of pulsed FUS with a 50 % duty cycle starting 150 ms before search display onset, center frequency 500 kHz, repetition frequency 2 kHz, pulse duration 0.25 ms, peak negative pressures of 250 kPa or 425 kPa, warming of brain tissue < 1.5 °C). Saccade response time and intracranial EEG recordings were acquired in two monkeys performing 9 sessions each.

Results
In both monkeys, event-related potentials (ERPs) associated with selective attention (N2pc) were significantly reduced during stimulation with both intensities. FUS stimulation attenuated the N2pc over the entire session block, rather than on a trial-by-trial basis. In one monkey with a craniotomy positioned directly over FEF, the mean saccade response times were reduced by 5 ms by FUS stimulation at 425 kPa (p < 0.001) when the target appeared in the upper hemifield contralateral to the FUS stimulation, while another animal with a craniotomy more ventrally did not show such a systematic behavioral modulation.

Conclusions
We are continuing to explore potential spatial relationships between stimulation location and behavioral modulations in ongoing work. Overall, our findings demonstrate prolonged FUS modulation of attention ERPs and suggest potential spatial selectivity based on the location of stimulation.

A26
Ultrasound-mediated modulation of motor and ocular responses in anesthetized mice in vivo
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Objectives
Focused ultrasound (FUS) has been identified as a non-invasive technique for modulating brain activity. Most studies involving sedate rodents utilize frequencies in the kilohertz-range, which allow for optimal transmission of acoustic power through the skull. The tradeoff with using lower frequencies involves producing larger acoustic foci and resultant poor target-specificity. Megahertz-range frequencies can therefore be used to improve target-specificity. This study demonstrates that Focused Ultrasound in the megahertz range can be used to evoke motor and ocular responses in mice under deep anesthesia by targeting cortical and subcortical structures, respectively.
Contralateral-paired hind limb movements were observed when stimulating cortical regions, demonstrating the ability of megahertz-range FUS to stimulate activity in highly-targeted regions. Additionally, pupil dilation was observed when deep-seated anxiety-related structures were targeted, demonstrating the ability of FUS to modulate activity in small subcortical structures.

Methods
For this study, wild-type adult male mice were anesthetized with intraperitoneal injections of sodium pentobarbital (65 mg/kg) and fixed in a stereotaxic frame. A single-element FUS transducer with fundamental frequency of 1.94 MHz was fixed to a 3D positioning system for accurate navigation through the brain. A 6x6 mm grid centered -2 mm anterior of the lambda skull suture was sonicated in a random order using a center frequency of 1.9 MHz, pulse repetition frequency of 1 kHz, 50 % duty cycle, 1 second pulse duration, 1 second inter-pulse interval for a total of 10 pulse repetitions. The acoustic pressure applied was varied in order to evaluate thresholds for eliciting physiological responses like motor movement, eye movement, or pupil dilation. Motor movements were validated using video recordings and intramuscular electromyography recordings from the biceps femoris in both hind limbs. Pupil movement and dilation from subcortical modulation were evaluated using a high-resolution camera aimed at the right eye and frame-by-frame processing technique.

Results
The minimum peak rarefractional pressure required to elicit hind limb movements was 1.45 MPa when targeting cortical regions, calibrated using an excised mouse skull. Higher pressures increased the success rate from 20 % (at the 1.45 MPa threshold) to 70 % (1.79 MPa) (Fig. 25). Targeting eye-motor and anxiety-related regions of the brain elicited eye movements and pupil dilations up to 20 %. Sonicating the superior colliculus resulted in both eye movement and pupil dilation at a lower threshold pressure (1.20 MPa) than the hippocampus and locus coeruleus, which required pressures greater than 1.80 MPa. A histological evaluation performed in five mice at 1.93 MPa and 3 MPa peak rarefractional pressure resulted in no red blood cell extravasation (Fig. 26).

Conclusions
This study successfully demonstrated that megahertz-range Focused Ultrasound can be used to elicit motor and oculor responses with high specificity in mice in vivo. It was also shown that the success rate of stimulation increased with acoustic pressure for motor movements associated with cortical modulation but depends greatly on the region of the brain targeted. These findings emphasize the complexity and yet to be determined mechanism of action involved in ultrasonic neuromodulation.

A27 Non-invasive neuromodulation via targeted delivery of neurotransmitter chemicals

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Objectives
Focused ultrasound (FUS)-microbubble treatment has been used to open the Blood-Brain Barrier (BBB) for targeted delivery of a wide variety of therapeutics. Here we propose to deliver neurotransmitter chemicals such as GABA or glutamate for the purpose of non-invasive neuromodulation. These chemicals function to transmit or suppress signals across the chemical synapses that connect neurons in the brain. This novel approach affects signaling between neurons, as opposed to existing neuromodulation techniques that affect the transmission of electrical signals along neurons. Such an approach could be an important new complimentary tool for basic neuroscience or lead to new therapeutics for neurological disorders.

Previously, we used electrophysiology measurements to demonstrate functional blockade via BBB disruption and GABA administration. Here we present initial results demonstrating the proof of concept in a rodent model using delivered GABA to modulate neuronal activity and functional MRI to measure the effects.

Methods
Sprague-Dawley rats underwent bilateral hindpaw electrical stimulation (1.5 mA, 0.3 ms duration, 2 Hz) to elicit a functional response of the somatosensory network. Varying levels of GABA were systemically injected under conditions No BBB opening and BBB opening. Functional activity in the thalamus and SI was measured using fMRI to quantify any effects of neuromodulation.

BBB opening: Microbubbles injected (Optison, 200 μl/kg), 274 kHz dual aperture transcranial FUS with 32 ms bursts applied at 4 Hz for 60 seconds.

GABA opening: Microbubbles injected (Optison, 200 μl/kg), 274 kHz dual aperture transcranial FUS with 32 ms bursts applied at 4 Hz for 60 seconds.

fMRI delivery: Systemic tail vein bolus injection in doses from 10 mg/kg to 50 mg/kg.

fMRI: Images acquired on a Bruker 7 T scanner with a single shot EPI sequence (TR = 1.5 s, TE = 18 ms, 18 slices, 300 images). Stimulation performed for 40 s OFF, 20 s ON block design over 7.5 total minutes. T-scores obtained using general linear model analysis in SPM 12.

Results
BBB Closed: Fig. 27 shows activation results in S1 for the case of No BBB opening. Compared to the baseline case of No GABA injected, a GABA injection of 10 mg/kg showed significant decrease in activity (p < 0.05) but GABA injections of 25 mg/kg and 50 mg/kg did not.

BBB Open: Fig. 28 shows activation results in the thalamus for the case of BBB opening. BBB opening was targeted, and confirmed through gadolinium imaging, in the right hemisphere. Bilateral activation was seen in the thalamus for the baseline case of No GABA.
injected. For GABA injection of 25 mg/kg, a significant decrease in activation was seen in the right (opened) ROI (p < 0.001), but not the left (unopened) ROI. For GABA injection of 50 mg/kg, a significant decrease in activation was seen in both ROIs (p < 0.001).

Conclusions
More experiments on a number of rats are necessary to confirm and expand these findings. However, these very preliminary results are a promising indicator that a neurotransmitter such as GABA can be delivered through the opened BBB for targeted manipulation of neuronal activity.

Objectives
Magnetic Resonance-guided Focused Ultrasound (MRgFUS) has been shown to reversibly open the Blood-Brain Barrier (BBB) for targeted drug delivery [1]. Research on animal models, including non-human primates [2], has been conducted to investigate the effectiveness and characteristics of BBB openings. Here we describe our initial experience in a pilot clinical study to establish the feasibility, safety and preliminary efficacy of Focused Ultrasound to temporarily open the BBB to deliver chemotherapy to brain tumors.

Methods
This phase-one clinical trial of BBB opening by Focused Ultrasound was approved by Health Canada. A modified clinical MRgFUS brain system (ExAblate 4000, 230 kHz, Insightec, Tirat Carmel, Israel) was used with a 3 T MR scanner (Signa MR750, GE Healthcare, Milwaukee, WI, USA). Two hours before the procedure, liposomal doxorubicin Caelyx (Janssen, Toronto, Ontario, Canada) was intravenously infused over 1 hour at a dose of 30 mg/m2. The patient’s head was then shaved and positioned in the FUS array with a stereotactic frame. Two targets close to the posterior margin of the glial tumor were chosen based on T2 images (Fig. 29). Each target consisted of a 3x3 grid of 9 spots at 3 mm spacing. For each spot, 2.6 ms on, 30.4 ms off FUS pulses were repeated for 300 ms before steering to the next spot. The pattern was repeated periodically resulting in an overall pulse repetition frequency (PRF) for each spot of 0.9 kHz. A bolus injection of 4 ul/kg of Definity microbubbles (Lanthus Medical Imaging, N. Billerica, MA, USA) was applied simultaneously with each sonication (1/5th of the clinical dose for ultrasound imaging). With the first injection of microbubbles, 10s short sonications at 5 W, 7 W and 9 W acoustic power were applied to find the appropriate power level based on feedback of cavitation signals. Cavitation signals were detected by two receivers and sampled at a rate of 2 MHz. Spectrum integration from 75 kHz to 155 kHz was calculated and two threshold levels of the spectrum integration were defined as a safety mechanism based on pre-clinical studies on a trans-human skull pig model [3]. 9 W was found to be adequate for these targets. 50 s sonications at 9 W were then applied at each target, with a separate bolus injection of microbubbles for each. Post sonication, Gd (Gadovist, Bayer)-enhanced 3D FSPGR images were acquired to verify the BBB openings, and T2*-weighted GRE images (TE = 15 ms) were collected to detect potential hemorrhage. After the treatment, the patient was released from the head frame and MR scans were repeated with an 8-channel head coil for better quality images. The patient underwent routine tumor resection the next day and tissue samples at the two BBB opening targets were collected for quantification of chemotherapy drug concentration.

Results
The opening of the BBB was successful at both locations with clear Gd enhancement in the 3x3 grid pattern. (Figs. 30 and 31). Despite using the same power level, the actual acoustic pressure at the 2nd target was lower than the first due to steering of the FUS beam. Low-level extravasation of red blood cells were seen as small dark signals within individual sonicated spots in the T2* image (Fig. 32). The quantification of drug concentration is pending further analysis.

Conclusions
The 3 mm spacing of the 9 spots was intentionally designed to form a grid pattern of Gd enhancement for easier confirmation in heterogeneous tumors for the initial cases. We do not expect an impact on other parameters if the spacing needs to be reduced for a more uniform drug distribution within the BBB opening volume. There was a small level of RBC extravasation but this was not a concern in the tumor environment. Our animal experiments have shown that the cavitation signal can be used during the sonications to control the power level for eliminating the RBC extravasations [4]. The current system did not use this method during sonications. The tumor in this patient was in the right temporal lobe adjacent to the skull. The two targets were ~4 cm lateral from the midline of the brain, and the 2nd target was also ~2.5 cm posterior. Thermal ablations by FUS at these off-centre locations are technically challenging due to excessive skull heating. However, successful BBB openings at these locations were demonstrated at low powers at 230 kHz. If these results can be repeated in other patients without complications, then the method may provide a new way to deliver
therapeutic agents into brain for the treatment of tumors and other brain diseases.

References

A29
Enhanced bevacizumab delivery to the CNS by Focused Ultrasound induced blood-brain barrier opening for malignant glioma treatment
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Objectives
Malignant glioma is the most severe form of primary brain tumors with an extremely high recurrence rate and the poorest prognosis. Current anti-angiogenic monoclonal antibody (mAb) treatment failed to show therapeutic efficacy due to transient “vascular normalization” stage
that restores BBB integrity in tumor regions and restricts anti-angiogenic mAb penetration, preventing angiogenic suppression of tumor cells in the CNS and diminishing the improvement in overall survival in clinical treatment observation. The purpose of this study is to demonstrate that transcranial Focused Ultrasound (FUS) enhances Blood-Brain Barrier (BBB) permeability of the antiangiogenic monoclonal antibody, bevacizumab, for glioblastoma multiforme (GBM) treatment.

Methods

Transcranial FUS in the presence of microbubbles was used to transiently open BBB, and enhance CNS penetration of bevacizumab in normal and glioma-bearing mice. Bevacizumab was quantitated by high-performance liquid chromatography (HPLC), and Western blotting confirmed bevacizumab in the CNS. Bevacizumab permeability was estimated in vivo via contrast-enhanced Magnetic Resonance Imaging (CE-MRI), and glioma progression was longitudinally followed via T2-MRI. Morphological changes and vascular inhibition were confirmed histologically with H&E and CD-31 immunohistochemistry (Fig. 33).

Results

HPLC confirmed that FUS significantly enhanced CNS delivery of bevacizumab from 5.7- to 56.7-fold. The high correlations between CE-MRI imaging indices and bevacizumab concentration (r² = 0.56-0.7378) suggested feasible non-invasive in vivo imaging of large-molecule BBB penetration. FUS-enhanced bevacizumab delivery significantly inhibited glioma progression, and improved median survival (ISTmedian = 135 %, compared to 48 % in bevacizumab-administration alone, Fig. 34).

Conclusions

In conclusion, anti-angiogenic glioma therapy is enhanced via our proof-of-concept study that FUS enhances large molecule bevacizumab BBB permeability, and combining Focused Ultrasound to open the Blood-Brain Barrier with bevacizumab delivery can overcome bevacizumab vascular normalization and potentiate bevacizumab’s anti-angiogenic tumor therapy effect.

A30

Closed-loop control of targeted drug delivery across the blood-brain barrier in rat glioma models

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Objectives

Microbubble-mediated Focused Ultrasound (FUS) can induce targeted drug delivery through Blood-Brain Barrier disruption (BBBD). Real-time feedback control of cavitation is critical to realize desired treatment outcome while avoiding tissue damage. Here, we propose an acoustic emissions-based controlling paradigm that can sustain stable cavitation (harmonic emission, HE) while suppressing inertial cavitation (broadband emission, BE). Our objective is to deliver desired drug dose by controlling HE strength during BBBD, while keeping the brain damage-free.

Methods

A dual-aperture FUS setup (f = 274.3 kHz) produced a subcentimeter focal depth in rats’ brain (n = 50) in vivo, and a passive detector (fcentral = 650 kHz) monitored cavitation activity. HE and BE were analyzed during 32-ms bursts in real time and fed back for control of the next pulse. The impact of multiple FUS parameters and microbubble (Optison) injection protocol on the controller performance was studied. To avoid inertial cavitation, the pressure was reduced if BE was detected and terminated if it crossed a set threshold. Both wild type and F98 glioma (ATCC # CRL-2397) models have been used in this study. Delivery of a model drug (Trypan Blue; 960 Da) and chemotherapeutic drug (Doxorubicin) was assessed using fluorescent imaging of formalin-fixed tissue blocks 1-h post sonication.

Results

Pilot study demonstrated the HE-pressure linearity (R² = 0.93) and found that the BE threshold decreased as bubble dose (up to 400 μl/kg) was augmented. To optimize controller performance in sustaining HE while suppressing BE, a Phase-1 study demonstrated that: 1) 4-Hz PRF (compared to 1-Hz, P < 0.001) significantly suppressed the HE signal variance; 2) Infusing microbubbles after an initial bolus prevented the decline of HE and a corresponding increase in pressure, and further improved HE stability (P < 0.05 for all comparisons, Fig. 35a) while reducing the likelihood of BE (33.7 % vs. 16.7 %). Using optimal settings, a Phase-2 study investigated HE control and Trypan Blue delivery (Fig. 35b). Integrated HE was exponentially correlated with epi-fluorescence intensity (R² = 0.82; red symbols in Fig. 35c). Based on this calibration, a Phase-3 study tested if we could deliver a desired amount of drug by sonicating until the HE reached a preset goal. The resulting fluorescent intensity matched well with the reference curve for three different goals (n > 5 per group, green symbols in Fig. 35c).

Conclusions

Our proposed controlling system and method has been demonstrated to effectively sustain the stable cavitation behavior while suppressing the inertial cavitation at a minimum level. Moreover, this real-time closed-loop controller can enable the reliable delivery of a pre-determined amount of drug to the brain.

Fig. 33 (abstract A29). Representative imaging indexes obtained from DCE-MRI analysis (T1-WI, R1-AUC, Ktrans, and Ve). a CD-31 IHC fluorescent microcopies. Bar = 100 μm

Fig. 34 (abstract A29). Kaplan–Meier plot to demonstrate animal survival among each experimental groups. BEV = Bevacizumab

Fig. 35a (abstract A30). Representative imaging indexes obtained from DCE-MRI analysis (T1-WI, R1-AUC, Ktrans, and Ve). a CD-31 IHC fluorescent microcopies. Bar = 100 μm

Fig. 35b (abstract A30). Kaplan–Meier plot to demonstrate animal survival among each experimental groups. BEV = Bevacizumab

Fig. 35c (abstract A30). Representative imaging indexes obtained from DCE-MRI analysis (T1-WI, R1-AUC, Ktrans, and Ve). a CD-31 IHC fluorescent microcopies. Bar = 100 μm

Fig. 35d (abstract A30). Kaplan–Meier plot to demonstrate animal survival among each experimental groups. BEV = Bevacizumab
Methods
The ultrasound treatment of the antibodies trastuzumab and pertuzumab in combination with BBB disruption was evaluated using MR imaging. BBB disruption was achieved using MR-guided focused ultrasound (MRgFUS) in combination with microbubbles. The treatment benefit was assessed by temporary disruption of the BBB using magnetic resonance (MR)-guided focused ultrasound (MRgFUS) in combination with microbubbles. The treatment benefit was evaluated using MR imaging.

Results
BBB disruption was successful in all sessions with an average ΔSI% of 21.2 % (range 4.5–77.6 %). The mean ΔSI% of two tumors before BBB disruption during the six treatment weeks were 0.4 % and 0.6 %, indicating that the tumors were not leaky before disruption.

In the FUS + antibody-group, 4/10 animals were classified as responders during the treatment period with an average growth rate of 0.010 ± 0.007, compared to 0.043 ± 0.013 for the non-responders. There was no difference in the average ΔSI% of the responding rats (21.8 % ± 16.7) and the non-responding rats (20.7 % ± 9.7). None of the control or antibody-only animals were classified as responder.

High-resolution T2w imaging showed that the tumor was homogenous in most animals till week 13–15, when cystic and necrotic areas started to develop. The tumors showed also a heterogeneous appearance on H&E-stained sections and the complete tumor was HER2-expressing in the examined brains (Fig. 37).

Conclusions
In this study, we demonstrate that BBB disruption using MRgFUS in combination with antibody therapy can slow down the growth of breast cancer brain metastasis. As the tumors were not leaky before BBB disruption and there were no responders in the antibody-only group, the disruption of the BBB is necessary for drug delivery to these brain metastasis. Part of the rats responded to the treatment, while the other animals had the same growth rate as the control-group. This is in line with a previous study [2], where antibody therapy was combined with FUS in a different breast cancer brain metastasis model and only in part of the animals a response was observed. The difference in response could not be explained by the tumor volume at the start of the treatment, nor was there a difference in contrast-enhancement after BBB disruption or in HER2-expression. Better understanding of why certain animals respond is needed and will help in translating this technique to the clinic.

References

Fig. 35 (abstract A30). a Signal stability assessment (*: P < 0.05, ***: P < 0.001, ****: P < 0.0001); b Cavitation control profile (HE in black and BE in red) and Trypan blue delivery; c Calibrated BBBD correlation (in red) and controlled delivery results (in green)

Fig. 36 (abstract A31). a T1-weighted images before contrast administration. The red arrow indicates the tumor. b No difference in enhancement of the tumor is observed after contrast administration (ΔSI = 0.4 %). c After focused ultrasound-mediated Blood-Brain Barrier disruption, the tumor enhances after contrast administration (ΔSI = 30.1 %)
A32
Phase I/IIa clinical trial of blood-brain barrier disruption by pulsed ultrasound
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Objectives
The Blood-Brain Barrier (BBB) limits the delivery and efficacy of systematically administered drugs to the brain. Pre-clinical studies have demonstrated that ultrasound-induced disruption of the BBB can significantly increase the intracerebral concentration of chemotherapy. To harness this approach, our group has developed an implantable ultrasound device, SonoCloud®, that allows for simple and repeated disruption of the BBB. The goal of this work was to determine the safety of repeated disruption of the BBB using the SonoCloud® device in patients with recurrent glioblastoma prior to receiving carboplatin chemotherapy.

Methods
A Phase I/IIa clinical trial was initiated at the Hospital Pitie Salpetriere in Paris, France in July 2014. Patients with recurrent glioblastoma were implanted with an 11.5-mm diameter biocompatible 1 MHz ultrasound transducer. The device was fixed to the skull bone in a standard burr hole, after which the skin was closed. Once a month, the device was connected to an external generator system using a transdermal needle connection and patients received a 150 second ultrasound sonication (25,000 cycles/burst, 1 Hz) to target regions at depths of 5 to 20 mm from the skull bone. During dosage studies, single 1.5 MHz focused transducer treatments were delivered to brains of 11 pigs using a 1.5 MHz focused transducer followed a craniotomy. Lesions generated were targeted to be contained within individual gyri to avoid perforations into the sulci. During dosage studies, single lesions were generated using between 1 and 200 histotripsy pulses delivered at ≤10Hz PRF with an estimated peak negative pressure of 45 MPa. Large lesions were generated by incrementally repositioning the histotripsy target site during treatment to ablate entire target volumes. 7 acute pigs were sacrificed within 6 hours of treatment and brains were removed for MRI and histology. 4 sub-acute pigs were sacrificed within 24 hours of treatment. 3 acute pigs, 5

and well tolerated in patients with recurrent GBM and may improve chemotherapy delivery in the brain.

A33
Preliminary report on sonodynamic therapy for C6 glioma rat model
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Objectives
Not released for publication

Methods
Not released for publication

Results
Not released for publication

Conclusions
Not released for publication

A34
In vivo porcine histotripsy brain treatments
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Objectives
Focused ultrasound thermal therapies have been explored for brain applications including treatment of essential tremors, Parkinson’s disease, and stroke. Histotripsy, a cavitation based ultrasound therapy, is being investigated for treatment of brain tumor and intracerebral hemorrhage, but there are concerns that excessive hemorrhage may be induced in the brain during histotripsy treatment by disrupting blood vessels. This study investigates the in vivo feasibility and safety of generating targeted lesions in the porcine brain using histotripsy.

Methods
Histotripsy treatments were delivered to brains of 11 pigs using a 1.5 MHz focused transducer following a craniotomy. Lesions generated were targeted to be contained within individual gyri to avoid perforations into the sulci. During dosage studies, single lesions were generated using between 1 and 200 histotripsy pulses delivered at ≤10Hz PRF with an estimated peak negative pressure of 45 MPa. Large lesions were generated by incrementally repositioning the histotripsy target site during treatment to ablate entire target volumes. 7 acute pigs were sacrificed within 6 hours of treatment and brains were removed for MRI and histology. 4 sub-acute pigs were sacrificed within 3 days after treatment. To assess lesion damage, hemorrhage and edema surrounding the lesions, MR images of the brains were acquired at 1 hour and at 3 days after treatment for sub-acute pigs. Ablation sites were dissected for histology. In 4 acute pigs, histotripsy was used to generate precise single lesions of 1.5x2.5 mm within the gyri. Dosage studies revealed that a single histotripsy pulse was sufficient to generate identifiable damage in the brain in MRI and complete cell fractionation in lesions was achieved by 50 pulses. In 3 acute pigs, 5
larger lesions measuring up to 7 mm diameter were generated in the brain. MRI and histology revealed that lesions confined to within the gyri show no hemorrhage or other major complication associated with treatment. In 4 sub-acute pigs, 6 single lesions were generated by 10 and 50 pulses, and two larger square lesions of 3.5 mm were created. MRI following treatment showed no hemorrhage or damage outside the target regions and at 3 days showed no additional hemorrhage and only minor edema near the lesion boundary. No midline shift or brain herniation was observed. Histology showed damage confined to within target volumes, with well demarcated boundaries between treated and untreated tissues and no evidence of hemorrhage, ischemic changes, encephalitis, or acute, sub-acute or chronic inflammatory infiltrate beyond the areas of the lesion.

Conclusions
The results of this study demonstrate that histotripsy may be used to generate targeted lesions in the brain without causing excess hemorrhage, edema, or other major complications associated with treatment. Lesions were observed to have well defined boundaries between treated and untreated tissues, with little damage beyond the confines of the lesion area. Tissues surrounding the lesions appear very viable with no acute, sub-acute or chronic inflammatory infiltrate, and tissues in adjacent gyri appear uninvolved and unremarkable. These results demonstrate that histotripsy may be applied in the brain without causing major complications and suggest the potential of histotripsy for use in brain therapy applications.

A35
Neuronavigation-guided Focused Ultrasound and real-time acoustic mapping: evaluation in non-human primates with blood-brain barrier opening
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Objectives
Focused ultrasound (FUS) has shown great promise for noninvasive brain treatment, including surgical ablation, Blood-Brain Barrier (BBB) opening and drug delivery, and neuromodulation. Accurate and precise targeting with personalized planning is the key for treatment success. HIFU, repetitive procedure is often required for BBB opening and drug delivery as well as neuromodulation, which entails the demand for a portable system apart from the current MR-guided FUS system requiring a dedicated and custom-built suite. In this study, a neuronavigation-guided FUS system was developed (Fig. 38), combined with in silico planning based on CT and MRI as well as real-time acoustic mapping in visualizing the location and intensity of the cavitation events associated with BBB opening. The system was evaluated in non-human primates with BBB opening, with the treatment preplanning and targeting accuracy validated both in silico and in vivo.

Methods
Three rhesus macaques were sonicated (N = 15) and the subcortical structure of the basal ganglia was targeted, which is associated with neurodegenerative diseases including Parkinson’s and Huntington’s disease. Both CT and MRI were acquired for personalized pre-planning and neuronavigation guidance (Brainsight). The 3D numerical simulation of the acoustic pressure field was performed for pre-planning and post comparison with the BBB opening. A single-element, 0.5-MHz FUS transducer (diameter: 64 mm) with in-house microbubbles were used for sonication, and a programmable data acquisition system (Verasonics) with an array of acoustic detectors for real-time passive cavitation mapping. Both the FUS and cavitation mapping were guided with the neuronavigation system in real time during the FUS procedure. After sonication, the contrast enhanced T1-weighted MRI was used to confirm the location and size of BBB opening. The accuracy of both the FUS targeting and cavitation mapping were assessed.

Results
The target shift was on average 2.0 mm laterally and 3.5 mm axially in the in vivo experiment, and the shift due to the skull was predicted to be 0-1 mm laterally and 1.0-5.5 mm axially in silico. Real-time cavitation mapping confirmed the sonicated area with and without BBB opening, and distance between the centroid of the cavitation map and that of the resulting BBB opening was under 2 mm. In order to achieve the desired BBB opening volume, the pressure and focal spot size in situ were estimated in silico and tailored for targeting in each subject. Simulation results showed a smaller focal spot size through the skull (2.6 mm laterally and 16.7 mm axially, compared with the original size of 4.0 mm laterally and 35.3 mm axially), which corresponded to the BBB opening volume under specific peak negative pressures (NHP 1 at 200 kPa, NHP 2 at 600 kPa). This pressure difference between individuals was due to the difference in skull attenuation, since the in silico pressure reduction was estimated to be 30.9% ± 12.4% and 53.9% ± 14.3% in NHP 1 and NHP 2, respectively.

Conclusions
In conclusion, the new portable neuronavigation-guided FUS with in silico preplanning and real-time cavitation mapping was shown feasible and could facilitate the BBB opening and neuromodulation applications while maintaining translational capability to a clinical setting.

A36
Definition of basic properties of physical immunotherapy in pancreatic cancer using HIFU and immune checkpoint inhibition
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Objectives
The latest advances in cancer immunotherapy have improved our understanding of how to stimulate the patient’s immune system to fight their own disease. Notwithstanding the unparalleled success of treating melanoma tumours and non-small cell lung cancer with immune checkpoint inhibitors, patients are not all getting the same benefit. In this context, pancreatic cancer presents unique challenges. Its dense stroma characteristics result in limited immune cell infiltration in the tumour microenvironment. In addition the low number of mutations in pancreatic cancer cells result in few tumour antigens, thereby reducing their immunogenicity. HIFU may address some of these limitations by ablating segments of the tumour to induce a stronger inflammatory response and degrade the stroma to facilitate enhanced diffusion of tumour antigens out of the tumour, and increased penetration of lymphocytes to the tumour. We propose to co-treat pancreatic tumours with HIFU and immune checkpoint inhibitors to elicit an effective anti-tumour immune
Methods
In vitro characterization of the effects of heat on the presentation of cell surface receptors associated with the immune response has been undertaken in human colon cancer HCT116 cells and mouse pancreatic cancer Panc02 cells using FACS analysis. A KPC-derived pancreatic cancer cell line will be transplanted in C57BL/6 subjects to form syngeneic subcutaneous and orthotopic pancreatic tumours. These tumours will be treated with HIFU, and within vivo monoclonal anti-CLTA-4 and anti-PD-1 antibodies. The phenotype of markers of the adaptive immune response will be assessed to identify whether HIFU exposures can initiate intrinsic anti-tumoural cellular immunity at the site of treatment. Analysis by flow cytometry and immunohistochemistry of both effector cells (cytotoxic CD8+ T cells, dendritic cells), and immunosuppressive cells (Tregs, myeloid derived suppressor cells), as well as cytokines and chemokines at various time points post HIFU will be performed. Tumour growth will be determined using high resolution ultrasound imaging, and the results will be correlated with overall survival.

Results
Thermal exposure of cancer cells results in a time-dependent regulation of the transmembrane receptor CD47. CD47 decreases immediately after, and 1 day after treatment, and increases 3 and 4 days after treatment relative to the sham-exposed cancer cells. Further data on the effects of thermal exposures on the regulation of immune-associated pancreatic cancer cell surface receptors will be presented.

Conclusions
Detailed pre-clinical studies in various mouse models of pancreatic cancer can provide a mechanistic understanding of the combinatorial effects of HIFU and immune checkpoint inhibitors. Strong evidence that HIFU stimulates endogenous immunity against pancreatic tumor cells, and augments the immunotherapy treatment of poorly immunogenic pancreatic tumours could be used as proof-of-concept for the initiation of Phase 1 clinical trials of HIFU-enhanced immunotherapy in patients with pancreatic cancer.

A37
Melanoma growth control via ultrasound depends on the adaptive immune system and surpasses αPD-1
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Objectives
Melanoma incidence continues to rise, while Stage IV 5 year survival remains below 20 %. Targeted immunotherapy approaches, designed to enhance natural anti-tumor mechanisms or inhibit the immunosuppressive tumor microenvironment, have shown promise in early clinical use. The programmed death receptor and its ligand (PD-1 and PD-L1) have drawn considerable attention. PD-L1 is known to inhibit the activity of anti-tumorigenic cytotoxic T cells, promoting a pro-tumor immune profile, and ultrasound (US) can promote the infiltration of immune cells into the tumor. Here, we tested the hypothesis that a combination of pulsed US, microbubbles (MBs) and anti-PD-1 will improve tumor growth control in a mouse model of melanoma in a manner dependent on the adaptive immune system. Melanoma incidence continues to rise, while Stage IV 5 year survival remains below 20 %. Targeted immunotherapy approaches, designed to enhance natural anti-tumor mechanisms or inhibit the immunosuppressive tumor microenvironment, have shown promise in early clinical use.

Methods
Eight week old immunocompetent C57BL/6 male mice were inoculated unilaterally with B16-F10 melanoma cells and divided into four groups: Control (untreated), US + MBs, αPD-1, or US + MBs + αPD-1. Immunocompromised Rag1-/- mice were similarly inoculated and divided into two groups: Control (untreated) or US + MBs. On day 11 post-inoculation, animals receiving US were anesthetized and sonicated with a 0.75" unfocused 1 MHz transducer as previously described1 for 60 minutes. MBs (105/g b.w.) were injected i.v. throughout the sonication. The αPD-1 groups were given an IP injection of 250 μg of αPD-1 on Days 10, 13 and 16. Tumors were monitored until Day 18, when tumors were resected for flow cytometry. Helper T cells (CD4+), cytotoxic T cells (CD8+), T regulatory cells (Treg), macrophages and natural killer cells (NK) were counted in immunocompetent mice, and dendritic cells (DC), natural killer cells, M1 and non-M1 macrophages were counted in immunocompromised mice.

Results
In immunocompetent mice, treatment with US + MBs provided improved tumor growth control compared to all other groups (Fig. 39a) as well as a significant survival benefit (not shown). Additionally, US + MBs generated a significant increase in macrophages (not shown) and Tregs (Fig. 39b) compared to control. Interestingly, the αPD-1 and US + MBs groups performed similarly in CD4+ and CD8+ cell counts (Fig. 39b), but the combined US + MBs + αPD-1 treatment generated fewer T cells than either treatment alone. In immunocompromised animals, treatment with US + MBs did not provide tumor growth control (Fig. 39c) or a survival benefit (not shown), but did generate a significant decrease in M1 macrophages within the tumor (Fig. 39d), although total myeloid cell counts remained the same (not shown).

Conclusions
US indiscriminately increases the numbers of immune cells within subcutaneous melanoma; therefore, we expected that US-enhanced delivery of αPD-1 would provide an additional benefit; namely, a shift towards an anti-tumorigenic immune profile. However, αPD-1 did not synergize with US + MB treatment. While the mechanism is not yet understood, low power noninvasive US + MBs provided excellent tumor growth control and may serve as a stand-alone clinical treatment. Furthermore, this anti-tumor effect is clearly dependent on the adaptive immune system, since US + MBs treatment in immunocompromised mice produced no benefit. A better understanding of this mechanism may identify adjunct treatments that would synergize with US and further enhance the anti-tumor effect demonstrated here.

Reference

Fig. 39 (abstract A37). a Tumor growth in immunocompetent C57BL/6 hosts. *P < 0.05 vs. control. b Immune cell representation in tumors in C57BL/6 hosts. c Tumor growth in immunocompromised Rag1-/- hosts. d Immune cell representation in tumors in Rag1-/- hosts.
A38
Stress response of cancer cells after low intensity Focused Ultrasound
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Objectives
Low intensity Focused Ultrasound (LOFU) results in sublethal stress to tumor cells. We investigated the cell surface expression of immunomodulatory molecules in three murine cell lines (Lewis lung, breast and prostate adenocarcinoma). The immunomodulation was examined by flow cytometry of cell surface stress proteins: HSP70 & calreticulin, of immunomodulatory co-activating signals: CD40, CD80, CD86 and other receptors: MHC I, Fas). Surface expression of these molecules trigger activation of or phagocytosis by dendritic cells. Additionally, the activation of the unfolded protein response (indicated by phosphorylation of PERK or IRE1α or cleavage of ATF6α) can trigger apoptosis – a programmed cellular death pathway which can also activate the immune system. We hypothesize that by altering the total acoustic power of LOFU, we can modulate the stress signals both on the cell surface and within the cells.

Methods
LOFU treatment was performed on the Philips Therapy and Imaging Probe System (TIPS, Philips Research Briarcliff, USA) using 3 W or 5 W, 100 % duty cycle, 1.5 seconds, 1 mm spacing. LOFU treatment was performed on a cell pellet which were then replated and incubated at 37 °C, 5 % CO2 for the specified time. At the end of the incubation time, cells were lightly scraped for flow cytometry staining or lysis buffer was added to the adherent cells and then scraped for Western blot analysis.

Results
The analysis of murine lung and breast cancer indicates that there is a significant surface localization of both HSP70 and calreticulin as early as 4 hours after LOFU treatment and this localization diminishes by 24 hours after treatment. For immunomodulatory co-activating signals: CD40, CD80, CD86 and other receptors: MHC I, Fas), Surface expression of these molecules trigger activation of or phagocytosis by dendritic cells.

Conclusions
Tumor stress response to LOFU treatment can be altered by the total acoustic power of LOFU treatment. The increased surface localization of HSP70, calreticulin, CD40, CD80 and CD86 have the capacity to augment the immune response to the tumor. Therefore, LOFU can be used in combination with ablative therapies in order to induce anti-tumor immunity and destroy the primary tumor.

A39
The “abscopal” effect after USgHIFU treatment of advanced pancreatic cancer
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Objectives
Pancreatic cancer is considered one of the main big “killers” in Oncology, with still a very poor prognosis, both in patients amenable to resection and of course in more advanced stage disease. More than 50 % of patients are diagnosed with an advanced stage disease, where chemotherapy is usually the main therapy option. External radiotherapy is the most common palliative loco-regional treatment in pancreatic cancer, but in the last few years HIFU has been proposed as an alternative option for palliation in advanced stage.

From our specific experience of USgHIFU for pancreatic cancer, we report few cases of ABSCOPAL effect: where after a local treatment not only the target tumor but also the distant deposits will shrink.

Methods
From 2008 until April 2016, 72 consecutive patients affected by pancreatic tumors were selected by a dedicated Tumor Board for receiving treatment with USgHIFU, for local tumor control and palliation. 64 % of patients were affected by pancreatic adenocarcinoma and 84 % of all the treated tumors were located at the level of pancreatic head. Majority of the patients had at least two lines of chemotherapy and 45 % of them received concurrent and/or consequent external RT. USgHIFU treatment was performed always during general anesthesia with GI catheter and abdominal compression with a dedicated water balloon in order to reduce the distance between the HIFU probe and the target and for push the bowel loop away from the acoustic treatment window. Patients were than evaluated by clinical and instrumental follow up.

Results
All the procedures were technically successful, with only one main complication due to a complete portal thrombosis developed 24 hrs after treatment. Objective response (OR) at MDCT, MRI and PET was 64 % with a disease control (DC) up to 90 % of all the patients. Pain was palliated in 23/27 of symptomatic patients. In four patients (4/46 with adenocarcinoma) the imaging follow up after the treatment revealed the shrinkage of metastases located in other sites: 2 retroperitoneal lymphnode (Figs. 40 and 41), 1 liver and 1 lung.

Conclusions
In advanced pancreatic carcinoma, HIFU can have a role in locally advanced unresectable disease. Based on our limited experience, HIFU could be also suggested in a metastatic setting (Stage IV), together with systemic therapy, with the potential benefit of being effective also outside the treated target (abscopal effect). There are few papers reporting the effect of HIFU as a booster of immune response in cancer patients, but there are still no data supporting the use of this non-invasive approach in advanced cancer patient just for this purpose. Specific dedicated clinical trial should be conducted for better understand the biological mechanism behind it.
Characterizing the immune response to boiling histotripsy ablation of renal carcinoma in the Eker rat

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Objectives
Boiling histotripsy (BH) is an experimental non-invasive Focused Ultrasound (US) technology. BH uses milliseconds-long US pulses at low duty cycle to mechanically homogenize targeted tissue. We aim to evaluate the adaptive immune response to BH ablation of spontaneous renal tumors in the Eker rat hereditary renal carcinoma (RCC) model.

Methods
Genotyped Eker rats (Tsc2 heterozygotes) were monitored for de novo RCCs with serial US until tumors were ≥8 mm. Syngeneic Wild-type (WT) (planned total n = 36) and Eker rats (planned total n = 48) were then randomly assigned to BH (n = 10 total to date) or an US SHAM procedure (n = 22 to date). BH was performed extracorporeally using US-guided small animal FUS system (VIFU-2000, Alpinion). The 1.5 MHz transducer was operated at duty cycle of 1 %, 10 ms pulses, 525-550 W electric power. Treatments targeted a ~0.5 cc area of the lower pole in WT rats and the largest RCC with a margin of normal kidney in the Eker rats. Blood samples were collected immediately before treatment and serially post-treatment. Rats were euthanized at 48 hrs, 7 days, 14 days, or 56 days following treatment. At euthanasia, ipsilateral and contralateral kidneys, hilar lymph nodes, and the spleen were collected. Intrarenal and circulating plasma cytokines were assessed using a 10 cytokine multiplex assay. Renal/tumor infiltrating leukocyte populations were assessed using immunohistochemistry. Leukocyte populations in tumor draining hilar lymph nodes and spleens were assessed with flow cytometry.

Results
BH treatment has been successful in all treated (n = 4 Eker, n = 6 WT to date) subjects, producing hypoechoic regions on US consistent with BH treatment effect. BH treatment was associated with significantly increased mean relative-plasma TNF-α vs. sham treatment at 0.25 (p = 0.02), 1 (p<0.04), and 24 (p = 0.05) hours. At 48 hours, significant differences in intra-renal cytokines were observed in BH (n = 4) vs SHAM (n = 4) treated rats: IFN-gamma, IL-10, and IL-8 with trend towards differences in TNF-alpha and IL-6 (Fig. 42). On immunohistochemistry, BH treated Eker rats had increased CD8+ T-cells in both the treated and contralateral kidney compared to SHAM treated rats at 48 hours (Fig. 43). Additionally, increased infiltration of f4/80+ M1 macrophages into the treated kidney, was observed in 2/4 BH treated rats vs. SHAM at 48 hours. Longer-term data is pending at this time.

Conclusions
These data represent preliminary findings indicating differences in cytokines and tumor infiltrating leukocytes between BH treated and SHAM treated Eker rats up to 48 hours post-treatment. Assessment of longer-term immune effects of BH treatment and their significance is on-going and will be reported.

Use of shock-wave exposures for accelerating thermal ablation of localized tissue volumes
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Objectives
In High Intensity Focused Ultrasound (HIFU) applications, nonlinear acoustic effects can result in the formation of high-amplitude shock fronts in focal waveforms, with amplitudes exceeding 100 MPa. The presence of such shocks leads to increased tissue heating which can be beneficial for HIFU thermal therapy. The goal of this work was to evaluate the efficacy of different shock-wave exposures to accelerate thermal ablation of tissue volumes while enabling safer conditions for intervening tissues.

Methods
Simulation studies were performed for a multi-element 1.2 MHz HIFU phased array of a clinical system (Fig. 44a, Sonalleve V1, Philips, Vantaa, Finland) for three different peak intensity levels at the array elements of 1.2, 8, and 15 W/cm2. A pulsing scheme was combined with discrete electronic steering of the array focus over a series of foci separated about 1 mm and arranged in 4 concentric circles (Fig. 44b). The circles were positioned in a plane at 25 mm depth in liver tissue of 50 mm thickness (Fig. 44a). The period between consecutive pulses was 20 ms. Pulse duration was varied to keep a constant time-average intensity at the array elements: 20 ms at 1.2 W/cm2, 3 ms at 8 W/cm2, and 1.6 ms at 15 W/cm2. The Westervelt equation combined with the bioheat equation was used for...
modeling temperature and thermal dose in tissue. Initial temperature in tissue was 20 °C, sonication was performed starting from the center and spiraling outward until a thermal dose of 1.76 s at 56 °C (equivalent to 240 CEM at 43 °C) was reached at each circle of the trajectory.

**Results**

The focal waveforms simulated in tissue for different peak intensities representing quasilinear sonication (1.2 W/cm²), sonication with a fully developed shock of 95 MPa at the focus (8 W/cm²), and sonication with higher focal shock amplitude of 118 MPa (15 W/cm²). Tissue ablation was achieved much faster at higher peak intensities with more clearly defined boundary between treated and untreated tissue. While temperature distributions and thermal dose contours in the focal plane were similar in shape for each trajectory and intensity level, corresponding distributions in the axial plane varied greatly. Slower heating at 1.2 W/cm² resulted in strong heat diffusion in the axial direction and much larger ablation volumes. On the contrary, with rapid shock-wave heating, the ablated tissue volume followed the geometry of the targeted heat deposition. Although overall ablated volumes were smaller at higher peak intensities, the ablation rate was the highest at 15 W/cm².

**Conclusions**

In comparison with conventional HIFU treatments, shock-wave exposures provide higher ablation rates at target sites with less heat diffusion to the surrounding tissues. Shock-wave heating regimes may therefore be clinically advantageous for accelerating thermal HIFU treatments, reducing the heating of surrounding tissues, and providing sharper margins of lesion volumes.

**Fig. 44 (abstract A41).** a Geometry of the numerical experiment: US beam is focused at the center of a 5 cm thick tissue layer. b Trajectory of the focus in the focal plane comprising 4 circles of 1, 2, 3, and 4 mm radii with 1 mm distance between the foci in each circle.

**A42**

**Effects of dosing and focal spacing in rapid ablation of large tissue volume using histotripsy with electronic focal steering**

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**Objectives**

Current tumor ablation techniques, typically thermal-based, are either limited to treating tumors <3 cm in diameter (RFA and microwave) or are very slow (HIFU). Because histotripsy uses microsecond-length pulses separated by up to seconds of off-time for a given focus, it is possible to electronically steer the therapy focus of a phased array transducer to excite cavitation events throughout a large volume consisting of many foci during the off-time. We hypothesize that histotripsy combined with electronic focal steering can achieve rapid ablation of a large volume. This hypothesis was investigated in this paper, and the effects of number of pulses per focal location and focal spacing on tissue damage generated using histotripsy with electrical focal steering were studied.

**Methods**

Histotripsy was applied using a 250 kHz, 256-element hemispherical phased array transducer with a 15 cm focal distance, generating 1.5-cycle, 6-microsecond pulses. Each element produced ~0.5 MPa P- individually, but focal pressure produced by all elements could not be measured due to cavitation. To establish treatment parameters including pulse repetition frequency (PRF) and number of pulses (dose) to deliver, a single-focus lesion was generated in tissue-mimicking phantoms containing a thin layer of red blood cells (RBCs) and monitored by optical imaging. Based on these results, 35 ex vivo bovine hepatic tissue samples were treated by electronically scanning the therapy focus at 200 Hz over 1000 sites (or 2 Hz per focal site) (Fig. 45). To evaluate the dose required for complete tissue fractionation, dose was varied between 20-120 pulses per focus. Focal spacing was varied from 2.5, 3.15, and 3.5 mm in the lateral direction and held constant at 4.1 mm in the axial direction to assess the extent of overlap between adjacent foci required to achieve complete homogenization. Lesion size was assessed by gross morphology and Magnetic Resonance Imaging (MRI). The degree of tissue fractionation was examined by histology (Fig. 46). Surrounding tissue temperature was measured by thermocouples 1 cm from the lesion.

**Results**

RBC phantom results suggested that fractionation efficiency was optimal near 0.2 Hz and that 120 pulses were sufficient to achieve complete homogenization. Using lateral spacing ≥ 3.15 mm or < 100 pulses, unfractionated tissue structures were observed between foci. Using 2.5 mm lateral focal spacing and 120 pulses per focal location, complete, homogeneous tissue ablation of 43 +/- 6 mL region was achieved within 10 minutes, resulting in an ablation rate of 4.3 mL/min. A temperature increase of ~4 °C was measured at the surrounding tissue. This work demonstrates that histotripsy combined with electronic focal steering can achieve rapid ablation of large volume at a rate more than double the rate of current available ablation techniques such as RFA.

**Conclusions**

Treatment of large and multiple tumor nodules remains a challenge for current tumor interventions, which are mostly thermal-based. This work demonstrates that histotripsy combined with electronic focal steering achieved homogenous and complete ablation of a large target volume at a rate greater than two-fold faster than microwave and RF ablation. Since histotripsy is non-thermal, the treatment should not be affected by the heat sink effect and is expected to remain effective and efficient even in highly vascular organs. With the capability of achieving rapid, homogenous cell disruption, histotripsy has the potential to substantially improve upon current tumor ablation methods.

**Fig. 45 (abstract A42).** MR of ex vivo sample following treatment. Scan plane orthogonal to transducer acoustic axis and positioned at geometric focus.
The TRANS-FUSIMO system — towards a prototype for HIFU treatment of the liver under breathing motion

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Objectives

The movement of the liver under breathing and its partial occlusion by the rib cage challenge the application of High Intensity Focused Ultrasound (HIFU/MRgFUS) for the thermal treatment of tumors. To explore the full potential of extracorporeal FUS to safely and precisely destroy tissue in the depth of the moving liver requires sophisticated technology. The European FUSIMO consortium (www.fusimo.eu) has shown in a proof of concept experiment that patient specific image processing, mathematical modeling of the organ’s motion, ultrasound propagation in the moving organ, and tissue heating is such enabling technology that empowers a physician to perform safe, effective and efficient ablation of tumors and to facilitate prediction of the outcome. In TRANS-FUSIMO (www.trans-fusimo.eu) the consortium is currently developing a prototype of a fully integrated system for the FUS treatment of the liver, which will be evaluated in an animal study and a patient study.

Methods

The TRANS-FUSIMO system comprises three sub-systems:

1. a planning software that incorporates image processing, motion modeling and prediction, as well as FUS simulations. This part of the system allows to plan and to assess the feasibility of the treatment;
2. a treatment system that performs the actual treatment control using previously evaluated treatment plans, as well as model based MR/US motion tracking and prediction and MR thermometry (Fig. 47);
3. a training system which acts as a treatment simulator that is based on real cases and the simulation of FUS propagation, tissue heating, and organ motion.

Thereby, the treatment system fully integrates with the GE MR scanner and the InSightec ExAblate Conformal Bone System (CBS). The control system and the model components have been validated in phantom and ex vivo experiments. Currently, the safety, efficacy and efficiency of the system are being evaluated in an in vivo animal study. Following, a two-arm study (neoadjuvant TRANS-FUSIMO MRgFUS + resection, TRANS-FUSIMO MRgFUS only) for human patients with metastases or HCC will show the feasibility of the TRANS-FUSIMO prototype for the clinical setting.

Results

The core of the TRANS-FUSIMO system comprises patient specific models for the motion of the liver under breathing, ultrasound propagation, and thermal damage. On the one hand the models are used for the planning of the treatment and in the training system. On the other hand the models are used to augment US/MR data during the treatment in order to achieve motion compensation and thermal monitoring.

Conclusions

The ex vivo validations show the safety and efficacy of the system under regulated motion of phantoms. Furthermore the validations show that the system fulfills all specification requirements. The currently running animal study will be the basis for a following two arm patient study.

The TRANS-FUSIMO system comprises three sub-systems:
(PAA) mixed with egg white and ex vivo material (i.e. porcine liver or breast chicken instead of PAA cylinders into an Agar structure).

The complete workflow of the FUTURA procedure is reported in Fig. 49; more specifically, the effective use of two US imaging probes during different phases of the lesion treatment has been carried out. The sonication is composed by 10 repetitions of 1 second with 90 % duty cycle. The value of frequency and acoustic power of the sonication are 1200 kHz and 120 Watts, respectively.

Results
During the target identification phase, both the US imaging probes are used. Although the US confocal probe is rigidly attached to the HIFU transducer, the FUTURA platform maintains US imaging flexibility thanks to the 3D US probe mounted on the second manipulator. The 3D US phantom reconstruction and the related 2D US confocal images acquired before sonication show the target without any lesion. In addition, the estimated HIFU transducer focus, located in the center of the target, assesses the correctness in positioning of the therapeutic manipulator (intersection of blue lines in pictures B.I-II-III). During the sonication process an on-line monitoring of the lesion is performed by exploiting a PWM signal for the HIFU transducer generator. Finally, the 3D US phantom reconstruction and the related 2D US confocal images acquired after sonication allow the lesion assessment (i.e., accuracy better than 1 mm). An optical assessment on the sectioned targets in proximity of the performed lesion was also carried out (Fig. 50). Finally, a specific analysis of the HIFU properties has been performed by observing the typical dimensions of the lesions induced by the transducer into the breast chicken tissue at different output power and duration of exposure (Fig. 51).

Conclusions
The most important features of FUTURA platform have been demonstrated. This work assesses the possibility to use a robotic platform for USgFUS treatment, using two US imaging probe for the different phases of the procedure. Synchronizing the HIFU transducer shot with the US imaging acquisition enables to monitor the lesion progress during the sonication phase, which is very important for patient safety. In addition, the use of both US probes during the sonication phase enables the tracking of moving 3D organs.

Future work will be carried out to assess the accuracy of USgFUS treatment under dynamic condition and a qualitative temperature estimation technique (e.g., by means of elastography) for the assessment of thermal effects will be integrated in the analysis.
Real time MR and US motion tracking for abdominal US/MRgFUS
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Objectives
The ablation with HIFU/FUS would be a fantastic tool for liver tumor
treatment and would lead to an invaluable benefit for patients suffering
from malignant liver metastasis or hepatocellular carcinoma. How-
ever, FUS of liver has still not yet reached a significant clinical spread,
particularly in the western world. There are two major challenges for
FUS ablation of the liver: the rib cage and the permanent motion of the
liver, e.g. due to respiration. Both, MRI as well as diagnostic US imaging
are suitable to provide real-time information about the liver position.
For several years, we have been developing and improving US and MR
based motion tracking for HIFU/FUS, e.g. currently in the European
TRANS-FUSIMO project (www.trans-fusimo.eu). A further goal of the
presented work is the automated detection of suited tracking features
and the assessment of feature quality on MR and US image data.

Methods
US or MR image streams are analyzed in real-time to track liver mo-
tion. In a first step, pronounced structures, like the diaphragm, or
landmarks/features like liver vessels are automatically detected or
manually defined on 2D US or MR-EPI images. For motion tracking, a
particle filter-based algorithm evaluates state hypotheses of local af-
line transformations to follow these features through the image stream.
Data fidelity is assured by not only tracking each landmark separately,
but also incorporating information about global motion. For each landmark, an estimate of the current position and the
current uncertainty of that estimate is provided. Features with high
uncertainty may then be ignored in following processing steps or re-
placed by more stable landmarks by the algorithm. For use in an ap-
plication, the output of the algorithm is sent to the HIFU/FUS
treatment unit, which can incorporate the data into further motion
compensating components. This may either be established through
direct beam steering of the FUS focus or by gating the therapy.

Results
US as well as MR based liver tracking was evaluated in different
phantom experiments and on data from volunteers. Figures 52 and 53 show US and MR liver images with sample data provided
by the algorithm. The arrows show the displacement of the fea-
ture related to a reference image due to liver motion. The discs indicate a relative measure of positional uncertainty. Automated
feature detection, e.g. of blood vessels in EPI liver images, en-
abled fast and reliable definition of tracking features. Automated
tracking quality assessment allowed a reliable detection of poor
quality features which were replaced to allow a stable and reliable
tracking over longer time periods. The mean tracking error on US
liver images was 1.5 mm in 2D and 2.79 mm in 3D with a process-
ing time of about 2 ms/frame. 1.7 mm was the mean error of MR
based liver tracking with a computing time of 2 ms/frame and
feature.

Conclusions
Quasi real-time liver motion tracking in 2D and 3D based on diagnostic
ultrasound image streams as well as on ultra-fast MR image data is
feasible, reliable and offers a sufficient precision for motion compen-
sated HIFU/FUS therapy. Besides regular and irregular respiratory mo-
tion also liver drifts due to peristalsis and gravity on longer time scales
can be compensated, since this technique is based on live image data.
Integration of the tracking into existing clinical USgFUS and MRgFUS
therapy units should be possible without great efforts. The presented
tracking method is currently being integrated into an MRgFUS therapy
setup for the treatment of liver tumors within the TRANS-FUSIMO pro-
ject. A first clinical trial is planned at the beginning of 2017.
Objectives
Pancreatic ductal adenocarcinoma (PDA) is one of the deadliest cancers with the lowest prognosis due to late diagnosis. PDA is characterized by an unusually dense stroma limiting chemotherapy perfusion. Harmonic Motion Imaging (HMI) assesses tissue mechanical properties by inducing localized oscillation resulting from a periodic acoustic radiation force. The amplitude of the induced displacement is directly related to the underlying tissue stiffness. The sonication is kept short for imaging (HMI) without tissue damage or the duration is prolonged for simultaneous HMI and HIFU treatment (Harmonic Motion Imaging for Focused Ultrasound or HMIFU).

This study first aimed at using HMI for characterizing the development of pancreatic tumors as a function of size and fibrosis extent in the KPC genetically-engineered mouse model. Secondly, the tumors were treated using HMIFU. HMI measurements were then resumed to monitor the mechanical changes resulting from the treatment.

Methods
A 4.5-MHz Focused Ultrasound transducer (FUS) generated an amplitude-modulated beam resulting in harmonic tissue oscillations at its focus. Axial tissue displacement was estimated using 1D cross-correlation of RF signals acquired with a confocally aligned, 7.8-MHz diagnostic transducer (P12-5, ATL) using a plane-wave beam sequence at a framerate of 1 kHz. Imaging was performed with 0.2 s sonication for each scan position, when treatment required 60 s long sonifications which were shown to generate lesion in this model according to previous work by our group. KPC mice were genetically-engineered to develop pancreatic tumors with pathophysiological and molecular features similar to those of human PDA. Pancreatic tumor growth was monitored for 15 days with HMI scans performed every two days. For the second part of the study, HMIFU was performed on 5-mm tumors. The success of the treatment was assessed by measuring both the tumor area and its elasticity up to 14 days.

Results
HMI demonstrated its capability to provide reproducible elasticity measurements in murine pancreatic tumors. Figure 54 shows that stiffening occurs progressively during pancreatic tumor growth from the very early stages. When plotting the HMI displacement against the tumor size, an exponential trend was fitted to the data with $R^2 > 0.87$ in accordance to general models. When ablated with HMIFU for 60s, the tumor stiffened displaying a decrease in HMI displacement (Fig. 55). The lesion was confirmed by histology. The follow-up of the HMIFU treatment is performed with HMI to assess long-term tissue mechanical changes after treatment.

Conclusions
These results demonstrate the capability of HMI to provide elasticity measurements in the KPC murine pancreatic tumor model. The technique monitored the tumor growth as well as its stiffening when treated with HMIFU. Post-treatment mechanical changes could also be assessed with HMI. This study underlines the potential of HMI for monitoring tumor growth, treatment and follow-up changes in elasticity.

Fig. 54 (abstract A46). Pancreatic tumor development in 3 KPC mice over time during the early growth. The HMI displacement which is inversely related to the tissue stiffness is plotted against the diameter measured on Bmode images.
implemented in COMSOL, was used to generate resulting temperature distributions in homogeneous and heterogeneous tissue models. Parametric studies were performed to investigate performance of different potential lens fluids, and to evaluate acoustic focal gain and heating as functions of applicator/reflector/lens dimensions and geometries. Hydrophone beam plot measurements were made for a proof-of-concept end-firing applicator, consisting of a mounted tubular transducer centered in a conical brass reflector and covered by a perfluorocarbon fluid lens with adjustable radius-of-curvature (ROC).

**Results**

Of the three potential lens fluids investigated (perfluorocarbon, silicone oil, and chloroform), perfluorocarbon offers the best acoustic transmission at lens focal lengths ranging from 0-100 mm, due to its greater speed-of-sound mismatch as compared to water or tissue. As demonstrated in Fig. 57, simulated achievable focal gain for either device configuration increases as a function of lens focal length to a maximum between ~20-50 mm, depending on the overall deployable reflector/lens size and tissue attenuation. Thermal simulations in homogenous muscle tissue with the end-firing configuration indicate capabilities of creating localized (~1 cm long X 5 mm wide) or volumetric (~3 cm long X 2 cm wide) ellipsoid thermal lesions at variable depths up to and beyond 5-6 cm in the tissue, as demonstrated in Fig. 58. Hydrophone beam plots of the preliminary proof-of-concept assembly illustrated capability of dynamically adjusting the focal plane depth between ~1-5 cm by adjusting the lens ROC, as shown in Fig. 59.

**Conclusions**

Preliminary theoretical and experimental investigations illustrate capabilities of using reflector and fluid lens assemblies coupled with endoluminal ultrasound to produce deeper and dynamically-variable acoustic focuses and thermal lesions.

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![Fig. 56 (abstract A47)](image_url)

**Fig. 56 (abstract A47)**. Endoluminal ultrasound applicators with deployable balloon reflectors and adjustable fluid lenses, in either a) end-firing or b) side-firing configurations.

![Fig. 57 (abstract A47)](image_url)

**Fig. 57 (abstract A47)**. Intensity gain along the central longitudinal axis for the end-firing assembly as a function of lens focal length, as simulated in water. The 1.5 MHz tubular transducer was 8 mm OD x 20 mm height, leading to a 45 mm reflector/lens aperture diameter.

![Fig. 58 (abstract A47)](image_url)

**Fig. 58 (abstract A47)**. Simulated temperature distributions and 52 °C contour (black) in muscle tissue using the end-firing assembly, with specifications as in Fig. 57. Sonication duration was for (a) 10 s and (b) 120 s at 5 W/cm² constant surface intensity input.
A48 Concurrent low dose Focused Ultrasound with chemotherapy in abdominal tumors

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Objectives
Systemic therapies for advanced stage IV abdominal tumors as standard chemotherapy regimens usually lack efficacy, and often this is substantially reduced by means of toxicity. Hyperthermic therapies have demonstrated efficacy alone by themselves and as well as concurrent treatments along with chemotherapy and/or radiotherapy. Focused ultrasound has been increasingly used as a method to enhance drug delivery. Several previous experiences have used this approach to improve tumor control in different types of tumors and with different chemotherapy drugs.

Methods
This is a feasibility study to increase delivery of the chemotherapy drug to the tumor. We have employed Ultrasound guided High Intensity Focused Ultrasound (USgFUS) along with concurrent standard chemotherapy treatments in a clinical setting with abdominal tumors. We have included advanced stage IV liver metastases from colo-rectal tumors, advanced stage IV pancreatic tumors and advanced stage IV retroperitoneal sarcomas. All patients were previously treated with and failed at least to 2 chemotherapy lines. Patients should not be candidates for tumor ablation. Patients will receive 3 FUS treatments concurrent with their chemotherapy infusions. FUS therapies are delivered the day of the chemotherapy infusion, the day before and the day after the chemotherapy. FUS treatments are on the range of 100 Watts for 45 minutes average. The objective is to cover maximum volumetric treatment.

Results
We measure performance status, complications rate, quality of life, symptoms control, pain control, and time to progression. Blood samples for functional studies and tumor markers as well as immunological studies: lymphocyte profile, immunoglobulins, complement, CRP were taken at daily basis during FUS treatments. MRI images were obtained previously and immediately after the chemotherapy. FUS treatments are on the range of 100 Watts for 45 minutes average. The objective is to cover maximum volumetric treatment.

Conclusions
Study is in progress. We will report updated results at the meeting.

A49
Clinical experience of intra-operative High Intensity Focused Ultrasound in patients with colorectal liver metastases. Results of a Phase II study

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Objective
Surgical resection is the only curative option for colorectal liver metastases (CLM), however, only 10 to 20 % of patients are suitable for surgery. High intensity focused ultrasound (HIFU) has been proven effective in a wide range of clinical applications. The liver is a particularly challenging organ for HIFU treatment due to the combined effect of respiratory-induced liver motion and partial blocking by the rib cage. A HIFU device based on toroidal transducers has been developed to enable the destruction of large liver volumes. Preliminary in vitro and in vivo studies demonstrated the feasibility, efficacy and safety of such HIFU ablations. This preclinical work is now translated into clinical practice. The aim of this study was to assess the feasibility, safety and accuracy of HIFU ablation in patients undergoing hepatectomy for CLM.

Methods
This study was a prospective, single-centre phase I/II study. The HIFU transducer has a toroidal shape and was divided into 32 concentric rings of equal surface (0.13 cm2). The diameter of the transducer and its radius of curvature were 70 mm. The operating frequency was 3 MHz. A 7.5 MHz ultrasound imaging probe was placed in the center of the device. The imaging plane was aligned with the HIFU acoustic axis. Nineteen patients were included in Phase I-Ila. In each patient two HIFU ablations were created. In step 1, ablations were centered in a target previously identified in ultrasound images. In step 2, ablations were created at distance from a target. Eight patients were included in Phase IIb and ten metastases (20 mm maximal diameter) were treated until now. HIFU ablations were created to ablate metastases with safety margins in all directions. The exposure time was adjusted from 40 seconds to 370 seconds according to the diameter of the metastases. For this first evaluation of the device, ablations were performed within the areas scheduled for resection. This study is registered with Clinical-Trials.gov (NCT01489787).

Results
The dimensions of ablations measured on ultrasound imaging were correlated (r = 0.88, p < 0.001) with dimensions measured on gross pathology. The average dimensions of HIFU ablations obtained in 40 seconds (Phase I) were a diameter of 21.0 ± 3.9 mm and a long axis of 27.5 ± 6.0 mm. The phase I-Ila study showed both that the area of ablation could be precisely created in the liver with a precision of 1-2 mm. It was possible to target about 90 % of the liver and all major anatomical structures. In Phase IIb, one metastasis of 10 mm in diameter was ablated in 40 seconds with safety margins. Using electronic focusing nine metastases of 2 cm in diameter were ablated with safety margins (>3 mm in all directions) in 370 seconds. The dimensions of these HIFU ablations were a diameter of 48.0 ± 4.9 mm and a long axis of 51.0 ± 3.4 mm. No lesions occurred in surrounding tissues. Two patients had a post-operative complication (pneumonia with pleural effusion and a urinary tract infection) not related to the HIFU treatment. There were no significant changes in hemodynamic and respiratory parameters.

Conclusions
This HIFU device safely achieved large and fast volume of liver ablation, with a precision of one to two millimeters. HIFU ablations of metastases with a diameter of less than 2 cm were successfully created with safety margins of at least 3 mm in all directions. This study is the first clinical use of intra-operative HIFU in patients with CLM. Reducing the risk of potential important complications associated with an extracorporeal approach was among the reasons we developed an intra-operative device. An open procedure seems also more appropriate since 15-20 % of additional metastases are discovered intraoperatively. From these preliminary results, intra-operative use of a HIFU toroidal transducer appears feasible, safe and effective in ablating liver metastases.

A50
[18 F]-FDG PET and contrast MRI for enhanced guidance of Focused Ultrasound ablation in a syngeneic orthotopic model of murine pancreatic adenocarcinoma

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Objectives
Intravenous injection of [18 F]-FDG and contrast MRI enable precise monitoring of liver metastases simultaneously. In this study, we used the technique to provide real-time monitoring of metastases for focused ultrasound ablation. The aim was to develop a method for precise monitoring of the placement of the transducer and the delivery of the focused ultrasound energy. The result was a technique for guidance of focused ultrasound ablation in a syngeneic orthotopic model of murine pancreatic adenocarcinoma.
Objectives
Pancreatic ductal adenocarcinoma (PDA) is the most lethal cancer and there is a desperate need for new therapeutic strategies. Due to its characteristic hypovascularity, dense stromal architecture, rapid progression, and local invasiveness, conventional therapies fail to cure PAC. Magnetic Resonance guided Focused Ultrasound ablation (MRgFUS) offers a platform to build combination therapies due to its ability to debulk tumors, enhance delivery of chemotherapeutics, and stimulate an immune response. Here, the feasibility of MRgFUS is evaluated in a syngeneic orthotopic mouse model of PAC developed from the KrasLSL-G12D/+; Trp53LSL-R172H/+; Pdx-Cre (KPC) model 1. Further, [18 F]-FDG positron emission tomography (PET) is used to enhance guidance.

Methods
Murine mT4-2D cells (gift from the Tuveson laboratory, Cold Spring Harbor, NY) were injected orthotopically by a sterile laparotomy (n = 5, C57BL/6 female mice). Mice were treated with MRgFUS on day 16 post-implantation with an MR-compatible anular array (Imasonic SAS, 3 MHz center frequency, 4.7 Watt acoustic power, 5.6 MPa peak negative pressure, 0.5 x 0.5 x 1.5 mm3 focal volume) and positioning system (Image Guided Therapy) in a Bruker BioSpec 7 T. Tumors were ablated with a diameter of 2 mm and scanning speed of 1 revolution per second for ~40s. In a separate group (n = 6), on days 10 (n = 3 + 1 wild-type) and 14 (n = 3 + 2 wild-type), mice were injected with 200 uCi of [18 F]-FDG, rested for 30 minutes, and scanned with a Siemens Inveon DPET for 30 minutes. Following PET acquisition, T1w images were acquired post gadolinium contrast (TE/TR/FA = 11.7 ms/750 ms/180°, 4.3 x 4.3 cm2 FOV, 256 x 256 matrix, 17 slices). Mice were euthanized, organs of interest were harvested, and radioactivity was measured by Wizard 1470 Automatic Gamma Counter (PerkinElmer). Maximum intensity projections were registered with MR images and regions of interest (ROIs) were drawn on the tumors and normal pancreases using Siemens Inveon Research Workplace.

Results
[18 F]-FDG-PET images enhanced the visualization of the tumor rim and demonstrated the heterogeneity of uptake within the tumor (Fig. 60a). In spite of the heterogeneity, ROI analysis indicated that primary pancreatic lesions (60.6 %ID/cc) exhibit a higher maximum uptake of [18 F]-FDG than the normal pancreas (17.8 %ID/cc) at day 14 of tumor progression (p < 0.05). Biodistribution confirmed these measurements, with 15.5 %ID/g and 16.9 %ID/g in the tumor-bearing pancreas at day 10 and day 14, respectively, as compared with 5.2 %ID/g in the normal pancreas (p < 0.05). Contrast enhanced T1w MRI facilitated visualization of lesions within both the pancreas and spleen (Fig. 60b-c). MRgFUS ablation was successfully completed on the tumors and normal pancreases using Siemens Inveon Research Workplace.

Conclusions
[18 F]-FDG-PET images enhance the visualization of the tumor rim. Feasibility of MRgFUS ablation of a syngeneic orthotopic murine model of PAC was established. Future studies will combine [18 F]-FDG-PET with MRgFUS to guide therapies and monitor disease progression.
Objectives
Histotripsy is a form of therapeutic Focused Ultrasound that mechanically ablates tissue. Pre-clinical studies have explored the potential for histotripsy to treat fetal septal defects, deep vein thrombosis, liver cancer, and benign prostatic hyperplasia (BPH). Histotripsy technology is also being tested in a clinical trial for the treatment of BPH. The bubble clouds generated by histotripsy are hyperechoic, enabling standard B-mode ultrasound imaging to be used for image guidance of tissue ablation. The mechanical oscillations of the bubbles within the cloud generate acoustic emissions. Information about the amplitude and location of these acoustic emissions would provide an additional means to monitor the treatment progress from histotripsy. Passive cavitation imaging (PCI) is an ultrasound imaging modality currently under development that spatially maps the power of acoustic emissions generated by cavitation. In this study, the ability of PCI and plane wave B-mode imaging to predict the location and spatial extent of histotripsy

Methods
Histotripsy pulses were generated in a prostate tissue phantom with a 1-MHz, 8 annular element array with a 10-cm aperture and 9-cm focal length. All elements of the transducer were simultaneously driven in parallel by a custom class D amplifier and matching network. Histotripsy pulses over a range of pulse durations (5-20 microseconds) and peak negative pressures (12-23 MPa) were explored in this study. During histotripsy insonation of the phantom, a 128 element linear array controlled with an ultrasound research scanner was used to monitor cavitation activity with both PCI and plane wave B-mode imaging (Figs. 61 and 62). After application of the histotripsy pulses, the phantom was sectioned and stained with a periodic acid-Schiff stain to delineate the ablation zone. The passive cavitation images and plane wave B-mode images were co-registered with the histological images. A receiver operating characteristic curve was used to quantify the comparison between each imaging modality and the spatial extent of the ablation zone.

Results
The area under the receiver operating characteristic (AUROC) was significantly greater than 0.5 for both plane wave B-mode and PCI, indicating both approaches can be used to predict the spatial extent of the ablation zone. The AUROC and accuracy were significantly greater for PCI compared to plane wave B-mode imaging. Furthermore, the sensitivity for predicting lesion formation was a factor of 3 larger for PCI compared to plane wave B-mode imaging.

Conclusions
Overall, these results indicate PCI provides an improved prediction of lesion formation from histotripsy.

A54
Real time Tissue Changes Monitoring during the treatment of prostate cancer with HIFU
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Objectives
Backscattered RF ultrasound signals have been used clinically in differentiating normal and infarcted heart condition. Similarly, backscattered signals from pre and post hifu site are significantly different in magnitudes that provide correlation on degree of tissue changes. This is a validation study of Tissue Change Monitoring (TCM) algorithm performed with real-time thermometry. TCM performs spectral analysis of the RF backscattered (pulse-echo) ultrasound signals acquired before and immediately following the HIFU exposure. The RF signal analysis generates energy spectra of these signals and the difference of the energy magnitude is used as an estimator for tissue changes caused by the HIFU tissue ablation. TCM results are color coded and displayed on the each HIFU ablative site in Orange, Yellow and Green indicating tissue temperatures in the range from 75-100, 60-75 and 37-60 degrees C respectively.

Methods
Five (5) patients with histologically confirmed, organ confined prostate cancer were enrolled with approved protocol for this study. Four patients with focal cancer at stereotactic saturation biopsy had hemi-ablation only and one had a whole gland ablation. The Sonablate 500 HIFU device with Tissue Change Monitoring (TCM) software was used for the ablative treatment. Specifically designed 18 gauge needles containing three thermocouples per needle separated by a 1 cm distance were placed transperineally under TRUS guidance in the prostate. 4 or 5 needles were placed to cover the ablative sites. Temperatures from all thermocouples were recorded simultaneously at 0.5 seconds interval by a multi-channel thermometry system. Temperature data were acquired from a total of 56 sites,(26 focal zone sites, 22 posterior sites to the focal zone and 8 sites in the lateral gland of the prostate where there was no HIFU applied). Sixteen RF signals lines, pre and immediately post HIFU, for each ablative site were acquired at 50 MHz sampling rate. All RF data lines were processed by the SB-500 computer in the frequency domain to derive average energy spectra and change in magnitude for pre and post HIFU. The magnitude change and temperature from the site were used to derive cross correlation function and color code to overlay on the ablative site.

Results
The measured temperatures (Average, Max, and Min) in the HIFU treatment zones were 84, 114 and 70 degree C respectively. TCM energy readings were 1.05, 2.6 and 0.4 resulting in 83 % orange (75-100 degree C) and 17 % yellow (60-75 degree C) indicating an estimated average temperature of 91 degree C. Outside the focal zone, average recorded temperature was 50 degree C. The temperature

Fig. 61 (abstract A53). Comparison of B-mode image with the histotripsy ablation zone (outlined in blue). The bar in the lower right corner corresponds to 1 mm

Fig. 62 (abstract A53). Comparison of passive cavitation image with the histotripsy ablation zone (outlined in blue). The bar in the lower right corner corresponds to 1 mm
A55

Focal therapy with High Intensity Focused Ultrasound in low risk prostate cancer

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Objectives

Prostate cancer (PCA) therapy undeniably exposes patients to a risk of impotence and incontinence. Therefore, low-risk localized prostate cancer is increasingly managed with watchful waiting or active surveillance, although this management bears some psychological burden and the risk of under staging and under grading. Alternatively, High Intensity Focused Ultrasound (HIFU) therapy allows non-invasive partial ablation of detected PCA in a single session (focal therapy).

Aim of this study was to evaluate clinical efficacy and side effects of a single focal HIFU session in low risk PCA patients.

Methods

66 patients with localized prostate cancer who refused both definitive radical therapy and conservative management and had a strong desire to preserve potency were treated with focal HIFU. Men with strong obstructive symptoms, severe calcifications in the target area or severe prostatitis/abscess history were excluded. Treatments were performed with Ablatherm® integrated imaging.

Results

53 % of the patients had positive staging biopsies on the right, 47 % on the left lobe. Initial PSA (PSAi = at diagnosis) was median 5.58 (1.02-10.43) and increased to a PSA at treatment (PSAt) of median 6.01 (0.2-11.4) within 4 months (period of first diagnosis until HIFU treatment), corresponding to a median pretherapeutic PSA velocity of 1.29 ng spinal anesthesia in median 50 (27.75) min with 16 french urethral catheter in place. Applied HIFU lesions were median 138 (86-338). There were no significant therapy related intra-/postoperative side effects within the follow-up period of median 4.4 (0.25-9.85) years, according to CLAVIEN Score. Based on patient interviews 3 months after HIFU treatment potency was: worse (4.5 %), equal (93.5 %); better (0 %). PDE-5 inhibitors were used by 74 % of patients. Urinary continence was preserved in all cases, 3 patients used a safety pad before and after treatment. There were no rectal disorder, fistulae or rectal incontinence. Post-HIFU catheter time was (83.94 ± 27.87) minutes, the mean sonication number respectively recorded before and 1-week, 1-month, 2-month, 3-month after the treatment.

Conclusions

Focal HIFU - a non invasive single session therapy performed under spinal anesthesia - demonstrated high potency- and continence preservation while treating diagnosed prostate cancer lesions. There were no significant therapy related intra-/postoperative side effects within the follow-up period. Focal HIFU results in a significant deceleration of PSA velocity from pre-HIFU 1.29 to post-HIFU 0.12 ng/ml/year by treating only 25 % of the prostatic volume. PCA diagnosis and consecutive watchful waiting and active surveillance bears an imminent psychological burden for the patient. The option of therapeutic efficacy, the wish to avoid therapeutic side effects and to preserve potency as well influences patients’ decision for focal therapy with HIFU.

A56

Factors affecting the efficacy of the Magnetic Resonance guided Focused Ultrasound ablation for painful bone metastases: results from a multicenter study in China

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Objectives

Bone is the third most common target organ that carcinomas metastasize to after lung and liver. Bone pain induced by cancer metastases contributes substantially to morbidity and mortality in patients with malignant carcinomas. Magnetic Resonance guided Focused Ultrasound (MRgFUS) is considered as one of the most promising therapeutic modality for tumor ablation. But, not all patients could obtain satisfied therapeutic efficacy. Some factors such as the number of bone metastasis, KPS and NPIR score can affects the results. The purpose of this non-randomized study was to evaluate the factors affecting the efficacy of the Magnetic Resonance guided Focused Ultrasound (MRgFUS) ablation for painful bone metastases treatment, and the safety and efficacy of this new non-invasive treatment modality.

Methods

One hundred and thirty one patients with painful bone metastases were screened for this study from June 2014 to September 2015. Seventy-one patients were finally enrolled in our study. All 71 patients underwent MRgFUS and none of them received radiotherapy or chemotherapy for pain palliation in the past two weeks. The Numerical Rating Scale for pain (NRS), the Brief Pain Inventory (BPI-QoL) score, the Karnofsky performance score (KPS), morphine equivalent daily intake dose (MEDID), and the adverse events (AEs) were respectively recorded before and 1-week, 1-month, 2-month, 3-month after the treatment.

Results

1) Seventy-one metastatic bone lesions in 71 patients were treated by MRgFUS and the treatment data was as follows: the mean treatment time was (83.94 ± 27.87) minutes, the mean sonication number was (12.74 ± 7.30) minutes. 2) Adverse events included: pain in therapy area in 3 patients and sciatia in 1 patient, which were obviously reduced after physiotherapy. 3) The NRS before treatment and at 1-week, 1-month, and 3-month after treatment was 6.70 ± 1.82, 4.37 ± 2.55, 3.96 ± 2.83, 3.70 ± 2.81 respectively. The NRS significantly decreased after treatment at all time points (P < 0.01). 4) The BPI-QoL score before treatment and at 1-week, 1-month, and 3-month after treatment respectively was 5.70 ± 2.09, 4.62 ± 2.36, 4.42 ± 2.60, and 4.42 ± 2.70. The BPI-QoL score significantly decreased after the treatment at all time points (P < 0.01). 5) After treatment, the mean ΔBPI-QoL score improvement in patients with solitary bone metastasis was statistically superior to that in patients with multiple bone metastases (P < 0.05). 6) At 1-week and 1-month, the mean ΔBPI-QoL score improvement in patients with osteogenic bone metastasis was statistically superior to other two groups (P < 0.05).

Conclusions

MRgFUS can be used as a non-invasive, safe, and effective method for treating painful bone metastases. The patients with solitary bone metastasis or osteogenic bone metastasis can achieve better therapeutic efficacy than those with multiple bone metastasis or osteolytic bone metastasis.

A57

Can MRgFUS achieve local control of bone metastases?

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Objectives

Magnetic Resonance Imaging guided Focused Ultrasound (MRgHIFU or MRgFUS) treatment of bone metastases is effective and safe for pain palliation. In a few circumstances, MRgFUS showed a partial or
complete local control of the tumoral mass affecting bone. The aim of the work was to understand the efficacy of MRFgFUS in terms of local tumor control of bone metastases and to explore this opportunity and the reasons for success or failure.

Methods

Patients with painful bone metastases were enrolled and submitted to MRFgFUS (ExAblate 2100, InSightec Ltd, Israel), with imaging (CT/MRI) before and 3, 6 and 12 months after treatment. The primary endpoint was the number of lesions with partial or complete response at 3 months according to MD Anderson criteria.

Results

Out of 65 patients, 18 were lost and missed the 3-month imaging check. Forty-seven patients with 49 lesions were evaluated. The procedure was successful in terms of local control at 3 months in 24 lesions (49 %) – complete in 11, partial in 13. A stable disease was observed in 20 lesions and a progression in 5. Lesions with osteolytic (or mixed) pattern, with complete accessibility of the margins to the ultrasound beam are perfect candidate for ablation with the intent to control the lesion. Results were statistically independent of histology (primary cancer), previous radiation therapy (59 %), and size (up to 14 cm), though smaller lesions were generally more associated with complete or wide accessibility. Body mass index, age and sex of patients did not influence the outcome.

Conclusions

At present, MRFgFUS for painful bone metastases may produce a local control of the lesion. In the future, patients with bone metastases should be selected for MRFgFUS with three different intents, and the target should be clearly established before the treatment according to patient condition, prognosis, symptoms and to the features of the lesion: 1) palliation of pain; 2) palliation and local tumor control; or 3) ablation / local control.

A58

Effect of heating duration on therapeutic ratio in MR-HIFU hyperthermia mediated drug delivery using thermosensitive liposomes in rabbit Vx2 tumors

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Objectives

Localized drug delivery using MR-HIFU hyperthermia to trigger the release of chemotherapeutic agents from thermosensitive liposomes (TSLs) has the potential to increase the therapeutic ratio between antitumor effect and systemic toxicity. For doxorubicin (DOX), which is active in many pediatric solid tumors, the late effects of cumulative dose to the heart are treatment-limiting. In this study, we investigate the effect of mild hyperthermia duration (10 vs. 40 minutes) on the ratio of DOX deposited in heated tumors compared to cardiac muscle, using a clinical formulation of TSL-DOX in a rabbit Vx2 tumor model.

Methods

Rabbits had Vx2 tumor cells injected into each thigh 12 days before therapy. For each rabbit, mild hyperthermia (10 or 40 minutes, 42 °C) was delivered to a 10 mm diameter region in one tumor using a clinical MR-HIFU system (Sonalleve V2, Philips Healthcare) incorporated into a 3 T MRI (Ingenia, Philips Healthcare). Feedback control of hyperthermia sonication was performed by modified software designed for mild hyperthermia, using MR thermometry data acquired in 6 slices every 3.2 seconds. During heating, TSL-DOX (Thermodox, Celsion Corporation) was administered intravenously at 2.5 mg/kg over 5-6 minutes. Rabbits were sacrificed and perfused with saline for tissue harvest 3 hours after the start of TSL-DOX infusion (Fig. 63). Tissue DOX concentrations were quantified using liquid chromatography-mass spectrometry against a daunorubicin standard following homogenization in cell lysis buffer and extraction in silver nitrate.

Results

Vx2 tumors treated with 10 vs. 40 minutes of mild hyperthermia (n = 10 per group) had largest diameters of 24.3 and 29.9 mm, and body temperatures of 36.6 °C and 36.6 °C before sonication. Heating quality in the two groups was similar: target region temperatures averaged 41.9 and 41.8 °C, respectively, with T09 of 41.1 and 41.0 °C, and T10 of 42.7 and 42.7 °C. The average duration that target region voxels exceeded 40 °C in the 10 and 40 min heating groups was 11.3 vs. 40.7 min. In non-targeted organs (heart, lung, spleen, muscle) DOX concentrations demonstrated no significant differences between the two heating groups (ANOVA with selected Bonferroni post-tests). This indicates that systemic toxicity does not increase with heating duration (Fig. 64).

In both heated and unheated tumors, tumor DOX concentrations were higher for 40 vs. 10 min (25.4 vs. 14.0 μg/g and 6.7 vs. 5.4 μg/g, Fig. 65). Similarly, therapeutic ratio (DOX in tumor / DOX in heart) increased with heating duration (3.9 vs 2.2 times). Heated tumors had higher DOX concentrations and therapeutic ratios than unheated tumors (p = 0.03, 2-way ANOVA with Bonferroni). Therapeutic ratio and DOX concentration for heated tumors had a significant correlation with duration above 40 °C.

Conclusions

Our results confirm the theory that longer heating duration increases doxorubicin deposition in the targeted tumor without increasing doxorubicin accumulation in critical structures such as the heart. This will have an impact on hyperthermia protocols used in clinical trials of MR-HIFU hyperthermia mediated drug delivery.
A59
Technical and imaging parameters predicting successful Magnetic Resonance guided Focused Ultrasound treatment of painful osseous metastases
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Objectives
Magnetic Resonance guided Focused Ultrasound (MRgFUS) has proven to be effective in the treatment of painful osseous metastases. The results of the phase III randomized clinical trial comparing MRgFUS to sham treatment of bone metastases demonstrated a 64 % response rate in the treatment group, compared to a 20 % response rate in the sham arm (Fig. 66). We hypothesized that this intention to treat analysis included patients who were suboptimally treated. We reviewed images from the patient procedures to identify imaging features that indicated technically successful treatment, and assessed technical parameters for predictors of pain relief.

Methods
All images were anonymized such that the reviewer was blinded to the treatment outcome. Imaging features (size of tumor, osteolytic vs blastic tumor, location of tumor, T2 signal intensity, presence of edema around the tumor after treatment, devascularization of the periosteum around the tumor, tumor devascularization, etc.) and technical parameters of the treatments (number of sonifications, total energy, average energy, sonication time, type of anesthesia etc.) were documented for each patient. After de-anonymization, imaging features and technical parameters were then correlated with successful pain relief (defined as at least a 2 point decline in pain score without significant increase in analgesic medication).

Results
We identified an imaging feature that we call the black band (BB), which represents devascularization of the periosteum around the tumor, as best seen on subtracted contrast enhanced T1 weighted fat saturated images obtained immediately after treatment (Fig. 67). The presence of the black band correlated with response to treatment (Odds Ratio of Complete or Partial Response with BB = 5.97; 95 % CI: 1.08 – 33.0; p = 0.041; sensitivity: 96 %, 95 % CI: 88-99 %; specificity: 22 %, 95 % CI: 9 – 40 %). The presence of the BB after treatment predicted a successful treatment (positive predictive value = 72 %, 95 % CI: 61 – 81 %).

We also identified a simple technical parameter, the Energy Density on Bone Surface (EDBS), which is the total energy delivered normalized by the bone surface area. EDBS was the only parameter that independently correlated with the likelihood of achieving complete or partial pain relief (OR of CR/PR with EDBS > 5 J/mm2 = 5.51; 95 % CI: 1.70 – 17.8; sensitivity = 57 %, 95 % CI: 44- 69 %; specificity = 81 %, 95 % CI: 61- 92 %). The black band was significantly correlated with high EDBS (EDBS mean with BB was 5.7 ± 2.8 J/mm2; without BB was 2.7 ± 0.9 J/mm2, p < 0.001).

Conclusions
With a technically successful treatment (BB+, EDBS > 5), the positive clinical response rate was 86 %, higher than in the published report. Those sites that had previous experience using MRgFUS to treat bone metastases and that used deep anesthesia used the highest mean EDBS (6.8 ± 3.1 J/mm2, p = 0.013), and also had the highest response rate (88 %, p = 0.009, Table 3). We also noted that the response rate (22 %) after suboptimal treatments (BB-, EDBS < 5) was the same as in the sham arm of the trial. Utilizing this information will help optimize MRgFUS treatment of osseous metastases.
Objectives
Osteoid osteoma (OO) is a benign bone tumor that causes dull, but often debilitating pain, affecting mainly children and young adults. It is most commonly treated with computed tomography (CT)-guided radiofrequency ablation (RFA). However, RFA is invasive and CT guidance requires exposure to ionizing radiation. These shortcomings may be addressed through the use of Magnetic Resonance Imaging-guided High Intensity Focused Ultrasound (MR-HIFU), which allows non-invasive thermal ablation under real-time MRI guidance, with high spatial precision and without ionizing radiation. MR-HIFU may be used to reduce or eliminate pain through destruction of periosteal nerves and the OO nidus, without need for incisions or drilling. In an ongoing Phase I clinical trial, eight of the planned total of 12 children with symptomatic OO have been treated at the Children’s National Medical Center (Washington D.C.). Herein we report on the technical aspects, feasibility, safety, and preliminary efficacy of OO MR-HIFU therapy.

Methods
Treatments were performed using the Sonalleve V2 clinical MR-HIFU system (Philips, Vantaa, Finland). The system includes a 256-element phased-array transducer, a positioning system with 5 degrees of freedom, and integrated MRI receive coils. Prior to therapy, patients were anesthetized or sedated, and positioned on the HI FI tableau using gel pads, ultrasound gel, and degassed water for acoustic coupling. T2w and T1w MR images were acquired as a 3D stack, and subsequently used for treatment planning. Sonifications of 16-48 s duration were performed at a frequency of 1.2 MHz, targeting regions 4-12 mm in diameter with acoustic powers of 20-160 W. During sonifications, real-time MRI temperature maps were acquired using a fast-field-echo pulse sequence and the proton resonance frequency shift method, which is based on the temperature-dependent proton magnetic resonance frequency shift. The temperature maps were assessed qualitatively by MRI experts.

Results
Patellar positioning on the HI FI table was feasible in all eight cases, with some modification to the shape of standard ultrasound standoff pads to provide acoustic coupling to extremities. Lesion locations included the tibia (n = 3), femur (n = 3), talus (n = 1), and the great toe (n = 1). Distance between OO nidus center and skin ranged 1.1-7.7 cm, with estimated 43 ± 16 % power remaining at the bone surface due to soft tissue attenuation. Average acoustic power was 50 ± 30 W per sonication and total energy deposition per therapy session was on average 10 ± 7 kJ, resulting in complete lack of MR contrast enhancement of the OO nidus on T1w-imaging in six of eight patients, and partial enhancement in two patients. Immediately after the treatment, nonperfused volume extended 7 ± 4 mm into the bone and 4 ± 6 mm into adjacent soft tissue. For seven of the eight patients, complete or nearly complete resolution of OO-related pain followed within days and all patients were able to cease using medication 28 days following treatment. Sonifications, including cool-down times, took 30-77 min, with 101-195 min of MRI suite time required. The treatments required less than 4 hours of total anesthesia and/or sedation and they were well-tolerated without any serious adverse events.

Conclusions
With complete symptom resolution in all patients evaluated thus far, MR-HIFU has the potential to offer a fast, safe, completely noninvasive, and radiation-free treatment option for children with painful OO. Sonication parameters were chosen conservatively in this initial clinical trial, and overall treatment quality may be improved and its duration reduced through optimization of sonication power, duration, and frequency, as well as improved, or more specialized MRI equipment.

A61
Bone thermal modelling for MRgFUS of osteoid osteoma — validation with human clinical data

Objectives
MR-guided High Intensity Focused Ultrasound (MRgFUS) has recently been applied to patients with bone lesions such as osteoid osteomas (OO), which are painful benign tumours. While initial results are promising, there is much uncertainty about the ideal target location, effective thermal dose, overall bone heating and possible inadvertent injury. Simulation of MRgFUS heat transfer within and around bone can be used as a pre-treatment planning tool that could optimize and improve the safety and efficacy of this therapy.

Methods
To address the uncertainties of MRgFUS treatment, a computer simulator was developed. The simulator has three main components: 1) an automatic bone segmentation tool; 2) an acoustic propagation simulation to calculate the acoustic velocity distribution at the target; and 3) a heat distribution model, calculated by the Pennes Bioheat Equation, which simulates the temperature distribution at the target location. Concurrently, a clinical pilot study is being conducted, to treat OO patients with MRgFUS. The MR imaging and MRgFUS sonation data from the first four patients from this study was reprocessed with the simulator and the results were compared with the measured clinical thermal results.

Results
Automatically segmented femurs were compared to the manually segmented ones. The average volume overlap (VO) and sensitivity (S) between the automatic and manual segmentations were 92.2 % and 96.3 %, respectively. The calculations for VO and S are as follows: VO = (FP + TN)/Vm, and S = TP/Vm, where TP is the number of false positives. FN is the number of false negatives, TP is the number of true positives and Vm is the manually segmented volume. The average maximum temperature difference between the simulation and clinical data at a region near the sonication focus, after 20 seconds of sonication, was 1.40°C, after 40 seconds of cool down time, the difference was 7.30°C. The max temperature is defined to be the average of the hottest 9 voxels. The average distance between ablation centroids was 2.9 mm. Figure 68 represents the clinical MR thermometry distribution (left) and...
the simulated temperature distribution (right) from a representative sonication in one patient. Figure 69 represents the max temperature comparison over 60 seconds for this sonication.

**Conclusions**

The calculated segmentation validation metrics suggests that the automatically segmented data is comparable to the manually segmented data. The average difference between simulated and thermometry data at peak temperature suggests that the simulator can accurately predict the extent of ablation near the focus during sonication. However, the average cool down temperature difference suggests the simulation is consistently under-predicting the temperature near the focus after 40 seconds of cool down. The distance between ablation centroids suggests that the simulator can predict the site of ablation to within 3 image voxels. A robust MRgFUS bone thermal simulation platform has the potential to significantly improve efficiency, safety and outcomes of patients with bone lesions.

![Fig. 68 (abstract A61). Comparison of MR thermometry (left) with simulated results (right).](image)

![Fig. 69 (abstract A61). Max temperature vs. time of simulated and MR thermometry treatment data](image)

**Objectives**

Percutaneous or surgical therapy can lead to non-negligible side-effects and limited efficacy for paediatric population or young adults with a benign, self-limiting pathology such as osteoid osteoma. Over a period of 6 years we followed treated patients for at least 3-year for clinical, imaging and rehabilitation process. The aim of the study was to demonstrate that a completely non-invasive radiation-free focal ablation of ostoid lesions with MRgFUS can be a safe, effective and durable treatment option.

**Methods**

Patients were eligible for this dual-centre prospective development study if they had both clinical and radiological diagnosis of non-vertebral ostoid osteoma (typical nocturnal-worsening pain relieved with NSAIDs, pain score [VAS] >4, CT and MRI specific findings), and if they could safely undergo MRI, localized US procedure and anesthetic. Patients received non-invasive therapy using MRI guided high-intensity Focused Ultrasound, delivered to all known ostoid lesions, identified on MRI, CT, or both. Feasibility, patient safety (adverse events), and clinical relevance profiles of MRgFUS were considered as primary outcomes; tumour control at imaging was considered as secondary outcome. Analyses were done on a per-protocol basis. This study is registered with ClinicalTrials.gov, number NCT02302651.

**Results**

Among 50 subjects recruited with both clinical and radiological diagnosis of OO, 45 underwent MRgFUS treatment and were included in the analysis. After intervention, no complications related to treatment or anaesthesia occurred and every patient leaved our hospital within 12-24 hours. Even during follow-up, no long-term complication related to treatment was reported. Clinical benefit was effective in all patients and overall VAS median value shifted from 8 (IQR 7-9) points before treatment to 0 at 1-week, 1-month, 6-month, 12-month, 24-month and 36-month follow-up. Similarly, VAS Scores median values for sleep disturbances, overall physical limitation, sport limitation and daily activities limitation dropped to 0 within the first month after treatment, and didn’t increase in subsequent controls. Quality of Life, assessed with FACT-BP Score, rose from a median value of 28 (22-34) before treatment to 55 at 1-week, 59 at 1-month, 60 at 6-month, 12-month, 24-month and 36-month follow-up. There was a complete response (a zero pain score one month after treatment, without recurrent symptoms, stable at follow-up) in 80.00 % of cases. 4 patients required a second line of intervention. Overall, patients who reached and maintained a stable 0 VAS score during our 3-year observation with MRgFUS treatment alone were 86.67 %. At final follow-up, 12-month data were available for 77 % of patients. In the remaining 23 %, tumour control was assessed using additional imaging at 6-month, 12-month, 24-month and 36-month follow-up. Among 50 subjects recruited with both clinical and radiological diagnosis of non-vertebral ostoid osteoma (typical nocturnal-worsening pain relieved with NSAIDs, pain score [VAS] >4, CT and MRI specific findings), and if they could safely undergo MRI, localized US procedure and anesthetic. Patients received non-invasive therapy using MRI guided high-intensity Focused Ultrasound, delivered to all known ostoid lesions, identified on MRI, CT, or both. Feasibility, patient safety (adverse events), and clinical relevance profiles of MRgFUS were considered as primary outcomes; tumour control at imaging was considered as secondary outcome. Analyses were done on a per-protocol basis. This study is registered with ClinicalTrials.gov, number NCT02302651.

**Conclusions**

MRgFUS has shown complete absence of adverse events, persistent clinical efficacy and great tolerance profile. These strong features may induce a change in the current paradigm of OO management, making MRgFUS the treatment of choice, whilst CTgRFA role may shift to a second-line intervention.

**A63**

**MRI-guided High Intensity Focused Ultrasound treatment of ostoid osteoma in pediatric patients: preliminary results**

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**Objectives**

Osteoid osteoma (OO) is the most common benign bone tumor in children. The vascularized, central nidus (Fig. 70) contains cells that produce pain-inducing prostaglandin hormones. While pain can be managed with non-steroidal anti-inflammatory drugs, many patients choose to undergo CT-guided thermal ablation using radiofrequency or laser, the standard-of-care for definitive treatment. Minimally invasive thermal ablation techniques include risks from ionizing radiation, fracture, infection and transmitted thermal damage along the needle. Non-target tissue injury is a concern as temperature cannot be measured with CT, and the treatment...
induces temperatures around 90 °C that are maintained for several minutes. Magnetic Resonance guided High Intensity Focused Ultrasound (MRgHIFU) is an alternative modality to successfully treat OO. There is no mechanical penetration of the bone, reducing the chance of pathologic fractures and reducing the potential for infection. We present our initial experience with OO MRgHIFU treatment in six pediatric pilot study patients.

**Methods**

The study was performed to determine the impact of focused ultrasound treatment on bone pain, medication usage and health-related quality of life (HRQL). Standardized metrics, such as physical, emotional, social, and school functioning were collected through the use of age-appropriate and validated surveys (Pediatric Ouch, and PedsQLTM). An initial planning MRI was performed to ensure lesion accessibility/patient eligibility. MR thermometry (Fig. 71) was used during treatment to measure temperature in the target and surrounding tissue to ensure patient safety. Procedures were performed under general anesthesia or deep sedation. Post procedure, contrast enhanced MRI, to assess the non-perfused tissues that correspond to the ablated tissue volume (Fig. 70). Pain, HRQL and drug usage surveys were recorded on days 1, 7, 14, 30, 90 and 180 post treatment. Clinical visits on days 30, 90 and 180 included physical examination and diagnostic MRI of the target lesion.

**Results**

Between July 2014 and May 2016, 6 patients underwent 7 MR-HIFU procedures. Subject 2 underwent MR-HIFU twice due to an inadequate pain response after the first procedure. The average age of patients enrolled was 13 ± 4.1 years (range: 8 – 17) and weight was 52.2 ± 19.4 kg (range: 24 – 74). During this period, a total of 12 patients were referred for OO treatment. Reasons for exclusion from the study included acoustically inaccessible locations (proximity to nerve or growth plate) in 6 patients and patient refusal in 1. The overall technical success rate was 83 % (5/6 subjects). Subject 4 could not be treated due to poor acoustic access to the lesion. Complete response (treatment success) was seen in of 80 % (4/5) and partial response in 20 % (1/5) of the treated subjects. Pain scores dropped from an average of 7.8 prior to treatment to 1.2 by day 7, as shown in Table 4. Prior to treatment, the time to fall asleep was reported as 30-60 minutes. Following MR-HIFU, this decreased to 7-38 minutes. There were no significant adverse events.

**Conclusions**

This study shows that noninvasive MRgHIFU treatment of OO can be used to successfully treat osteoid osteoma in pediatric patients. When the cortex is intact, MR-HIFU heats the bone surface. Treatment of OO lesions results from heat conduction through bone from the cortical surface. In intact bone, MR-HIFU heats the cortical surface with deeper areas being heated by conduction. Subjects 1, 3 and 5 had superficial lesions and responded very well to MRgHIFU therapy. Subject 2 had a large, sclerotic medullary lesion that presented a challenge for heat conduction through the bone and required a second session with a more aggressive MR-HIFU treatment. Further research is necessary to define the optimal treatment parameters and limitations of MRgHIFU. A prospective registry is currently being designed to compare MRgHIFU with other types of thermal ablation.

**Table 4 (abstract A63), Pain scores pre and post HIFU treatment (at days 1, 2, 7, 14, 30 and 90) for all treated patients. Patient 4 could not be treated, and exited the study. Only the data from the first treatment of Patient 2 was used**

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age</th>
<th>Weight (kg)</th>
<th>1 Day</th>
<th>2 Day</th>
<th>7 Day</th>
<th>14 Day</th>
<th>30 Day</th>
<th>90 Day</th>
</tr>
</thead>
<tbody>
<tr>
<td>HIFU-O001</td>
<td>16</td>
<td>60</td>
<td>8</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>HIFU-O002</td>
<td>17</td>
<td>64</td>
<td>5</td>
<td>3</td>
<td>2</td>
<td>6</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td>HIFU-O003</td>
<td>8</td>
<td>24</td>
<td>7</td>
<td>4</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>HIFU-O004</td>
<td>17</td>
<td>74</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>HIFU-O005</td>
<td>10</td>
<td>58</td>
<td>10</td>
<td>6</td>
<td>5</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>HIFU-O006</td>
<td>10</td>
<td>33</td>
<td>9</td>
<td>2</td>
<td>3</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Average</td>
<td>13</td>
<td>52.2</td>
<td>7.8</td>
<td>3.2</td>
<td>2.2</td>
<td>1.2</td>
<td>1.2</td>
<td>0.8</td>
</tr>
<tr>
<td>SD</td>
<td>4.1</td>
<td>19.4</td>
<td>1.9</td>
<td>1.9</td>
<td>1.9</td>
<td>2.7</td>
<td>2.2</td>
<td>1.8</td>
</tr>
</tbody>
</table>

**Fig. 70 (abstract A63). Representative pre and post HIFU sagittal MRI of an osteoid osteoma lesion with gadolinium enhancement. The lesion is fully enhanced before HIFU treatment, and shows non-perfused regions following treatment.**

**Fig. 71 (abstract A63). Representative MRI thermal imaging during HIFU treatment from a 30 W, 20s exposure (axial and sagittal views). Temperatures in the nidus reached >70 °C, indicating thermal ablation of the lesion.**

**A64**

**Treatment of breast fibroadenoma with High Intensity Focused Ultrasound: update of feasibility study IDE #G130252**

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**Objectives**

Fibroadenomas are common in the United States and many patients find their fibroadenoma bothersome, and thus opt for surgical excision.

Study objectives are to evaluate the safety and feasibility of Ultrasound guided High Intensity Focused Ultrasound (USgHIFU) delivered by the Echopulse device (Theraclion, Paris) for treatment of breast fibroadenomas. General patient safety, cosmetic outcome, tumor response, patient experience, physician/operator experience, and device performance are being assessed.

**Methods**

Twenty female patients diagnosed with palpable breast fibroadenomas 1 cm or larger were enrolled in a single arm study and underwent treatment of their tumor utilizing a computer-driven, continuously cooled, extra-corporeal HIFU probe mounted on an arm moved by motors, and guided in real-time with an integrated ultrasound imaging scanner. The integrated probe is positioned by the operator and the lesion is imaged. Treatment planning is automated and presented for review and approval on an integrated computer screen. Patients had tumors meeting the following criteria: Distance from the skin of ≤ 23 mm to the posterior border of the
fibroadenoma, ≥ 5 mm from the anterior border of the fibroadenoma, and ≥ 11 mm from the focal point of the HIFU treatment. The chest wall must be more than 1 cm from the posterior margin of the tumor, and tumor volume must be between 0.3 cc and 10 cc. Subjects are assessed immediately after treatment and at 3, 6, and 12 months.

Results
The study remains open to accrual, the following are PRELIMINARY RESULTS, thus the denominators vary by data point: Enrolment at the time of the writing of the abstract is 16 patients. Six patients remain in follow-up. To date, there have been no grade 3 adverse events, nor any skin burns, persistent changes in skin appearance, or other significant toxicities/morbidities observed in the patients treated. Mean patient-rated pain score during treatment on a scale from 0 to 100 was 17.8. One patient has reported persistent mild pain at 6 months follow-up (2 on a scale from 0 to 100). The most common toxicity observed was pain (reported by 9 of 16 patients). Preliminary patient satisfaction was 4.65 on a scale of 1-5 (5 = most satisfied), 10 of 11 patients reported they would undergo the procedure again, and 11/11 reported they would recommend the procedure to a friend or family member. Reduction in the size of the palpable mass was reported by both the patient and evaluating physician in almost all cases. Similar findings were found on ultrasound in the majority of cases. Cosmesis can be excellent, and unchanged from baseline in all cases to date.

Conclusions
To-date, USgHIFU, delivered by the Echopulse device for treatment of breast fibroadenomas in the IDE G130252 study has been well tolerated by patients, resulted in minimal toxicity, and appears to have been effective.

A65
Long-term efficacy and tolerability of one or two US-guided HIFU treatment of breast fibroadenoma
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Objectives
Breast fibroadenoma (FA) is the most prevalent benign tumor, accounting for up to 70% of benign breast lesions [1,2]. They affect females in the reproductive period with two peaks of incidence in the third and in the fifth decade of life. During the follow-up, a minority of FA decrease in size or disappear, more than half of them remain unchanged, and some of them significantly increase [3]. Ultrasound (US-guided) high-intensity focused ultrasound (HIFU) is the only non-surgical and non-invasive procedure, where thermal destruction is achieved by precisely delivered energy to the target, without interrupting skin integrity. Recently, a multicentre study established that US-guided HIFU treatment of 51 FA resulted in 72.5% volume reduction at 1 year [4].

The purpose of our study was to compare the long-term efficacy and tolerability of one or two HIFU treatments in patients with breast FA.

Methods
Twenty patients with 26 FA were selected for US-guided HIFU. The therapy was performed with the system EchoPulse (TheracIon, France) on an outpatient basis, in one or two sessions, under conscious sedation. FA volume was assessed before and followed up to 24 months after the last HIFU treatment. After each procedure, adverse events were evaluated. Written informed consent was acquired from all patients.

Results
In 19/26 FA (73.1%) one HIFU was performed (group 1), whereas 7/26 FA (26.9%) received second HIFU (group 2) 6–9 months (median, 7 months) after the first session. In group 1 and 2, FA volume decreased significantly at 1-month (p < 0.001) and 3-month follow-up (p = 0.005), respectively, and continued to reduce until 24-month follow-up (p < 0.001 and p = 0.003, respectively). At 24 months, mean volume reduction was 77.32% in group 1 and 90.47% in group 2 (p = 0.025). Mild subcutaneous oedema was observed in 4 patients and skin irritation in 3 patients.

Conclusions
US-guided HIFU represents a promising non-invasive method with sustainable FA volume reduction and patient’s tolerability. Although one treatment is highly efficient, the volume reduction can be increased with second treatment.

References

A66
Harmonic motion imaging for characterization and Focused Ultrasound ablation monitoring of post-surgical human breast tumors
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Objectives
Each year in the United States, over 50,000 women are diagnosed with breast cancer. High-Intensity Focused Ultrasound (HIFU) holds promise as a non-invasive and targeted therapeutic technique for breast cancer patients. To facilitate its widespread translation to the clinic, however, there is still a need for a real-time and cost-effective device that can reliably monitor HIFU ablation. Harmonic Motion Imaging for Focused Ultrasound (HMIFU) is an all-ultrasound radiation-force-based technique, which can be used for real-time HIFU ablation monitoring by tracking relative stiffness change at the treated area without interrupting HIFU.

Methods
Specimen collection and handling of post-surgical breast tissues were approved by the Institutional Review Board (IRB) board of Columbia University and informed consent was obtained from all enrolled patients. HMIFU was performed in 10 normal, 10 malignant tumors (invasive ductal carcinoma, namely IDC) and one benign tumor (fibroadenoma, namely FA) specimens. The HMIFU setup consists of a 93-element, 4.5-MHz HIFU transducer and a confocally-aligned 64-element, 2.5-MHz phased array imaging probe, which is connected to an ultrasound imaging research system. All HIFU elements were synchronously excited by a 25 Hz amplitude-modulated signal to vibrate the tissue at 50 Hz. A GPU-based fast image reconstruction method was used to monitor lesion development in real-time.

Results
For real-time monitoring, the displacement map and lesion map were streamed on the computer screen at a display frame rate of 2.4 Hz during treatment without interruption. Following a 2D raster scan, 3D HMI displacement maps were reconstructed representing the relative stiffness of the tissue (Fig. 72). The mean HMI displacement within the ROI decreased by 60% from 24.73 ± 10.97 μm to 9.83 ± 6.46 μm in normal breast tissue (n = 10, p = 0.0048), decreased by 38% from 12.77 ± 3.50 μm to 7.35 ± 2.74 μm (n = 10, p = 0.045) in IDC and decreased by 20% from 2.56 μm to 2.06 μm (n = 1) in FA. There were statistically significant differences between before and after HMIFU ablation in both the normal and tumor specimens.

Conclusions
HMIFU is shown to be capable of differentiating relative stiffness between normal and abnormal breast tissues prior to treatment. HMIFU
Enhancing volume following the initial treatment. However, first treatment was complicated by a third degree burn along the far-field skin. Despite the complication, the family and referring clinicians wished to proceed with additional treatments. Enhanced safety measures were implemented to protect the far-field skin including water bags and fiber-optic temperature probes, as well as interval skin checks. The patient had four subsequent treatments over 14 months, without complication, with non-perfused volume of 90% most recently.

Case 2: Pretreatment tumor volume 740 cc. Notably, the sciatic nerve courses along the tumor’s anterior margin. Left lateral decubitus position was used to minimize the amount of energy through the sciatic nerve. Enhanced safety measures as above. First treatment resulted in a relatively low non-perfused volume of 30%, likely related to a prominent fascial band. Second treatment resulted in 75-80% ablation of the target.

Case 3: Pretreatment tumor volume was approximately 440 cc. The sciatic nerve was encased by the anteromedial portion of the mass. Left lateral decubitus position and enhanced safety measures again used. First treatment resulted in a relatively low non-perfused volume of 30%, likely related to low energies. Second treatment resulted in 70-80% ablation.

Conclusions
MRgFUS is an effective treatment for desmoid tumors with a favorable side effect profile, allowing for repeated treatments if necessary. Ablation size and safety can be improved by use of far-field coupling devices, careful patient positioning, and optimized sonication planning.

Consent to publish had been obtained from the patients and guardians.

A67
MRgFUS for desmoid tumors within the thigh: Early clinical experiences
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Objectives
Desmoid tumors are benign but locally aggressive tumors derived from fibroblasts. Surgery, chemotherapy, and radiation therapy have historically been the mainstay of treatment but recurrence is common and side effects can result in significant morbidity. Desmoid tumors of the thigh are particularly difficult to treat with conventional approaches and have a higher recurrence rate. MRgFUS has increasingly been used for management of desmoid tumors because of the favorable side effect profile. In this case series we highlight our experiences performing treatments in the thigh, including strategies for optimizing ablation size and safety.

Methods
Since December 2014, 14 MRgFUS treatments for desmoid tumors were performed at our institution in seven patients. 9 of these treatments were completed in three patients with large tumors within the posterior thigh. The first was a 7-year-old boy who had previously been treated with surgical resection, intraoperative radiation, along with courses of vinblastine/methotrexate and sorafenib. The second was a 21-year-old woman who had previously taken sulindac and celecoxib but had no other therapy. The third patient was a 14-year-old girl with no prior treatment. Treatment efficacy was evaluated by calculation of post-contrast ablation volume immediately following treatment and in diagnostic studies between treatments. Side effects and complications were also documented for each treatment.

Results
Case 1: Pre-treatment tumor volume 770 cc with 75% non-enhancing volume following the initial treatment. However, first treatment was complicated by a third degree burn along the far-field skin. Despite the complication, the family and referring clinicians wished to proceed with additional treatments. Enhanced safety measures were implemented to protect the far-field skin including water bags and fiber-optic temperature probes, as well as interval skin checks. The patient had four subsequent treatments over 14 months, without complication, with non-perfused volume of 90% most recently.

Case 2: Pretreatment tumor volume 740 cc. Notably, the sciatic nerve courses along the tumor’s anterior margin. Left lateral decubitus position was used to minimize the amount of energy through the sciatic nerve. Enhanced safety measures as above. First treatment resulted in a relatively low non-perfused volume of 30%, likely related to a prominent fascial band. Second treatment resulted in 75-80% ablation of the target.

Case 3: Pretreatment tumor volume was approximately 440 cc. The sciatic nerve was encased by the anteromedial portion of the mass. Left lateral decubitus position and enhanced safety measures again used. First treatment resulted in a relatively low non-perfused volume of 30%, likely related to low energies. Second treatment resulted in 70-80% ablation.

Conclusions
MRgFUS is an effective treatment for desmoid tumors with a favorable side effect profile, allowing for repeated treatments if necessary. Ablation size and safety can be improved by use of far-field coupling devices, careful patient positioning, and optimized sonication planning.

Consent to publish had been obtained from the patients and guardians.

A68
Anatomical distribution of pediatric sarcoma and neuroblastoma: Targetability with MR-HIFU
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Objectives
Not released for publication

Methods
Not released for publication

Results
Not released for publication

Conclusions
Not released for publication

A69
A prospective study on the efficacy of single High Intensity Focused Ultrasound treatment of patients with benign symptomatic thyroid nodule
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Objectives
Benign thyroid nodules are prevalent among the general population and some nodules may exhibit growth over time leading to compression symptoms or cosmetic concerns. Although surgery remains the treatment of choice for symptomatic thyroid nodule, it is associated with a 2%-10% risk of complications and requires a general anesthesia. The aims of the present study were to assess the efficacy of a single treatment of High Intensity Focused Ultrasound (HIFU) in reducing benign thyroid nodule volume and to evaluate the changes in health-related quality of life (HRQL) following a single HIFU treatment.
Methods
After obtaining IRB approval, consecutive patients with symptomatic thyroid nodule were assessed for eligibility. Inclusions were nodule(s): 1) without signs of malignancy (i.e. no suspicious clinical and ultrasonic features and benign cytology on fine needle aspiration); 2) measuring ≤10 mm on ultrasound (USG) in three orthogonal dimensions; and 3) malleable to HIFU. Exclusions were nodule(s): 1) measuring < 40 mm (by largest dimension); and 2) located < 2 mm from trachea, esophagus or recurrent laryngeal nerve (where ablation might pose thermal injury). Eligible patients were offered a choice of HIFU treatment, active observation or surgical resection. HIFU treatment was conducted with the USG-guided Echopulse (Theraclion SA, France). Primary outcome was a change in index thyroid nodule volume 6 months after HIFU. To have 80 % power and 95 % confidence interval (two-sided) to detect minimal important difference of 20 %, 20 patients were needed. Assuming a 10 % incomplete and withdrawal rate, 22 patients were required. Thyroid volume (mL) was assessed at baseline (i.e. before ablation), 1-week, 3-month and 6-month while HRQOL was assessed by the Chinese version of the SF-12v2 at baseline and 6-month. The HRQOL measured by both summary scores (physical and mental component summary) was then compared between those who received HIFU and those who chose active observation (controls) at 6-month.

Results
Over this period, 22 (52.4 %) chose to receive a single course of HIFU (HIFU group) while the other 20 patients chose to have active observation (controls). Among the HIFU group, the majority were females (90.9 %) and the majority (77.3 %) had their index nodule as the dominant nodule in a multinodular goiter. The mean base index nodule volume was 6.48 ± 4.34 mL (range: 0.92 - 16.76 mL). At 6-month follow-up the mean nodule volume decreased to 1.38 ± 1.31 mL (n = 22, p < 0.001). The average extent of nodule reduction was 71.58 ± 11.81 % (range: 54.45 – 90.09 %). However, there was no significant correlation between extent of 6-month volume reduction and basal volume (r = 0.096, p = 0.806), total ablation time (r = 0.503, p = 0.168), total energy delivered (r = 0.122, p = 0.755) or mean delivered energy per treated volume tissue (r = 0.150, p = 0.700). Compared with controls, the HIFU group achieved a significantly greater improvement in four quality-of-life domains (17.57 ± 6.46, p = 0.009 for role physical; 18.01 ± 6.78, p = 0.011 for bodily pain; 24.77 ± 8.15, p = 0.004 for general health; 32.61 ± 8.93 for social functioning), and physical component summary of the SF-12v2 (7.89 ± 2.83, p = 0.001).

Conclusions
USG-guided HIFU ablation is not only an effective and safe treatment option for patients with benign symptomatic thyroid nodules but has the potential of improving the HRQOL of patients who do not wish to undergo surgical resection.

A70
The effect of different treatment regimens with US-guided HIFU on thyroid nodule volume
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Objectives
Thyroid nodules prevalence can reach 67 % on ultrasound (US) examination. Although 95 % of them are benign, 1/3 show continuous growth and should be treated because of clinical symptoms or cosmetic concerns (1). Surgery is still the main therapeutic strategy, in spite of the fact that it carries 2–10 % risk of complications (2). US-guided high-intensity focused ultrasound (HIFU) is a non-invasive ablative technique, designed to decrease thyroid nodule size (3, 4). The purpose of our work was to compare the long-term efficacy and safety of one and two consecutive HIFU sessions for the ablation of benign solid thyroid nodules.

Methods
Twenty patients (mean age, 44.5 years) with euthyroid benign nodular goitre were treated with US-guided HIFU device (EchoPulse, Theraclion, France) under procedural sedation and analgesia. US volume measurement was performed at baseline, 3 and 12 months after the final treatment. Three months after the first treatment, 12 patients (group 1) continued the follow-up and 8 patients (group 2) received second HIFU treatment. Adverse events were evaluated after each HIFU procedure. Written informed consent was acquired from all patients.

Results
Concerning the baseline nodule volume and the energy applied per nodule volume, there was not significant difference between group 1 and group 2 (5.04 ± 2.70 mL and 4.83 ± 2.74 mL respectively; 3.5 ± 1.4 kJ/mL and 4.1 ± 1.6 kJ/mL, respectively). At 12-month follow-up the mean nodule volume decreased significantly in both groups (2.35 ± 2.44 mL, p = 0.003, and 2.63 ± 1.85 mL, p = 0.017, respectively) with a maximal volume reduction of 95.4 % and 66 %, respectively. The mean percent of volume reduction at M3 after the first HIFU and at M12 after the final HIFU differed significantly between group 1 and 2 (47.4 % ± 20.8 vs 24.2 % ± 15.8, p = 0.02 at M3, and 55.5 % ± 28.4 vs 46 % ± 22, p = 0.011 at M12). In two patients, transient subcutaneous oedema and mild skin redness were registered after the first HIFU procedure and after the second procedure one patient developed transient Horner syndrome, which resolved in 6 months.

Conclusions
The long-term effect of one and two HIFU sessions in solid benign thyroid nodules is comparable. Larger studies are needed to explain the different thyroid nodule susceptibility to HIFU ablation.

References

A71
Prostate cancer HIFU — novelty or innovation?
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Objectives
Not released for publication

Methods
Not released for publication

Results
Not released for publication

Conclusions
Not released for publication

A72
MR-HIFU ablation of osteoid osteoma: Experience in a pediatric tertiary care center
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Objectives
To describe our experience with Magnetic Resonance-guided High Intensity Focused Ultrasound (MR-HIFU) ablation of painful Osteoid Osteoma (OO) in children.
Methods
Nine children with OO (7 M, 2 F; 16 ± 6 years) underwent MR-HIFU ablation through an FDA and IRB-approved safety and feasibility clinical trial using the Sonalleve V2 MR-HIFU system. Treatment feasibility, patient safety, and clinical response were evaluated in all patients over 28 days. Additional follow-up was performed to evaluate longer-term safety and durability of clinical response.

Results
MR-HIFU therapy was feasible in all nine patients without any serious treatment-related adverse events. Eight out of 9 patients reported complete response in terms of pain resolution and cessation of medication usage and one out of 9 patients reported partial response. All treatments were performed on an outpatient basis without overnight admission.

Conclusions
These findings show that MR-HIFU ablation of OO is feasible and safe in pediatric patients and offers clinical response rates similar to those seen with radiofrequency ablation, the standard of care therapy at most US hospitals. However, the completely noninvasive and radiation-free nature of MR-HIFU is advantageous over RFA, particularly in children and adolescents for whom collateral damage and radiation exposure may cause long-term morbidity.
tail artery. One month post-RD procedure, animals were euthanized and kidney medulla norepinephrine concentration, a proven, robust metric of successful denervation, was obtained using an ELISA (Rocky Mountain Diagnostics). Histological analysis was performed on both renal arteries and surrounding tissues and kidney function, as measured by blood urea nitrogen and urinary albumin secretion, was evaluated.

**Results**
Acute blood pressure response: A transient decrease in MAP (>5%) was observed during 11 of the 36 monitored sonications (~30%). The normalized MAP measurements obtained during these 11 sonications are seen in Fig. 75. In all animals the effect was dependent on sonication location. In addition, the change in MAP occurred several seconds after the sonication start time indicating it was an accumulated effect. Treatment efficacy: Kidney medulla norepinephrine concentration was reduced by 36% (p = 0.05) one-month post ablation when compared to the sham animals, indicating RD was successfully performed. Efficacy was further confirmed by the presence of inflammatory cells, pyknotic nuclei as well as degraded nerve architecture in the histological results (Fig. 76). Treatment safety: Comparison of the treated and sham RD animal’s blood urea nitrogen and urinary albumin/creatinine ratio resulted in no significant changes indicating kidney function was not affected by MRgFUS renal denervation. In addition, mean T1 values in both kidneys did not significantly change during the MRgFUS RD procedure. The peak mean temperature rise measured by real-time MR thermometry in the back muscle (Fig. 77) located in the near field of all animals ranged from 15.5 to 25.0 °C, confirming energy delivery and targeting accuracy.

**Conclusions**
Controlling resistant hypertension with renal denervation using Focused Ultrasound has been demonstrated in this and other studies to be a safe and potentially efficacious procedure. This work indicates that an acute, systemic response that is a function of sonication position can be detected during MRgFUS ablation of the renal sympathetic nerves. Future work will further evaluate this response in a hypertensive animal model and assess the effect of ultrasound parameters.
A75

Trans esophageal HIFU for cardiac ablation: first experiment in non-human primate

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1INSERM U979, Paris, France; 2Francis Bessière, Alléz, France; 3IHU Liryc, Electrophysiology and Heart Modeling Institute, Pessac-Bordeaux, France; 4Station de Primatologie-UPS846-CNRS, in vivo tested acoustic coupling through the inflation of a latex balloon. The probe was 55 mm range from the transducer. Circulation of water at 5 °C inside the focal length. The focus could be steered electronically over a 15 to 40 mm elements truncated at 14 mm, 3 MHz operating frequency and 40 mm (Constanciel 2013) allowed setting the features of the HIFU transducer: 8ducer was built. Anatomical configuration and numerical simulations for that purpose, an endoscope integrating a 5 MHz 64-element com-
tination with trans esophageal HIFU in the heart of a non-human primate.

Methods

For that purpose, an endoscope integrating a 5 MHz 64-element commercial transesophageal echocardiography (TEE) probe and a HIFU transducer was built. Anatomical configuration and numerical simulations (Constancei 2013) allowed setting the features of the HIFU transducer: 8 elements truncated at 14 mm, 3 MHz operating frequency and 40 mm focal length. The focus could be steered electronically over a 15 to 55 mm range from the transducer. Circulation of water at 5 °C inside the endoscope ensured cooling of the front face of the HIFU transducer and acoustic coupling through the inflation of a latex balloon. The probe was tested in vivo in a 30 kg-baboon after preliminary TEE and CT-scanner demonstrated the acoustic access from the esophagus to the heart in this animal model. HIFU were delivered to the interatrial septum, the inferior wall of the left atrium and the upper posterior wall of the left ventricle (LV). A multi channel amplifier allowed delivering a focal intensity (Ipsa) of 3000 W/cm2 applied 4 times for 16 s at each selected locations. B-mode, shear-wave (SWE) and passive elastography were performed before and after HIFU with an ultrafast scanner. MR imaging (T1 mapping, contrast) was performed one-day post HIFU. Animal experimentation approved under agreement 04938.04.

Results

The endoscope was successfully inserted in the esophagus of the baboon. The inflation of the latex balloon allowed acoustic transmission from the esophagus to the heart and visualizing satisfactorily the cardiac structures by TEE. HIFU could be delivered at the locations identified on the CT scans even though the heart was slightly shifted in the chest due to different positions of the animal during CT and HIFU procedures. The ECG was not impacted by the treatments. No change in echogenicity of treated zones could be evidenced after HIFU. SWE showed a stiffening of the LV myocardial wall after ablation (two-fold increase of the myocardial stiffness), increase of stiffness could be observed simultaneously in the beating heart on passive elastography. MRI of the atrial wall for detecting thermal ablation could not be achieved. A hypo signal zone could be evidenced on T1 maps in the lateral wall of the LV where one treatment was performed. Contrast MRI did not show local fibrosis in the heart. Endoscopic examination did not reveal esophageal damage. The baboon recovered properly from the protocol and no significant side effect could be evidenced for one month after treatment.

Conclusions

This experiment was a first attempt to perform thermal ablation in the heart with a trans esophageal probe in vivo in a non-human primate. While the procedure was safe and lesions seemed to be induced in the heart, further development will be necessary in order to 1- Deliver more power locally by HIFU, to compensate for motion and heat sink by blood circulation and to generate more pronounced lesions, 2- Improve the performance of monitoring techniques for the atrium (sensitivity and resolution) and for the ventricle (difficult to generate shearwave farther than 50 mm from the esophagus).

A76

Ex vivo and in vivo non-invasive ultrasound-based cardiac pacing

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Objectives

Currently, no non-invasive cardiac pacing device acceptable for prolonged use in conscious patients exists. The main approach is invasive, employing intravascular catheters, which has associated risks. Focused ultrasound can be used to perform remote pacing using reversibility of electromechanical coupling of cardiomyocytes. This technique might be useful in the short term in the clinical settings in various conditions: temporary pacing for bradyarrhythmia or any clinical condition with risks of asystole; terminating or examining the inducibility of tachyarrhythmia; screening and optimization of cardiac resynchronization therapy. Here we described an extracorporeal cardiac stimulation device and study its efficiency and safety.

Methods

Ex vivo acoustic stimulation threshold was determined performing 756 sonications in 10 pig beating hearts. In vivo non-invasive stimulation was performed using 314 sonications in 4 anesthetized pigs. The animals were injected with ultrasound contrast agents. Experiments were performed using a Focused Ultrasound device (256 elements, 13/13 cm aperture/focal, operating at 1 MHz) under MR-guidance. At the end of each in vivo experiment, a navigated delayed inversion-recovery 3D Flash sequence was performed. Masson's staining was performed to assess acute damages screening from acoustic stimulation of both ex vivo and in vivo experiments.

Results

Using HIFU it was possible to perform ventricular continuous pacing (Fig. 78a) or to induce ventricular tachycardia (Fig. 78b). Consecutive stimulations of different heart chambers with a single ultrasonic probe was shown, allowing to modify the resulting atrio-ventricular delay (Fig. 78c-d). The results of the 756 stimulation sites performed in the right atrium, and the left and right ventricles in 10 ex vivo beating hearts from pigs were processed to determine stimulation threshold for pulse duration ranging from 30 μs to 10 ms. Two different pressure thresholds were highlighted: one around 4 MPa peak negative for HIFU pulses ranging from 50 μs to 1 ms (Fig. 78e). The same setup was used in vivo in 4 pigs to show clinical potential (Fig. 78f). Electrophysiological changes were also confirmed by arterial pressure modifications (Fig. 78g). The maximal peak negative pressure was estimated to be around 2 MPa at focus during in vivo experiments, due to the limited acoustic window. At this pressure level, stimulation of the LV was observed but with an insufficient success rate. Using ultrasound contrast agents, consistent cardiac stimulation was achievable for up to 1 hour sessions in 4 different animals. No damage was observed in the 4 animals.

Conclusions

To the best of our knowledge, this study is the first ex vivo and in vivo proof of feasibility of controlled non-invasive ultrasound-based cardiac stimulation in large animals. Preliminary safety results showed that this novel technology offers good prospects for clinical developments.
A77
Restoring perfusion after critical hindlimb ischemia using pulsed Focused Ultrasound and mesenchymal stem cell in aged mouse
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Objectives
Critical limb ischemia (CLI) is associated with a 5 year mortality rate in excess of 70 % with limited effective therapies. The goal of this study was to determine if pulsed Focused Ultrasound (pFUS) would enhance homing of mesenchymal stromal cells (MSC) to CLI in an aged mouse model and reestablish perfusion compared to pFUS or MSC alone.

Methods
CLI model was created by cauterizing the external iliac artery (EIA) on C3H mice (age = 10-12 months). Laser Doppler perfusion imaging (LDPI) was performed on lower extremities to confirm surgery and subsequently performed for 7 weeks post surgery. At 14 days post surgery, mice were divided into 4 groups: saline (n = 8), pFUS (n = 8), MSC (n = 8), and MSC + pFUS (n = 17). Mice received either 3 consecutive days of saline, pFUS, MSC, or MSC + pFUS starting on day 14 post surgery. 106 human MSC were administered IV. CLI mice (n = 3/group) were euthanized after one dose of either MSC + pFUS or MSC alone to determine if there was increased cell homing to ischemic muscle. pFUS exposures was performed with VIFU 2000 with a 1 MHz transducer; at 4 MPa; pulse repetition frequency, 1 Hz; DC 5 %; 101.0 ± 67.4 compared to mice that received pFUS + MSC, MSC in the ischemic limb was 306 ± 175 (p < 0.01 for MSC alone vs. pFUS + MSC). We also observed ~4-6 fold increases (p < 0.05) in human VEGF and IL10 expression in MSC + pFUS compared MSC alone groups.

Conclusions
This study demonstrates that pFUS enhanced homing of IV MSC to targeted muscle resulting in reperfusion and neovascularization in CLI model compared to MSC alone, pFUS preconditioning effects in the ischemic muscle stimulated local molecular changes in the tissue microenvironment that when combined with MSC infusion increased stem cell numbers and potency by producing increased amount anti-inflammatory faction that lead to increased perfusion compared to MSC alone. The ability of pFUS to modulate the molecular microenvironment in chronic diseased tissue opens the possibilities for enhancing cellular therapies in regenerative medicine and potentially improves clinical outcomes in patients suffering from CLI.
United Kingdom, Czech Republic, Germany, New Zealand, and Poland. 81 eligible study participants have been treated to date.

Conclusions

The Surround Sound system, which is developed to provide a one-time, non-invasive procedure to treat hypertension, has the potential to greatly reduce cost, lower risk, and improve access to millions of hypertension patients worldwide who are not adequately controlled by drug therapy. The initial clinical data have shown that ablation of renal nerves using Surround Sound system can result in profound and lasting reduction in blood pressure. The technology platform in which the system targets specific tissue while automatically tracking movement and continuously enabling therapy delivery through a custom shaped high power therapeutic phased array, could be easily adapted and used in other clinical applications.

A79

Ultrasound ablation enhances nanoparticle accumulation and survival in mammary carcinoma models

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Objectives

Magnetic Resonance-guided Focused Ultrasound (MRgFUS) ablation is a noninvasive method for treating solid tumors. However, ablation of the entirety of the tumor may not be possible due to constraints imposed by surrounding tissue, resulting in a small rim of viable tumor following thermal therapy. Therefore, augmentation of MRgFUS with chemotherapeutics such as liposomal doxorubicin may be necessary. These formulations reduce systemic toxicity by encapsulating non-bioactive copper-doxorubicin crystals which dissociate in the low pH of the tumor microenvironment. Our ultimate goal is to use Focused Ultrasound (US) within curative protocols in the mouse and in human medicine.

Methods

All studies were approved by the UC Davis Institutional Animal Care and Use committee. Tumors were generated in FVB/n mice expressing an activated form of ErbB2/neu, a system modeling human HER2 amplified breast cancer. These tumors were then transplanted into the mammary fat pads of wild type FVB/n mice. The results were validated in another mammary carcinoma model (4 T1 in BALB/c mice).

US ablation was accomplished under Magnetic Resonance (MR) guidance via either a 20 s point ablation or a 60 s point insonation in a circular pattern (16 element annular array with a 3 MHz central frequency). Temperatures >65 °C and CEM43 > 5000. Hyperthermia (41 °C) was induced using a 192-element transducer with 128 therapeutic elements. The desired regions of ablation produced durable response in 100 % of mice within 4 treatments (up to 50 °C). In the presence of MBs, application of FUS with lowpeak power density (up to 50 °C) and directional focused ultrasound (DFUS) with high peak intensities (up to 7.0); the encapsulated DOX is slowly released under hyperthermic conditions.

Results

Ablation enhanced accumulation of 64Cu-labeled liposomes as assessed with positron emission tomography (PET) and autoradiography following each of the ablation protocols. Accumulation of 64Cu-labeled liposomes was assessed with positron emission tomography (PET) and autoradiography following each of the ablation protocols.

Conclusions

In conclusion, initial results for the combination of hyperthermia, ablation, and liposomal doxorubicin in a fully immunocompetent aggressive syngeneic model demonstrated multiple opportunities to achieve a durable response in local disease.

A80

Targeted drug delivery with cyclodextrin-based nanocarriers and Focused Ultrasound triggering

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Objectives

The Nanoporation project set out to explore specific solutions to overcome the current challenges of targeted drug delivery (TDD) to tumours using Magnetic Resonance Imaging guided Focused Ultrasound (MRgFUS) to cavitate microbubbles (MBs) for increasing cell permeability and to open ‘drug nano-capsules’ to release proven active anticancer drugs directly to the tumour site with reduction of systemic drug dosage needed for the desired therapeutic effect. The work reported here aimed to develop novel nano-carriers for existing anticancer drugs, by establishing of human cancer cell models to evaluate the carriers’ encapsulation efficiency in vitro and in vivo, by using animal models and a clinical MRgFUS system to investigate the carrier-drug vehicles’ in vivo distribution and localised drug release / cellular drug uptake.

Methods

A novel y-cyclodextrin (y-CD) based carrier for encapsulation of doxorubicin (DOX) was synthesised and fully characterized (Figs. 81 and 82). The encapsulation efficiency was assessed under various temperatures and pH levels by both chemical analysis and in vitro human cancer cell modelling with KB and HCT116 cells. A computer-controlled high-throughput in vitro FUS device facilitating exposure to High Intensity FUS fields in a standard 96-well plate was designed and applied. It allows maximum flexibility in choosing experimental parameters in combination with carrier-DOX inclusion. Sonovue® MBs was used to investigate TDD in cell monolayers. An extensive in vitro study had been carried out to investigate both mechanical and thermal effects of ultrasound. An unique MRgFUS system compatible animal house were designed and built to allow ex vivo and in vivo experiments by using small rodents (Fig. 83). Ex vivo and in vivo trials were carried out with a clinically approved ExAblate MRgFUS system (InSightec, Israel) to establish a safe and efficient clinical TDD protocol on small rodents (Fig. 84).

Results

The desired y-CD based carrier greatly reduced DOX’s toxicity and the carrier-DOX inclusion was highly stable under physiological temperature conditions as well as under a wide range of acidic conditions (pH 1.0 ~ 7.0); the encapsulated DOX is slowly released under hyperthermic conditions (up to 50 °C). In the presence of MBs, application of FUS with low mechanical indexes, under which no thermal effect was observed, enhanced the drug uptake into tumour cells for both encapsulated and free DOX. Optimal setups of MR parameters and FUS parameters were identified ex vivo and in vivo, allowing application of MRgFUS treatments to 4 live mice bearing tumours (human colorectal carcinoma, up to 1059.71 mm3) under anaesthesia with full recovery.

Fig. 80 (abstract A79). T1w MR localize ultrasound for (a) point, (b) circle, (c) grid ablation. MR thermometry following (d) 20 s point, (e) 60 s circle ablation. Autoradiography 48 hours after (f) point, (g) circle, (h) grid ablation. (i) Survival plot for all animal groups
Conclusions
The study demonstrated the possibility of translation of the constructed γ-CD derivative to potential clinical use as a delivery vehicle for DOX using combined thermal and mechanical release mechanisms by clinically applicable MRgFUS–triggered TDD with the potential for cancer therapy. It provides better understanding of the mechanism of and potential for this novel delivery approach: MRI-guided, ultrasound-mediated, site-specific drug delivery assisted by MB contrast agents. The chemical modifications and in vitro, ex vivo and in vivo preclinical studies discussed here represent only a glimpse into the future of clinical application. It is likely that this novel technology will enter the clinical arena in the near future, based on the ever-increasing scientific contributions from researchers.

Objectives
Thermosensitive liposomes (TSLs) have been reported to accumulate and deliver an effective therapeutic to tumours with the application of hyperthermia. However, there is limited understanding of the effect of pharmacokinetics, the time frames and the duration of hyperthermia applications. For instance, the timing between drug injection and application of focused ultrasound (FUS) needs to be optimised considering nanoparticle biodistribution in the tumour and drug release. Near-Infrared Fluorescence (NIRF) imaging offers an easy and sensitive method to follow the biodistribution of dye-labelled theranostic nanoparticles and drug release in real time and could act as a useful preclinical surrogate for other forms of imaging to help understand the mechanism of drug kinetics in the tumours. In the present study, we examined the biodistribution of novel dual MRI and NIRF-labelled thermosensitive liposomes (image guided; iTSLs), and we modulate their distribution with hyperthermia induced by a high-intensity focused ultrasound transducer (FUS).

Methods
We have synthesized lipids that can be used for imaging in mice by MRI and NIRF. We have prepared iTSLs with the optimum concentration of imaging lipids for thermally triggered release and we developed them in combination with a FUS treatment regimen (thermal dosing). Labeling of liposomes for imaging can provide substantial information of the mechanism of tumor uptake (post injection) and provide insight on the reasons why this uptake is more enhanced in FUS-treated tumors. We prepared iTSLs to encapsulate the anticancer drug topotecan (Hycamtin®), a chemotherapeutic agent which when released in vivo can be monitored by its intrinsic drug fluorescence. We have optimized drug encapsulation for maximum fluorescence signal difference (before/after release) for both in vitro and in vivo. FUS (TIPS Phillips) was applied using temperature feedback via subcutaneously placed fine-wire thermocouples to maintain hyperthermic temperatures. NIRF imaging was performed using multispectral analysis bioimaging (Maestro EX) following the emission of the NIRF lipid and topotecan. FUS was applied using imaging as guidance. FUS was applied 30 min post injection and the tumors were monitored in a separate group of animals FUS was applied at 30 and 90 min post injection. MRI imaging confirmed observations obtained with NIRF imaging.

Results
iTSL tumour accumulation was detected using NIRF imaging immediately after liposome administration, this is due to the enhanced permeability and retention effect. Mice were bearing tumours at both flanks allowing one tumour to act for control (no FUS treatment). FUS-induced hyperthermia (3 min at 42 °C, 30 min post i.v.) greatly enhanced liposomal uptake as seen by imaging. A co-localised, enhancement of topotecan fluorescence emission was also observed immediately after application of hyperthermia, indicating rapid thermally triggered drug release.
within the area of the tumour. Topotecan is used as an anticancer drug model and its intrinsic fluorescent properties allow to easily follow thermal drug release. The phenomena of increased iTSL accumulation and concomitant topotecan release appeared to be amplified by a second mild hyperthermia treatment applied 1 hour after the first. NIRF imaging detected the signal coming from the liposomes (NIRF incorporated lipid) indicating that liposomes accumulated in tumours post injection. This accumulation was substantially enhanced after each application of FUS. Topotecan fluorescence appeared transient in the tumour indicating phenomena of cell uptake and DNA binding. MR imaging also confirmed enhanced iTSLs uptake due to the FUS treatments in the same animals that were used using NIRF imaging (Fig. 85).

Conclusions
In this study we have formulated a theranostic novel dual MRI/ NIRF labeled thermosensitive liposome (iTSL) that encapsulates the anti-cancer agent topotecan and enables real time/diagnostic imaging, making use of pre-clinically and clinically relevant imaging modalities. Image-guidance in turn enables the application of brief, moderate intensity FUS treatments that greatly increase iTSL tumor uptake and set up the possibility for substantial FUS triggered topotecan release within the tumor volume. FUS hyperthermia applied during the half-life of the iTSL in plasma promotes iTSL accumulation in the tumors and takes advantage of the high concentration of the liposomal drug in plasma seen immediately after injection. Should these effects be translated to the clinic, this suggests substantial benefits to cancer patients and improvements of the standard of the chemotherapy treatments for both primary and metastatic tumors.

A82
Near infrared fluorescence imaging for Focused-Ultrasound mediated local drug delivery of doxorubicin
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Objectives
High-intensity Focused Ultrasound (FUS) is valuable clinical technique for non-invasive thermal tissue ablation. Its capability for localised drug delivery has also been investigated recently, with application to cancer therapies. The ability of thermosensitive liposomes (TSLs) to accumulate and deliver a therapeutic drug to tumours under application of hyperthermia has been well reported but there is limited understanding of the time frames and kinetics involved. For instance, the optimal delay (if any) between injection of the therapeutic and application of heat. Near-Infrared Fluorescence (NIRF) imaging offers a sensitive method to follow the biodistribution of dye-labelled theranostic nanoparticles in near real time and acts as a useful preclinical surrogate for other forms of imaging (such as PET or MRI).

Methods
Our iTSLs (imageable TSL) containing doxorubicin (0.8 mg/mL) were formulated with different molar ratios of key lipids, and a dye-labelled lipid - XL750-DSA. In vitro studies were performed to characterise the temperature-triggered doxorubicin release by measuring changes in its intrinsic fluorescence. In vivo experiments were performed on SHO mice bearing subcutaneous MDA-MB-231 tumours on both flanks. For bioimaging experiments, iTSLs were injected i.v. with and without any hyperthermia induced by FUS (TIPS, Philips) and controlled by three thermocouples placed around the tumour (Fig. 86). TSLs and doxorubicin were respectively monitored by recording the emitted fluorescence in the NIR and the blue regions of the spectrum. Finally, tumour sizes were measured by calliper for 3-4 weeks following treatment.

Results
In vitro release studies showed minimal leakage of the drug at 37°C over a 10 min period, while the release was almost immediate when the temperature was raised to 41°C. In vivo, a first hyperthermia event (3 min at 43°C) was applied on one of the two tumours before iTSLs administration. After 30 min, the iTSLs were injected i.v. (tail vein) and their tumour accumulation was monitored by tracking the NIRF signal. A second hyperthermia treatment (3 min at 42°C) was then applied 45 min later in order to release the doxorubicin. As a consequence of this regimen, the mice showed a very significant NIRF intensity increase in the treated area. At the same time, doxorubicin release was observed but only for a short period of time. The combination of this protocol & doxorubicin-iTSLs showed a significant impact on tumour growth, including tumour eradication in some cases (Figs. 87 and 88).

Conclusions
NIR fluorophores carried by the iTSLs allowed the monitoring of the liposomes’ biodistribution over a period of several weeks post-injection in our mouse model. Mild hyperthermia applications on the tumour greatly favoured nanoparticle uptake. The hyperthermia also had the effect of releasing encapsulated doxorubicin, therefore improving the drug delivery to the tumour. Finally, a strong correlation between thermal dosing and tumour targeting was observed with an increase of the NIRF signal in the heated area with a subsequent enhanced therapeutic effect.

Fig. 85 (abstract A81). NIRF and MRI imaging after dual FUS application

Fig. 86 (abstract A82). TIPS focused ultrasound (FUS); (left) overview of the equipment showing the water-filled transducer chamber, the thermocouple interface, and the control PC; (right) schematic of the in vivo configuration with the transducer (a) raised such that the ultrasound biconic (b) focuses just above the skin surface over the tumour (c). The mouse is surrounded with warmed, degassed ultrasound gel (d) and placed on an ultrasound absorbing mat (e) to prevent reflections off the table. Temperature monitoring is via two or three fine-wire thermocouples (f) implanted around the tumour.
A83
Interleaved mapping of temperature and longitudinal relaxation rate monitor drug delivery from temperature sensitive liposomes during MR-HIFU induced hyperthermia
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Objectives
Not released for publication

Methods
Not released for publication

A84
Ultrasound ablation transiently increases accumulation of small molecule gadoteridol within the ablated volume
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Journal of Therapeutic Ultrasound 2016, 4(Suppl 1):A84

Objectives
Magnetic Resonance guided Focused Ultrasound (MRgFUS) ablation is a noninvasive method to deliver a precise, and lethal, thermal dose to solid tumors. During thermal ablation, tissue temperature typically exceeds 50-60 °C and results in rapid coagulative necrosis. Such ablation often results in “heat fixing” of the tissue which is thought to restrict the diffusion of molecules into and out of the ablated region. In this study we used gadoteridol (MW 558D) to evaluate the extent to which small molecules could be delivered to tissue following thermal ablation.

Methods
All studies were approved by the UC Davis Institutional Animal Care and Use committee. MRgFUS ablation of syngeneic murine mammary carcinoma (FVB/n mice expressing an activated form of ErbB2/neu) was accomplished via a 60 s insonation in a circular pattern (16 element annular array with a 3 MHz central frequency, 3.1 MPa PNP). This ablation protocol resulted in temperatures >65 °C and CEM43 > 5000.

Animals were injected with intraperitoneal gadoteridol either immediately before or after treatment (0.05 mmol/kg) or with intravenous gadoteridol before or after treatment. Mice were then imaged with a T1w scan (TE/TR = 12.5/750 ms, FOV = 3.2 cm × 3.2 cm, MTX = 256 × 256, ST/SI = 1/1 mm, and 9 slices) at 0.5, 1.5, 3, 6, 20, and 48 hours following ablation. The ablated region and quadriceps were then manually segmented and the tumor-to-muscle (T/M) ratio measured. In addition, two control groups of mice were also imaged with T1w MRI, a set treated with circle pattern MRgFUS and but no gadoteridol and a set not treated with ultrasound, but injected with intraperitoneal gadoteridol.

Results
Immediately following ablation, the T1w signal of the ablated volume increased, beginning with the rim. The T1w T/M signal ratio progressively increased until 1.5 hours post-ablation where it peaked (Fig. 89). The T/M ratio remained elevated compared to pre-ablation values for up to 6 hours post-ablation. Control animals treated with the same ablation protocol without gadoteridol injection did not exhibit an increase in T1w signal post-ablation. At 20 and 48 hours post-ablation, the T1w signal had decreased to pre-ablation levels and re-injection of gadoteridol at these time points had no effect on tumor T1w signal.

Thus, the small molecule gadoteridol accumulated within the tumor, specifically within the ablated volume, for several hours following ablation.

Conclusions
The results suggest that there is a window of time post-ablation when small molecule drugs can be successfully delivered to ablated tissue.
A85

Pulsed Focused Ultrasound to kidneys induces Molecular changes that potentiate mesenchymal stem cells to further improve acute kidney injury

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Objectives

Pulsed Focused Ultrasound (pFUS) enhances homing of IV-infused mesenchymal stem cells (MSC) to kidneys during cisplatin (CIS)-induced acute kidney injury (AKI). pFUS serves as a neo-adjuvant to MSC therapy where sonicating kidneys with IV MSC leads to better AKI outcomes than MSC therapy alone. Nearly twice as many MSC home to pFUS-treated AKI kidneys, but >10 times more inter leukin (IL)-10 is produced by MSC that home to pFUS-treated kidneys. This result suggests that pFUS sonications modify the renal micro-environment to increase potency of MSC that home to pFUS-treated kidneys. Interferon-g (IFNg) increases MSC potency and is upregulated in kidneys by pFUS. This study investigated pFUS-induced IFNg expression in kidneys as an activator of MSCs that improves their therapeutic efficacy during AKI.

Methods

C3H or IFNg-ko mice received CIS (15 mg/kg ip), kidney pFUS (4 MPa; 5% duty cycle) and/or MSC (10^6 human MSC). IV MSC injections were performed 3-4 hr post-pFUS. siRNA was used to knockdown IL-10 in some experiments. Experimental groups included AKI only, AKI + pFUS, AKI + MSC, AKI + pFUS + MSC, and healthy mice. Mice received CIS on Day (D)0 and pFUS/MSC on D1. Mice were euthanized on D4. One-way analysis of variance used Bonferroni post-hoc with p-values <0.05 considered significant.

Results

Following pFUS to the kidneys of IFNg ko mice, more MSC homed to sonicated kidneys (~2 fold) but did not lead to improved AKI [serum blood urea nitrogen (BUN) or serum Creatinine (Scr)] outcomes compared to mice that received MSC injections alone (Fig. 90). Levels of BUN and Scr, as well as expression of kidney injury molecule 1 (KIM1), were all significantly reduced by MSC treatment alone, but not further reduced by combination pFUS/MSC treatment like was previously observed in wild-type mice. Furthermore, the production of IL-10 by MSC seen in wild-type mice treated with pFUS + MSC was absent in IFNg-ko mice. Lastly, the improved AKI outcomes using pFUS + MSC in wild-type mice were erased when mice were given MSC that had been pretreated with siRNA to knockdown IL-10.

Conclusions

pFUS creates a molecular zip code in AKI kidneys that enhances MSC homing. While MSC infusions alone improve AKI in IFNg KO mice, pFUS + MSCs did not replicate the additional improvements in outcomes seen in wild-type mice. While pFUS-independent mechanisms of AKI repair by MSCs do not require renal IFNg, the pFUS-dependent mechanism that yields improved recovery does depend on preconditioning of the parenchyma. IFNg upregulated by pFUS mediates the neoadjuvant function of pFUS. Furthermore, the IFNg upregulated by pFUS serves to enhance MSC function by increasing MSC production of IL10, which has previously been shown to improve AKI. These data demonstrate that an IFNg/IL10 molecular axis is activated by pFUS and critical for improved AKI outcomes. Moreover, these results provide justification in incorporating pFUS as a treatment to improve MSC therapy during AKI, which often has limited clinical therapeutic options.

A86

Use of focused beams of special structure for pushing and trapping kidney stones with acoustic radiation force

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Objectives

Focused ultrasound not only causes lesions in biological tissue, but can effectively push the medium in the focal region. This effect, associated with the ability of ultrasound to remotely deliver and transform momentum to an absorbing or scattering target, is called radiation force. Until recently, therapeutic applications of this effect did not exist, although it has been employed in ultrasound imaging, e.g. shear wave elasticity imaging. Previously, our team developed a technology to reposition kidney stones with radiation force. In a clinical trial, the technology was used to transcutaneously facilitate passage of small stones and to relieve pain by dislodging larger obstructing stones. While successful, the trial revealed a need for optimization of the ultrasound beam structure, frequency, and intensity to make it more effective.

Methods

Two aspects of the kidney stone repositioning were studied. First, we investigated the optimal ultrasonic beam diameter vs. stone size. Numerical modeling was performed to calculate the force acting on an elastic sphere in liquid from a focused beam. Simulations were
performed for a given acoustic power, changing the ratio of the beam width to the diameter of the stone. The second study investigated the feasibility of trapping and moving a stone in a direction transverse to the beam axis. Toward this goal, vortex beams were considered that produce a ring-like intensity distribution in the focal region with zero on-axis intensity. Theoretical modeling was performed using a previously developed algorithm. Acoustic trapping and manipulation of kidney stones in water was investigated using both a single-element transducer (combined with a sector-shaped phase plate) and sector arrays with operating frequencies 0.3–1.5 MHz. Human stones approximately 3–5 mm, as well as glass and aluminum beads, were displaced in a water bath.

**Results**

The numerical modeling indicated that pushing a kidney stone is strongest when the beam width is slightly wider than the stone diameter. This can be explained by more effective generation of shear waves inside the stone resulting from their effective coupling with the acoustic waves in liquid at the stone edges. During exposures with the vortex beams, stones could be drawn to the beam axis and then controllably translated along the surface in any direction transverse to the beam. The phase between sector elements could be used to control the vortex size and thus adjusted for trapping different-sized stones.

**Conclusions**

The intensity distribution in the focal region significantly influences the ability to manipulate kidney stones using radiation force. To achieve the strongest force with a predetermined acoustic power, it is necessary to select a mode in which the focal waist diameter slightly exceeds the stone diameter. For transverse confinement, it is convenient to use vortex beams radiated by sector arrays or single-element sources with an attached phase plate. In such an approach, the beam diameter can be controlled by choosing a proper phase distribution on the source surface.

**A87**

**Modulated Focused Ultrasound for treatment of de-myelinating axons in multiple sclerosis lesions — pilot animal studies**

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*Journal of Therapeutic Ultrasound 2016, 4(Suppl 1):A87*

**Objectives**

Multiple sclerosis (MS) is a debilitating disease of the central nervous system whose symptoms arise from de-myelination of axons within brain and spinal cord tissue. De-myelination can lead to loss of central and peripheral function, including vision and muscular problems. There currently exists no cure for multiple sclerosis. In this exploratory study, we seek to induce re-myelination of axons de-myelinated by an MS model using pulsed Focused Ultrasound (pFU). There exists a rich history targeting the use of ultrasound to temporarily and non-destructively activate central neural circuits. Our own work has shown that pFU can induce detectable electroencephalography (EEG) signals in the brain. Recent work by Gibson et al. (2014) who showed increased myelin thickness of neurons that had been activated by laser light in an optogenetic mouse model. It seems plausible, therefore, that ultrasound induced neural activation could affect myelin growth and thickness.

**Methods**

We hypothesize that pFU activation of axons within MS lesions in a rodent model will decrease de-myelination and increase re-myelination. To test this, we selected the cuprizone model for MS, which causes de-myelination in the corpus callosum of the mouse brain and is reversible, allowing us to assess the effects of ultrasound during the phase of disease progression and during the phase of disease recovery. Rodents undergoing the MS model received five consecutive days of pFU therapy and underwent EEG monitoring. We applied pFU at two different time points, first during the demyelination process and second (in separate groups of animals) during recovery from demyelination. The pFU therapy protocols were inspired by the optogenetic stimulation protocols of Gibson et al (2014), and consisted of 30 seconds of pFU application followed by 90 seconds of rest, repeated 15 times. We tested three different pFU transducers with center frequencies of 0.65 MHz, 1.09 MHz, and 2.0 MHz, all with Ispta of 1.2 W/cm²s. For each of those frequencies, we verified that each pFU protocol could activate brain. We collected Magnetic Resonance (MR) images at three time points and collected the brains for histological analysis at the end of the study.

**Results**

Initial results show that mice undergoing the MS model exhibit different shapes in their pFU-induced EEG signals than normal healthy mice. Further results of the therapeutic pFU treatment following analysis of MR and histological results will be presented.

**Conclusions**

If successful, this non-invasive therapy may lead to rapid advancements in the treatment of MS and other de-myelinating neurological disorders.

**A88**

**Noninvasive peripheral nerve stimulation via Focused Ultrasound in vivo**

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*Journal of Therapeutic Ultrasound 2016, 4(Suppl 1):A88*

**Objectives**

The leading technique to treat peripheral neurological disorders is currently through implantation of electrodes along the peripheral nerve and stimulating the nerve with electrical current. Recently, pulsed Focused Ultrasound (FUS) has been shown to elicit modulation of both brain and peripheral neurons. While the effects of the FUS brain stimulation experiments have been shown in vitro, the majority of the peripheral nerve stimulation studies have been in vitro, or only investigating the effects of FUS stimulation on the nerve itself. Thus far, there have been no studies determining if non-invasive stimulation of peripheral nerves with FUS can elicit physiological effects in vivo.

**Methods**

Peripheral nerve stimulation was conducted on both hind limbs of 13 mice under general anesthesia (pentobarbital 65 mg/kg). Both the imaging and stimulation transducers were mounted interchangeably on a 3-DOF positioning system with 1-mm resolution. Targeting of the sciatic nerve was achieved using a 1.28x2 element imaging probe with a center frequency of 18.5 MHz. A single element transducer (3.57 MHz center frequency, 90 % duty cycle, 1 kHz PRF, 5-10 ms sonication duration, 3.65 x 0.45 mm focal area) was used for stimulation of the sciatic nerve. Evoked EMG activity in the Tibialis Anterior muscle was recorded via two stainless steel electrodes sampling at 2 kHz. Gross pathology was used to assess safety of the procedure. Behavioral testing occurred an hour before, and 24 hours after FUS stimulation. Electrophysiology experiments were used as positive controls with electrodes directly stimulating the sciatic nerve while recording EMG from the Tibialis Anterior muscle. Negative controls were conducted via administration of 5 mg/kg lidocaine to the stimulation region as well as clipping the nerves downstream from stimulation.

**Results**

Stimulation of the sciatic nerve was successful eliciting both muscle movement and EMG activity. Two distinct responses to stimulation were observed, a fast and slow activation (2.1 ± 2.6 ms and 18.8 ± 5.5 ms average delay following onset of FUS stimulation). The fast activation was accompanied by an average higher peak-to-peak EMG response than the slow responses (0.86 ± 0.87, 0.19 ± 0.63 mV respectively). EMG spike duration for both responses were similar and not significantly different (6.9 ± 2.6, 7.1 ± 2.2 ms respectively). Administration of lidocaine IM to the target region eliminated EMG responses from FUS stimulation. An hour following lidocaine administration EMG responses were detected again with FUS stimulation of the nerve. Electrical stimulation resulted in similar EMG spike durations to FUS stimulation, although at significantly higher peak-to-peak responses (7.3 ± 1.6 mV average). Gross pathology and behavioral testing revealed no significant damage (no red blood cell extravasation, significant...
difference in distance traveled, rotations or gap distance) with FUS stimulation at the powers used for stimulation.

**Conclusions**

In this study, we show physiological effects of in vivo FUS stimulation of peripheral nerves in the mouse for the first time. FUS stimulation can safely elicit muscle activation with no short-term behavioral effects. Positive controls verified EMG responses similar to that of electrical stimulation while negative controls verified FUS was stimulating the nerves and not the muscle tissue. These results support the further investigation of FUS-based techniques for the treatment of peripheral neurological disorders.

**A89**

The mechanosensitive TRPC1 channel is activated by pulsed Focused Ultrasound to induce stem cell homing

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**Objectives**

Stem cell therapies are promising regenerative medicine approaches. Pulsed Focused Ultrasound (pFUS) induces microenvironmental changes in normal and diseased tissues that enhance homing and efficacy of intravenously-infused mesenchymal stromal cells (MSC) to further improve disease outcomes. pFUS/tissue interactions that induce molecular changes is unclear. Mouse muscle and kidneys were sonicated at increasing powers while passive cavitation and tissue displacement were measured. Tissue was measured for cyclooxygenase-2 (COX2) to correlate with ultrasound sounds, as COX2 signaling is critical for MSC homing. To test that mechanotransduction causes the necessary molecular changes, inhibitors of mechanosensitive channels were given before pFUS as well as pFUS to TRPC-knockout mice, as TRPC can function as a mechanosensitive receptor.

**Methods**

C3H or TRPC-ko mice received hamstring or kidney pFUS from a VIFU 2000 system. pFUS was at 1 MHz, 5 Hz pulse repetition frequency, 5 % duty cycle, and varying transducer output powers (ranging from 10-80 W). A built-in hydrophone passively detected cavitation. Mice were euthanized 4 hr post-pFUS and COX2 expression was measured by ELISA. GdCl3 (0.04mmoles/kg) or ruthenium red (RR) (0.01mmoles/kg) was given by intravenous injection before pFUS. One-way analysis of variance using Bonferroni post-hoc with p-values <0.05 were considered significant.

**Results**

Statistically significant increases in COX2 were measured at 20, 40, 60, and 80 W compared to untreated muscle. Mean tissue displacement correlated with COX2 expression. Statistically significant increases in cavitation were only observed at 60 and 80 W.

**Conclusions**

Mechanical pFUS forces interact with the mechanostretch receptor TRPC1 to drive molecular changes in tissue that are critical to stem cell homing processes. We have previously determined that COX2 expression is an acceptable proxy for molecular outcomes. At lower powers (20 and 40 W), cavitation from the sonifications was not detectable, suggesting that cavitation-independent mechanical forces (i.e., acoustic radiation forces) drive COX2 expression and thus cell homing. At higher powers (60 and 80 W), cavitation was detectable and COX2 expression was elevated compared to sonications at 20 and 40 W. It is unclear whether cavitation at these elevated powers drives the additional COX2 expression, or if it is the result of increased acoustic radiation forces at higher powers. Regardless, 40 W was maximum power we previously determined not to cause detectable tissue damage and from the point of view of pFUS use in regenerative medicine, would be the maximum power.
legs (Fig. 92). The number of MSCs that remained in treated legs was nearly 10 times greater than in contralateral muscle.

**Conclusions**

The application of TUS to skeletal muscle containing transplanted MSCs extends the resident time of live MSCs in tissue by slowing MSC death/clearance rates. This has profound implications for a wide range of MSC therapies because TUS is noninvasive, safe, and inexpensive. Therefore, such a routine and uninvolved procedure to extend the survival of transplanted MSCs could easily and dramatically improve clinical cell therapies, extending the functional lifetimes of transplanted cells and reducing the number of transplantations/injections required for disease treatment.

**Results**

Fluorescent dyes were loaded into RBCs with good stability (minor leakage over many hours). Fluorescent PFC nanodroplets demonstrated stable adherence to the RBC membrane. Upon a single ten-cycle 10 MHz ultrasound pulse, dye leaked from the RBC within a fraction of a second. Specific adhesion (~65-fold over control) of targeted RBCs to alphaVbeta3-coated surface was observed. Magnetic targeting demonstrated efficient pulling of magnetic RBCs in a tube and phantom, confirmed by photoacoustic imaging of ICG-loaded RBCs. RBC-nanodroplet complexes remained in the bloodstream for at least two hours following intravenous injection, as determined with Cr-51-radiolabeling.

**Conclusions**

Red blood cells decorated with perfluorocarbon nanodroplets rapidly release entrapped fluorescent dye after a short ultrasound pulse. As a drug carrier, modified RBCs stay in the bloodstream for extended time; they can be targeted by molecular or magnetic means, and monitored by photoacoustic imaging. Overall, PFC-RBCs may be an ideal drug carrier system for ultrasound-triggered drug delivery.

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**Objectives**

Red blood cells (RBC), when modified to operate as drug carriers, combine the biocompatibility of a patients' own cells, long circulation time, large volume to sequester the drug, and ability to accumulate at the target sites (e.g., tumor vasculature) via molecular targeting. Recently point-of-care devices for RBC drug encapsulation have received CE approval mark in Europe. However, triggered release of the drug at the desired site is still complicated; therefore, we present in this study ultrasound-triggered release of entrapped materials from RBCs, by the preparation of RBC decorated with superheated perfluorocarbon (PFC) nanodroplets, and subsequent rupture of RBC membrane with contents release.

**Methods**

Fluorescent dyes, calcein and indocyanine green (ICG) and magnetic nanoparticles were loaded inside RBCs via hypotonic loading. After incubation in hypotonic media, the loaded RBCs were resealed by PIGPA buffer (adenine, glucose, inosine and pyruvate). Unincorporated material was removed by centrifugation (1000 g, 5 min). Nanoparticles made of decalfluorobutane and dodecafluoropentane were prepared by sonication following by the repeated Nucleopor filtration under elevated pressure; they were stabilized with a positively charged lipid shell, made of PEG stearate, DSPC and distearyl trimethylammonium propane. Nanoparticles were attached externally to the membranes of dye-loaded RBCs. For molecular targeting, RBCs were decorated with biotin or alphaVbeta3-specific peptide cyclo(RGDfK), via insertion of the molecules anchored to the cell membrane via a PEG spacer. To characterize the efficacy of ligand attachment, fluorescein-PEG-lipid was used as a model. Magnetic sensitivity of iron oxide-loaded RBCs was confirmed with a N52 neodymium magnet. Peptide-mediated targeting was confirmed in a solid phase binding system, where RGDfK-PEG-DSPE-RBCs were incubated with polystyrene dishes carrying recombinant alphaVbeta3, or control albumin surface. Photoacoustic imaging of ICG-loaded RBCs was performed with a custom-built apparatus (Verasonics Vantage mated with a SpectraPhysics laser). Ultrasound-triggered contents release was monitored by fluorescence microscopy: dye-loaded RBCs carrying perfluorocarbon nanodroplets were subjected to sononation.

**A91**

**Red blood cells as ultrasound-triggered drug delivery vehicles: Activation via perfluorocarbon nanodroplets, targeting and photoacoustic imaging**


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**Objectives**

Red blood cells (RBC), when modified to operate as drug carriers, combine the biocompatibility of a patients' own cells, long circulation time, large volume to sequester the drug, and ability to accumulate at the target sites (e.g., tumor vasculature) via molecular targeting. Recently point-of-care devices for RBC drug encapsulation have received CE approval mark in Europe. However, triggered release of the drug at the desired site is still complicated; therefore, we present in this study ultrasound-triggered release of entrapped materials from RBCs, by the preparation of RBC decorated with superheated perfluorocarbon (PFC) nanodroplets, and subsequent rupture of RBC membrane with contents release.

**Methods**

Fluorescent dyes, calcein and indocyanine green (ICG) and magnetic nanoparticles were loaded inside RBCs via hypotonic loading. After incubation in hypotonic media, the loaded RBCs were resealed by PIGPA buffer (adenine, glucose, inosine and pyruvate). Unincorporated material was removed by centrifugation (1000 g, 5 min). Nanoparticles made of decalfluorobutane and dodecafluoropentane were prepared by sonication following by the repeated Nucleopor filtration under elevated pressure; they were stabilized with a positively charged lipid shell, made of PEG stearate, DSPC and distearyl trimethylammonium propane. Nanoparticles were attached externally to the membranes of dye-loaded RBCs. For molecular targeting, RBCs were decorated with biotin or alphaVbeta3-specific peptide cyclo(RGDfK), via insertion of the molecules anchored to the cell membrane via a PEG spacer. To characterize the efficacy of ligand attachment, fluorescein-PEG-lipid was used as a model. Magnetic sensitivity of iron oxide-loaded RBCs was confirmed with a N52 neodymium magnet. Peptide-mediated targeting was confirmed in a solid phase binding system, where RGDfK-PEG-DSPE-RBCs were incubated with polystyrene dishes carrying recombinant alphaVbeta3, or control albumin surface. Photoacoustic imaging of ICG-loaded RBCs was performed with a custom-built apparatus (Verasonics Vantage mated with a SpectraPhysics laser). Ultrasound-triggered contents release was monitored by fluorescence microscopy: dye-loaded RBCs carrying perfluorocarbon nanodroplets were subjected to sononation.

**A92**

**HIFU for targeted antibiotic delivery and therapy of chronic wounds and osteomyelitis**

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**Objectives**

Chronic non-healing wound infections require long duration antibiotic therapy, and are associated with significant morbidity and healthcare costs. This results in both systemic and local infection to deeper tissues (e.g bones), thereby requiring long duration treatment (generally >6 weeks), resection of tissues, and emergence of drug resistance. Novel approaches for efficient, readily-translatable targeted and localized antimicrobial delivery, as well as targeted ablation of pathogens are a critical clinical need. The objectives of this study were to: 1) develop low temperature-sensitive liposomes (LTLSs) containing an antimicrobial agent (ciprofloxacin) for induced release at mild hyperthermia (~42 °C) (Fig. 93), 2) characterize in vitro ciprofloxacin release, and efficacy against Staphylococcus aureus plankton and biofilms, and 3) determine the feasibility of localized ciprofloxacin delivery in combination with MR-HIFU hyperthermia in a rat model.

**Methods**

LTLSs were loaded actively with ciprofloxacin, and their efficacy was determined using a disc diffusion method, MBEC biofilm device, and scanning electron microscopy (SEM). Ciprofloxacin release from LTLSs was assessed in a physiologic buffer (serum and PBS), and in vivo in a rat model using MR-HIFU by fluorescence spectroscopy.

**Results**

Results indicated that >95 % loading of ciprofloxacin was achieved. Further, < 5 % was released from the LTLS within 15 min at baseline (25 °C) and body (37 °C) temperature, while > 95 % was released at 42 °C. Sonication of rat muscles resulted in accurate and homogeneous temperature
control within the heated ROI (42.0 ± 0.2 °C), with a 90th percentile (T90) and 10th percentile (T10) of 43.2 ± 0.3 °C and 41.0 ± 0.3 °C respectively. Precise hyperthermia exposures in the thigh of rats using MR-HIFU during IV administration of the LTSLs resulted in a 4-fold greater local concentration of ciprofloxacin compared to controls (free ciprofloxacin + MR-HIFU or LTSL alone). The biodistribution of ciprofloxacin in unheated tissues was fairly similar between treatment groups. Triggered release at 42 °C from LTSL achieved significantly greater S. aureus killing and induced membrane deformation and changes in biofilm matrix compared to free ciprofloxacin or LTSL at 37 °C.

Conclusions
This technique has potential as a method to deliver high concentration antimicrobials to chronic wounds.

A93
Effectiveness of MR-guided Focused Ultrasound treatment for the tender points in patients with painful bone metastasis or with chronic osteoarthritic pain
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Objectives
MR-guided Focused Ultrasound surgery (MRgFUS) has been recently demonstrated as a promising tool for pain alleviation of painful bone metastases (pBM) and lumbar facet joint pain. The mechanisms of pain alleviation are believed to be most likely local denervation of the nociceptors at the ablated area using the Focused Ultrasound. However, few previous studies have suggested an assessment method to estimate the denervation. Peripheral nociceptive sensitization resulting in bone metastatic pain and osteoarthritic pain is thought to make pain on pressure (tenderness) at the lesion sites too sensitive. Therefore, our hypothesis is that MRgFUS treatment of the sensitized tender points could improve the refractory chronic pain in patients with bone metastasis or osteoarthritis (OA). The aim of the present study is to clarify changes in the tenderness and the worst pain intensity at the lesion treated with MRgFUS.

Methods
We have conducted MRgFUS treatments using ExAblate® system (InSightec Ltd, Haifa, Israel) for 8 patients (mean age; 64 years old) with pBM, 9 patients (72 y.o.) with chronic lumbar facet joint osteoarthritic pain (LFP), and 11 patients (78 y.o.) with chronic knee osteoarthritic pain (KP). Patients with the worst pain intensity (worst) ≥ 4 on the Numerical Rating Scale (NRS) and tenderness at the lesion site were enrolled. Treatments were applied to the bone surface of the most painful tender point of the metastasis in pBM group, the dorsal area of these joints which selected on the basis of the 70 % reduction of pain with diagnostic block in LFP group, the bone surface around osteophyte of medial femorotibial joint tenderness in KP group. We assessed pressure pain threshold (PPT) using an electronic pressure algometer to evaluate tenderness at the lesion sites, and the subjective worst pain intensity with NRS. PPT is defined as the minimal amount of pressure where a sensation of pressure first change to pain. We evaluated changes in the PPT on the treated site and at the contralateral non-treated site as a control and NRS before and 1 week, 1 month and 3 months after the treatments.

Results
PPTs at the treated sites were significantly increased from 107 kPa [40-432] (median [min-max]) at the baseline to 271 kPa [94-534] at the final follow-up in pBM group, 280 kPa [66-427] to 462 kPa [207-818] in OALFJP group, and 156 kPa [50-249] to 246 kPa [114-427] in OAKP group. At the control sites, the PPTs were significantly higher than at the treated sites and there was no significant difference of PPTs between before and after the treatment in all groups. The NRS were significantly decreased from 6.5 [4-8] to 1 [0-3] in pBM group, 8 [4-9] to 1.5 [0-7] in LFP group, and 6 [5-9] and 3 [1-6] in KP group. The treatment responder, defined as reduction of 2 or more in the NRS without increase analgesic intake in pBM, and a 50 % or greater decrease in the NRS in OA group, were all patients in pBM group, 14 of 20 patients in OA group. In OA group, the PPTs at the treated sites were significantly increased from 223 kPa [50-427] to 326 kPa [184-731] at 1 week after the treatment in responders, but there were no significant differences in non-responders.

Conclusions
These results illustrate the excellent pain-relieving effects of MRgFUS treatment for the tender points in patients with painful bone metastasis or with chronic osteoarthritic pain. Significant increase of PPTs speculated successful denervation effect on the nociceptive nerve terminals of the treated area. PPTs measurement is a useful tool for quantitative evaluation after MRgFUS and might be a useful parameter for assessing a treatment’s effect.

A94
Passive acoustic mapping and MR thermometry for real-time multi-modality imaging of Focused Ultrasound ablation
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Objectives
MRI provides outstanding soft-tissue contrast and ability to map temperature which are ideal for monitoring of Focused Ultrasound therapy. However, at the high intensities required for ablation acoustic cavitation may be nucleated, leading to an enhancement in ultrasound-mediated heating. The likelihood of cavitation may also vary significantly between adjacent tissue regions, which has frustrated efforts to control ablation over large volumes. As the onset of cavitation is accompanied by characteristic acoustic emissions,
simultaneous acoustic mapping during treatment can provide critical information that may used for control of ultrasound output to achieve ablation of entire tumors. Previous work has shown that such monitoring is feasible within an MR scanner using post-processed ultrasound data. The objective of the current study was to optimize ultrasound data acquisition and processing to allow rapid real-time display of acoustic mapping data alongside MR thermometry.

**Methods**

An ultrasound imaging array was aligned with the focus of a 1.5 MHz HIFU transducer (Fig. 94). The two components were clamped together, immersed in degassed water, and connected to a MR-compatible positioning stage. Rabbits with subcutaneous VX2 tumors were anaesthetized and placed on a platform with a cutout for ultrasound propagation and acoustically coupled using a bag filled with degassed water. A receive-only MR surface coil was placed in the air gap between the water and animal platform. The entire setup was placed within a clinical 3 T MR scanner (GE Signa Excite). The HIFU transducer was then driven from a waveform generator via a power amplifier. During sonication the MR scanner performed temperature mapping while an ultrasound research platform (Verasonics) performed B-mode and passive acoustic mapping (PAM) using the array. Temperature maps, B-mode and passive acoustic images were displayed in real time. The process was repeated in rabbits injected with phase-shift nanoemulsions (PSNE) to promote cavitation or saline as a control under a range of ultrasound conditions.

**Results**

Real time passive acoustic mapping was successfully implemented using a rapid frequency-domain beamformer and employed simultaneously with MR thermometry for imaging of tumor ablation. Illustrative results are shown in Fig. 95. Over 50s (40s sonication, 10s baseline and cooling time) 477 frames of ultrasound data were acquired, processed and immediately displayed at an average frame rate of 10 fps. MR temperature maps were also acquired and displayed every 2.3 s (0.4 fps). Comparison of the two modalities (Fig. 95a) shows accelerated temperature rise following the onset of cavitation (30s) and decline thereafter as the amplitude of cavitation dropped (40s), perhaps due to depletion of cavitation nuclei or changes in tissue structure. MR temperature maps (Fig. 95b) and passive acoustic maps (Fig. 95c) show that heating and cavitation activity were both well localized to the intended focal region. Extraction of tumors after sonication showed lesions were successfully produced in the tumors.

**Conclusions**

These results show that rapid, real-time passive acoustic imaging coupled with PSNE for nucleating cavitation locally may be applied simultaneously with MR thermometry for enhancing Focused Ultrasound ablation. Future work will apply the method for optimization and control of PSNE enhanced ablation of whole tumors.
high spatial resolution scanning of the acoustic field inside the MR scanner.

Conclusions
In conclusion, HIFU pressure field measurements both outside and inside an MRI scanner provides a systemic method for ensuring safe and consistent treatment in a clinical MR-guided HIFU system. Moreover, the proposed inside MR scanner calibration method allowed characterization of the acoustic fields of a clinical MR-guided HIFU system without the need of moving the HIFU table outside of MR scanner, which is not feasible for most clinical systems.

Preservation of endometrium after Magnetic Resonance Imaging-guided High-Intensity Focused ultrasound (MR-HIFU) ablation for uterine fibroids of submucosal type
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Objectives
To evaluate the endometrial integrity after MR-HIFU ablation for submucosal uterine fibroids based on MRI findings and to know the risk factors associated with endometrial injury

Methods
A total of 117 submucosal uterine fibroids (diameter: 5.9 ± 3.0 cm, 1.0-12.9 cm) in 101 women (age: 43.6 ± 4.4 year, 33-55 year) treated with volumetric MR-HIFU ablation between November 2010 and December 2015 were retrospectively analyzed. Endometrial integrity was assessed with contrast-enhanced T1WI of immediate (n = 101), 3-month (n = 62), and 12-month (n = 15) follow-up. Endometrial injury was classified into one of grade 0 (continuous involved endometrial lining), grade 1 (pin-point, full-thickness discontinuity of involved endometrium), grade 2 (between grade 1 and 3) and grade 3 (full-thickness discontinuity of involved endometrium over 1 cm in size). Potential risk factors for endometrial injury (age, duration from LMP, history of full term delivery, GnRH agonist pretreatment, FIGO grading of endometrial protrusion, T2 signal intensity and perfusion degree of fibroid, and average acoustic power) were assessed with a generalized estimating equation (GEE) analysis.

Results
Among 117 fibroids, endometrial injury was of grade 0 in 66 (56.4 %), grade 1 in 29 (24.8 %), grade 2 in 16 (13.7 %), and grade 3 in 5 (4.3 %) at immediate follow-up (Fig. 99). Among 37 fibroids of which endometrium was injured and underwent follow-up studies, 30 (81.1 %) showed improvements at 3- and/or 12-month follow-up. Among 14 fibroids in which endometrium was injured and followed-up till 12 months, 12 (85.7 %) were completely normalized (Fig. 100). GEE analysis revealed that the degree of endometrial protrusion were significantly associated with severer endometrial injury (p < 0.0001, B = -0.687).

Conclusions
After MR-HIFU ablation of submucosal uterine fibroids, endometrium is preserved or minimally injured in majority of the cases. Injured endometrium which is more frequently encountered after treating intracavitary fibroids usually recovers spontaneously.
Results

The results of the type 2 fibroid treatment using MR-HIFU showed the potential of targeted vessel ablation. A similar strategy was used for type 3 fibroids. Based on the first treatment the high power sonication (maximum power of 230 W, 42-55 s) was repeated prior to treatment, spatially matched with the screening DCE and TimeSLIP sequence, and used for planning of the high power (450 W) sonications. A Philips Sonalleve V2 system with skin cooling (water temperature ~ 15°C) is used. After sonicating the inflow areas with high power ultrasound the bulk volume is treated with ‘low power’ sonications (£300 W). NPV is assessed on contrast-enhanced T1w-images. Follow-up MRI is performed after three months.

Conclusions

As a preparation a patient with a type 2 fibroid was treated in a similar manner except that all sonication were at a power (£300 W). Philips software was used to determine areas with a high wash-in rate (WIR) indicating the areas of high perfusion and these regions were treated.

Potential factors (age, BMI, history of full term delivery, multiplicity of fibroids, respectively) and retrospectively analyzed. Fibroid volumes and T2 signal intensity changes were compared between before and 2-3 weeks after the last GnRHa injection. Potential factors (age, BMI, history of full term delivery, multiplicity of fibroids) were evaluated for the association with greater volume and T2 signal intensity changes. All statistical tests were performed using model analyses.

Fig. 99 (abstract A96). Follow-up results of endometrial injury after MR-HIFU ablation for submucosal uterine fibroids.

Fig. 100 (abstract A96). Follow-up results of endometrial injury after MR-HIFU ablation for submucosal uterine fibroids.
A non-contrast method for evaluating non-perfused volume after MRgFUS treatment of uterine fibroids

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Objectives

The efficacy of MRgFUS treatment of uterine fibroids is evaluated post-operatively by measuring the non-perfused volume (NPV) on post-contrast images as an estimate of the ablated region. NPV is not a direct measurement of ablation but rather of non-perfused tissue. This may lead to an underestimate of the ultimately ablated volume, as some cell death may be delayed, or to an overestimate of the treatment effect, for example, in cases where temporary vasoconstriction contributes to the non-perfused volume. In addition, this can be confusing in cases where non-targeted tissue appears non-perfused; since there is no thermal dose predicted in these regions, it may be unclear whether the tissue is actually ablated.

A hypointense contour, which we call the Black-Line Volume (BLV), is present around the margin of fibroids on non-contrast images obtained at follow-up. Our aim was to quantitatively compare the BLV to the NPV.

Methods

Five patients with follow-up imaging after treatment were included in this study. Four of them had a second treatment within 7 to 14 days either because the fibroid was too large for a single treatment, or it appeared incompletely ablated. Patient 2 had no additional treatment. In three of the retreated patients, a reperfused volume was noted on follow-up images in regions that the NPV indicated as non-perfused at the end of the first treatment.

The NPVs obtained at the end of each treatment (NPV1 and NPV2) and the reperfused volumes were measured and compared by multiplying slice-by-slice measurements of the manually contoured area by the slice thickness. Using the same methodology, the BLV was calculated using EPI thermometry magnitude images acquired during each sonication in follow-up images.
Results
The results are provided in Table 5. In these patients, the BLV contour is not present prior to treatment, but is reliably visualized on follow-up imaging obtained 7-43 days after treatment. Quantitation of the BLV demonstrates that it is very similar to the NPV. In cases where the BLV was less than NPV1, the discrepancy is explained by reperfusion (Fig. 105d).
In three cases, at the end of a second treatment, portions of the fibroid that were part of NPV1 were seen to have reperfused. In these cases, BLV consistently identified the reperfused volume, which was outside of the BLV.

Conclusions
These results show that BLV correlates well with the ablated region, and the non-perfused volume. In cases in which NPV1 overestimated the truly ablated volume, as defined during the follow-up imaging, BLV was a better predictor because it excluded the reperfused volume. Thus, BLV may prove to be a powerful tool to disclose adequately treated regions during the planning session of a second treatment, allowing more tailored targeting of the area to ablate and preventing undertreatment.

We are currently assessing how soon after the initial treatment the BLV becomes visible. If further research demonstrates a BLV at the end of the first treatment, then it may be used to identify untreated parts of the fibroid and guide further, more complete treatment without contrast administration.

Table 5 (abstract A99). Comparison of non-perfused volume after first fibroid treatment (NPV1) and black line volume (BLV) observed on follow-up imaging

<table>
<thead>
<tr>
<th>Patient</th>
<th>NPV1 (cc)</th>
<th>Days to Follow Up Imaging</th>
<th>BLV on Follow Up (cc)</th>
<th>Reperfused Volume on Follow Up (cc)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>412</td>
<td>14</td>
<td>419</td>
<td>none</td>
</tr>
<tr>
<td>2</td>
<td>42</td>
<td>43</td>
<td>42</td>
<td>none</td>
</tr>
<tr>
<td>3</td>
<td>314</td>
<td>14</td>
<td>295</td>
<td>16</td>
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<tr>
<td>4</td>
<td>257</td>
<td>7</td>
<td>208</td>
<td>88</td>
</tr>
<tr>
<td>5</td>
<td>85</td>
<td>14</td>
<td>82</td>
<td>5</td>
</tr>
</tbody>
</table>

Fig. 105 (abstract A99). Comparison of (a) NPV1 and (b) NPV2 demonstrates an area of reperfusion (green outline). The BLV contour (c, pink), matches the NPV2 contour (d, pink), and excludes the reperfused contour (green) in Patient 3.
capable of preserving the functionality of myometrial structures, hence may reduce the length of duration before pregnancy. MRRgFUS is a very promising technique for Adenomyosis patients who desired fertility, while its short and long term impacts in the fertility and pregnancy outcomes are still required.

A102
Endoluminal ultrasound applicators for thermal ablation of pancreatic cancer under MR-guidance: Preliminary investigations in an in vivo porcine model
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Objectives
While conventional treatments remain limited, thermal ablation has been demonstrated to provide pain palliation and survival benefit to patients with advanced pancreatic cancer. In light of the anatomical challenges that limit safe and effective thermal ablation to the pancreas, endoluminal ultrasound may serve as a minimally invasive option for treating pancreatic tumors adjacent to the gastrointestinal (GI) tract, and could be compatible with MR-guidance for target confirmation and real-time thermometry. This study’s objective was to perform a preliminary investigation of endoluminal placement and heating performance of prototype ultrasound applicators and temperature monitoring capabilities in MR-guided in vivo porcine pancreas ablations.

Methods
A family of MR-compatible endoluminal applicators were fabricated and characterized using radiation force balance and hydrophone measurements. Each applicator possessed a distinct two-element transducer array (~3.2 MHz) covered by a cooling balloon at the tip of a flexible catheter assembly, as shown in Fig. 106. Three transducer geometries were considered: 10x10 mm (planar), 8x10 mm with 20 mm radius of curvature (ROC) along the short dimension (lightly-focused curvilinear), and 9.3x1.14 mm with 25 mm ROC along the long axis (strongly-focused curvilinear). In each in vivo porcine experiment, an applicator was orally introduced and positioned in the stomach adjacent to the splenic lobe of the pancreas. Sonication were performed through the luminal wall into the pancreas for ablation, at durations ranging from 3-16 minutes and applied intensities of 5-7 W/cm². MR-guidance techniques were developed and evaluated for anatomical target identification, tracking/placement of the applicator, and PRF-based MR temperature imaging (MRTI), implemented in the real-time RTHawk software environment.

Results
Placement of the endoluminal applicators in the stomach adjacent to pancreatic tissue, as demonstrated in Fig. 107, was achieved in all porcine studies (n = 5). In vivo MRTI-guided heating trials demonstrated capability of ~15-20 °C temperature elevation in pancreatic tissue at 1-2 cm depths using the planar and lightly-curvilinear applicators (6-16 mins, 5-7 W/cm²). Strongly-focused curvilinear applicators were capable of reaching higher temperature elevations of ~25-35 °C at 2-3 cm depths and more spatially localized lesions in shorter treatment durations (3-5 mins, ~5 W/cm²), as shown in Fig. 108. Dimensions of thermal lesions in excised pancreas ranged from 9-28 mm, 3-10 mm, and 5-10 mm in length, width, and depth, respectively, as verified through histopathological analysis of tissue sections (Fig. 109). Multiple-baseline reconstruction and respiratory-gated acquisition were effective in suppressing motion artifacts and improving temperature precision for multislice MRTI in the in vivo studies.

Conclusions
This study demonstrates the technical feasibility of generating volumetric ablation in porcine pancreatic tissue using endoluminal ultrasound applicators positioned in the GI tract, with integrated MR-guidance for target identification, device positioning/alignment, and MRTI treatment monitoring. Future applicator development will involve incorporating endoscopic steering capabilities to enhance control over endoluminal placement, fixation, and coupling to the gastric wall to improve positional accuracy and luminal sparing.
A randomized, sham-controlled trial of transcranial MR guided Focused Ultrasound thalamotomy trial for the treatment of tremor-dominant, idiopathic Parkinson’s disease

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Objectives

Traditional stereotactic RF thalamotomy has been used with success in medication refractory tremor-dominant Parkinson’s disease. Recently, transcranial MR guided Focused Ultrasound (MRgFUS) has been used to successfully perform thalamotomy for essential tremor. We designed a double blinded, randomized controlled trial to investigate the effectiveness of MRgFUS thalamotomy in tremor-dominant PD.

Methods

Patients with medication refractory, tremor-dominant Parkinson’s disease were enrolled in the two center study and randomized 1:2 to receive either a sham procedure or treatment. After the 3 month blinded phase, the sham group was offered treatment. Outcome was measured with blinded CRST and UPDRS ratings. The primary outcome compared improvement in hand tremor between the treatment and sham procedure at 3 months. Secondary outcomes were measured with UPDRS and hand tremor at 12 months. Safety was assessed with MRI, adverse events, and comprehensive Neurocognitive assessment.

Results

Twenty-seven patients were enrolled and six were randomized to a sham procedure. For the primary outcome assessment, there was a mean 50% improvement in hand tremor from MRgFUS thalamotomy at 3 months compared to a 22% improvement from the sham procedures (p = 0.088). The 1 yr tremor scores for all 19 patients treated with 1 year follow up data (blinded and unblinded) showed a reduction in tremor scores of 40.6% (p = 0.0154) and a mean reduction in medicated UPDRS motor scores of 3.7 (32%, p = 0.0326). Sham patients had a notable placebo effect with a mean 21.5% improvement in tremor scores at 3 months. Twenty seven patients completed the primary analysis, 19 patients completed the 12 months assessment, 3 patients opted for DBS, 3 were lost to follow up, 1 patient opted for no treatment, and 1 is pending 12 m evaluation.

Conclusions

Transcranial MRgFUS demonstrates a trend towards improvement in hand tremor, and a clinically significant reduction in mean UPDRS. A significant placebo response was noted in the randomized trial.

A104

MR Image-guided delivery of non-viral miRNA-34a gene vectors via Focused Ultrasound inhibits tumor growth in a mouse glioma model

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Objectives

Glioblastoma (GBM) is remarkably difficult to treat with systemically administered therapeutics because the blood brain/blood tumor barrier (BBB/BTB) blocks their diffusion from the bloodstream, while the bioadhesive and nanoporous ECM hinders their tissue dispersion. In addition, GBM is characterized by genetic complexity, molecular adaptability, and multiple deregulated pathways, all of which markedly limit conventional therapies. Here, we address these obstacles by combining Focused Ultrasound (FUS)-mediated disruption of the BBB/BTB with brain-penetrating non-viral gene vectors (DNA-BPN) bearing the plasmid for miRNA-34a, a miRNA that is downregulated in human GBM and predicted to regulate multiple tumor suppressive pathways.
U87 malignant glioma cells were injected into the brains of athymic nude mice. Five days later, anesthetized mice were coupled to an MRI compatible 1.14 MHz FUS system. T1 contrast MR imaging was used to visualize and target tumors. miR-34a plasmid-bearing DNA-BPN that had been densely PEGylated to permit brain penetration were i.v. co-injected with microbubbles. FUS was applied to tumors with a 0.5 % duty cycle for 2 minutes at either 0.4 or 0.5 MPa peak negative pressure (PNP). Immediately after sonication, additional contrast MR images confirmed BBB/BTB opening. Untreated mice and mice that were i.v. injected with miRNA-34a BPNs and MBs without FUS application served as controls. At 7 days and 15 days after treatment, corresponding to 12 and 20 days after tumor implantation, tumor volumes were determined from 7 T MR images.

Results

Figure 110a shows representative pre- and post-FUS T1-weighted contrast MR images when using PNPs of 0.4 and 0.5 MPa. Enhanced signal in the post-FUS images indicates increased leakage of gadolinium contrast agent across the vasculature due to FUS-mediated BBB/BTB disruption. Note that BBB/BTB disruption appears to be further enhanced at 0.5 MPa when compared to 0.4 MPa. Figure 110b shows representative 7 T images of U87 tumors for each group taken at 12 and 20 days post tumor implantation. These images were used to quantify post-treatment tumor volume (Fig. 110c). At Day 20, mice treated with 0.5 MPa FUS + MB + miRNA-34a BPN (n = 3) showed a marked and statistically significant (p < 0.05) decrease in tumor volume when compared to the untreated (n = 5), miRNA-34a BPN + MB (n = 4), and 0.4 MPa FUS + MB + miRNA-34a BPN (n = 4) groups. For all groups, T2* MRI (not shown) confirmed that there was no leakage of blood products across the BBB/BTB.

Conclusions

We show that a single MR image-guided treatment, entailing the FUS (1.14 MHz; 0.5 MPa) mediated delivery of brain-penetrating non-viral miR-34a gene carriers across the BBB/BTB, can significantly inhibit glioma growth in mice. To the best of our knowledge, this represents the first time that therapeutic miRNA plasmid has been complexed into a non-viral gene nanocarrier and delivered using Focused Ultrasound for treatment of gliomas. Ongoing studies are centered on optimizing miRNA-34a expression levels, verifying changes in miRNA-34a oncogenic target protein expression, and assessing tumor cell apoptosis and proliferation.

Fig. 110 (abstract A104). a Pre- and Post-FUS contrast MR images of U87 tumors (yellow arrows). b 7 T contrast MR images of U87 tumors (yellow arrows) at 12 and 20 days after tumor implantation. c U87 tumor volumes. P < 0.05 vs. all other groups at Day 20

Neurorestoration of the nigrostriatal pathway through multiple treatments with FUS-facilitated brain drug delivery

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A105

Objectives

Current Central Nervous System (CNS) drug delivery techniques are confined to either targeted but invasive or to non-targeted and non-invasive. Focused Ultrasound (FUS) coupled with the systemic administration of microbubbles has been proven to open the Blood-Brain Barrier (BBB) locally, transiently and non-invasively, thus facilitating the diffusion of neurotrophic factors. IV injection of Neurturin, a member of the glial derived neurotrophic factors (GDNF) family has been demonstrated to activate the downstream signaling pathway following BBB opening. The promising previous findings set the milestone for investigating the effect of Neurturin on the depleted dopaminergic neurons in vivo. The aim of the current study was thus to investigate the neurorestorative effects of single and triple delivery sessions of the neurotrophic factor Neurturin in a Parkinsonian mouse model.

Methods

For this study a single control group was employed and three groups per treatment cascade: single and triple. Wild type mice (12 months old) were infused with sub-acute dosages of MPTP causing apoptotic degeneration in the nigrostriatal pathway. After stabilization of the MPTP lesions and the decontamination period, the treatment groups were sonicated on the left hemisphere targeting twice the Caudate Putamen region (CPu) and once the Substantia Nigra region (SN) while one received an IV injection of 0.5 mg Neurturin. The survival period after the last treatment was equal to 28 days allowing the neurotrophic factor to develop its restorative effects followed by coronal sectioning for tissue processing. The brain slices of both the SN and the CPu were stained for tyrosine hydroxylase positive cells (TH+) with a custom protocol. Brain sections were imaged to count the TH+ nerve cell bodies on the SN while the dendritic and terminal areas were quantified by a custom MATLAB algorithm by computing the percentage of the relative difference (RD) between the two hemispheres as for each mouse the contralateral side was compared to the ipsilateral side to eliminate across-mice variation in the number of nuclei and projections.

Results

Comparison between the neuronal cells on the contralateral side and the ipsilateral side revealed an increase in the cell number on the treated side only for the group that received Neurturin with FUS. The RD was found to be significantly higher in the SN region for the groups that received Neurturin, both in the cases of single and multiple treatments. An increase in RD for the group that received multiple treatments with Neurturin compared to the single administration group was not observed. On the other hand, significant increase in the RD was only found in the case of the triple-treatment group in the CPu region.

Conclusions

The increased RD at the SN region revealed the immense potential of the current treatment and the necessity of expanding the study into multiple administrations of Neurturin. The comparable findings in dendritic density at the SN region between the two treatment regimen points towards the direction of examining the treatment effect at the CPu region. The significantly increased CPu terminal density could be explained as the restoration of the neuronal processes through collateral sprouting. Finally, despite the increase in neuronal cells on the ipsilateral
side, no statistical significance was observed. This finding is in accord-
ance with our knowledge of Neurturin restoring impaired neurons and
not regenerating them. The presented findings are essential considering
the therapeutic effect of multiple treatments with FUS enhanced drug
delivery in patients.

Fig. 111 (abstract A105). Substantia Nigra: a Atlas 3D representation
of the nigrostriatal pathway and the involved structures, Substantia
Nigra (SN) and Caudateputamen (CPu), b Coronal section at a cross-
section of the SN in increasing magnification. c Fluorescent im-

Fig. 112 (abstract A106). a T1wGd, T2w, T2* MRI (3.0 T), and b
T2*w and DTI MRI (9.4 T) of a representative brain after 6 weekly
sonications

A106
Long-term effects of Blood-Brain Barrier opening with pulsed
Focused Ultrasound and microbubbles
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Joseph Frank
National Institutes of Health Clinical Center, Bethesda, Maryland, USA

Objectives
The objective of this study is to use advanced Magnetic Resonance
Imaging (MRI) techniques to characterize the long-term effects of
pulsed Focused Ultrasound (pFUS) and microbubbles (MB) in the rat
brain. We evaluated the effects of single and repeated Blood-Brain
Barrier (BBB) disruptions (D) on morphology in the rat striatum and
hippocampus as monitored by MRI and histology over 12 weeks.

Methods
Female Sprague Dawley rats (n = 6/group) received either pFUS + MB
once (Group A) or six weekly exposures (Group B) to the striatum and
the contralateral hippocampus along with 100 μl of intravenous MB
(Optison™, GE Healthcare). Prior to the first sonication rats received 3
daily doses of 300 mg/kg 5-Bromo-2′ deoxy-uridine (BrdU, Sigma Al-
drich) intraperitoneally to label proliferating (neurogenesis) cells
in vivo. 0.3–0.5 MPa acoustic pressures were applied in 10 ms burst length and
1 % duty cycle (9 focal points, 120 sec/9 focal points – striatum, 120 sec/
4 focal points – hippocampus) using a single-element spherical FUS
transducer (center frequency 589.636 kHz; FUS Instruments). T2w, T2*w
and Gadofosveset (Gd)-enhanced T1w images were obtained by 3.0 T
MRI (Philips) and T2w, T2*w, diffusion tensor imaging (DTI) was per-
formed by 9.4 T MRI (Bruker). Animals were euthanized 6 or 12 weeks
after the first pFUS treatment. Histological evaluation of brain and track-
ing of BrdU tagged cells was performed and compared to untreated
contralateral brain.

Results
In groups A and B, contrast enhancement on T1w images was de-
lected post sonication in the striatum and the hippocampus. Gd-
extravasation, T2 and T2* abnormalities were not seen in the brain
1 day post pFUS + MB at 9.4 T MRI. Approximately 50 % of Group A
and 100 % of Group B rats had hypointense voxels appeared on T2* 3 T MRI 2-3 weeks post pFUS + MB (Fig. 112a) consistent with
microhemorrhages or influx of metallophagocytic cells from the
spleen within the parenchyma. Fractional anisotropic (FA) changes in
white matter fiber structure- and gray matter abnormalities on DTI
MRI were detected in regions with the T2* abnormalities suggestive of
increased astrogliosis and transient axonal damage along with
increased cell density. Group B also demonstrated ventriculomegaly
and meningeal abnormalities (Fig. 112a, b). Histologically, increase in
numbers of Nissl positive cells along with activated microglia and
BrdU labeled cells were detected in both cortex and hippocampus in
greater amounts in Group B compared to Group A rats, consistent
with multiple episodes of induced sterile inflammatory response (SIR)
within the parenchyma.

Conclusions
We have observed a complex graded molecular and cellular SIR with
increase in the brain up to 24 hrs after pFUS + MB. However, little
has been reported on using advanced imaging techniques at high
magnetic field strengths on the long-term effects of single and re-
peated pFUS + MB in the brain. Based on changes in FA, DTI MRI
demonstrated low degree of structural injury at the location of both
single and multiple sonication exposures. Multiple pFUS + MB expo-
sures resulted in increased ventriculomegaly and numbers of acti-
vated microglia, Nissl positive cells and BrDU in both striatum and
hippocampus. Increased numbers of BrdU+ cells would be consistent
with the stimulation of neurogenesis indicative of increased damage
secondary to repeated SIR episodes in the parenchyma.

A107
Activatable nanodelivery combined with CpG-ODN and anti-PD-1
achieves a complete response in directly-treated and contralateral
tumors in a murine breast cancer model
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Objectives
Activatable nanotherapeutics are attractive since the toxicity of chemo-
therapeutics can be constrained to a small region; combining such a
strategy with immunotherapy is the goal of this study. We have previ-
ously shown that administration of CpG-ODN as an adjuvant, together
with local release of doxorubicin from temperature sensitive liposomes
(TSL) resulted in regression of directly-treated tumors, suppressed growth
of contralateral tumors and reduced chemotherapeutic-mediated toxicity
in a murine breast cancer model.

Methods
The following is the protocol explored for the addition of anti-PD1:
immune intact FVB/n mice with bilateral invasive neu deletion syngen-
ethic transplanted tumors were treated with a combination of anti-PD1
(aPD-1, 200 μg, i.p.) and intratumoral administration of CpG-ODN
(100 μg, i.t.) on days 0, 7, 14 and 0, 3, 7, 10, 17 and 24, respectively.
Doxorubicin TSL were prepared from DPPC:MPPC:DSPE-PEG2k, 86:10:4
in the presence of copper (II) gluconate and triethanolamine at 0.2 mg-
drug/mg-lipid and administrated i.v. at 6 mg doxorubicin/kg body
weight on days 10, 17, 24. The formation of a complex between doxorubicin and copper was created to enhance the circulation and stability of TSL and to reduce systemic toxicity. To trigger drug release, hyperthermia was induced in the primary tumor with ultrasound (peak ultrasound pressure of 1.1 MPa at a frequency of 1.5 MHz) at 42 °C for 5 min prior to and 20 min post drug injection with a variable duty cycle. Immediately afterwards, 100 μg of CpG-ODN 1826 was administered intratumorally to the isononated tumor.

**Results**

Upon treatment with this combination of locally-released doxorubicin, local administration of CpG-ODN and systemic aPD-1, 100 % of treated and contralateral tumors regressed by at least 80 %; further, all of the directly-treated tumors and 50 % of the contralateral tumors were eliminated without recurrence. Thus, a 50 % complete response rate was achieved, with tumor regression observed immediately after the incorporation of the doxorubicin treatment. By contrast, administration of CpG-ODN and systemic aPD-1 alone resulted in regression of 66 % of treated and contralateral tumors.

**Conclusions**

We demonstrate for the first time that blocking of the programmed death-1 (PD-1) pathway in conjunction with immunogenic cell death induced by CpG-ODN and activatable nanodelivery of doxorubicin can generate curative responses in both primary and contralateral tumors. Increases in cytotoxic CD8+ T lymphocytes and a reduction in regulatory T cells and myeloid-derived suppressor cells were observed in both directly treated and contralateral tumors. This combinatorial approach was curative for directly-treated tumors and overall survival was significantly extended, however, the contralateral tumor returned in all treated mice.

**A108**

Towards FUS lung cancer ablation: Aspects of MR guidance in flooded lung

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**Objectives**

One-Lung flooding (OLF) enables HIFU application under sonographic guidance in lung. Additionally MRgFUS is superior for non-invasive ablation under thermal dose monitoring. FUS treatment of lung tumours requires complete gas- liquid exchange in the treated lung section, which we likely affect also imaging modalities. MRI is seldom used for diagnosis and staging of thoracic tumours because of low proton density with short T2* and field inhomogeneities in ventilated lung. So far no data is accessible for MR imaging of lung in flooded condition and therefore its use for FUS guidance on lung isonatum is required. This study was aimed to investigate in vivo OLF in MR environment and to evaluate the capabilities of MR imaging on lung tumours (NSCLC).

**Methods**

OLF was performed on six pigs (female 35-60 kg) in 3 T MR (Prisma, Siemens AG, Germany). After narcosis, mechanical ventilation was performed with an ICU respirator (Servo 900, Siemens AG, Germany) through a 39 Ch double-lumen tube (Mallinkrodt, Ireland). After ventilation with FIO2 = 1.0 for 30 min the left lung wing was flooded with isotonic saline (0,9 %@ 35 °C). Ventilation of the right lung was maintained for 60-90 min, followed by re-ventilation for 30 min. MR imaging was performed with spine and body array coils in lateral position using T2 HASTE, and T1 weighted GRE sequences. As a cancerous ex vivo model, MR imaging was performed on ten human lung lobes containing lung cancer (NSCLC). Lobes where flooded immediately after resection with saline (35 °C) and imaged using GRE and T2 HASTE sequences in 1.5 T MR (Achieva, Philips, NL).

**Results**

OLF was successfully performed in MR environment on all animals (6/6). Flooded lung appears hyperintense in T2 weighted images (Fig. 113) and hypointense in T1w. Bronchial wall and vascular structures in flooded lung clearly appear as hypointense structures in T2 weighted image at high level of detail. GRE phase images show a homogenous phase in bronchial and alveolar tissue of the flooded lung wing. NSCLC tissue appear in flooded lung as strongly hyperintense lesions in T1 (Fig. 114) and hypointense in T2 weighted images with clear demarcation from lung parenchyma.

**Conclusions**

This study shows that in vivo OLF is operable in MR environment, all animals survived the procedure. The absence of susceptibility changes in GRE phase images suggests a complete gas to saline exchange in the flooded lung wing. For FUS guidance a demarcation of tumor to parenchymal structures is essential. In T1w images the lung tumour appears bright hyperintense to surrounding lung in all tumours. Further in T2w images a good demarcation between bronchus and parenchyma can be used for anatomic orientation. It can be concluded that MR imaging based on general fast GRE and TSE sequences is sufficient for FUS guidance on lung cancer during OLF. Further research investigating nodule size detection limits and the use of PRFS thermometry in flooded lung is required.
Towards patient-specific acoustic scattering models for use in transcranial HIFU treatments
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Objectives
In transcranial HIFU treatments, scattering that occurs in the porous skull can contribute to a significant portion of the ultrasonic attenuation. In many acoustic models, scattering is neglected or considered homogenous across the skull. However, significant variations in skull anatomy may require patient-specific models of scattering for treatment planning. These patient-specific models are based on clinical CT scans that do not have high enough resolution to resolve many of the scattering sites. In this work, we relate the scattering properties of the skull, as simulated using a high-resolution microCT scan, to quantitative Hounsfield Units (HU) as measured by a clinical CT.

Methods
A high-resolution (23.6 micrometer isotropic) microCT scan was obtained of an ex vivo human skull section in a holder and thresholded into cortical bone and pores. The same skull was imaged with a clinical-resolution (0.23 x 0.23 x 0.4 mm) CT scan. The images were registered using hand-selected correspondence points. A database of pores was created from the microCT scan via connected components analysis and associated with the corresponding HU values in the clinical CT scan based on the center of mass. Additionally, the ratio of cortical bone versus porous material in the microCT was calculated for each associated HU range in the clinical CT. Models for various HU ranges were constructed by randomly selecting hole shape and sizes associated with the HU range and placing them in uniform bone until the ratio of cortical bone to porous material was the same as measured for the HU range. Simulations were then performed using the Hybrid Angular Spectrum technique assuming no absorption. The ratio of power at the focus compared to the input power at the face of the model is representative of scattering and reflections.

Results
Figure 115 shows example models for 301-400 and 801-900 HU ranges respectively. For simulations, the transducer is to the left of the model with the focus in the water to the right. These representative models clearly show the differences in shape distribution of the pores as well as overall density of the different HU ranges. Figure 116 shows the ratio of power at the focal plane compared to the power input to the simulation as a function of HU range. The power lost is due to both scattering and reflections.

Conclusions
The work presented here is the first step towards patient-specific models of acoustic scattering. The models created represent the differences in scattering and reflections for various HU ranges. Future work will be focused on applying these results in simulations at a clinical resolution to validate the method.

Reference
A110

Study of aberrations at the focus of an ultrasonic beam due to the propagation across different areas of the skull
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Objectives
Over 95% of useful therapeutically molecules cannot penetrate to the brain parenchyma in central nervous system due to the Blood-Brain Barrier. In recent years it has been developed a promising ultrasound assisted technique for noninvasive, targeted and safe BBB disruption for drug delivery. This technique employs a transcranial focused ultrasound beam focalized on targeted BBB areas to disrupt. However, there are many limitations of the technique: the skull shows very high acoustic absorption, the acoustic impedance of the skull bone is highly different than the inner brain and outer media and the shape of the skull produces aberrations in the acoustic field generated by focused acoustic beams that cause unexpected targeted focal areas. The aim of this work is to study the transcranial targeting of whatever part of the brain taking into account normal skull surface incidence, in order to find the best incidence zone which generates less beam aberrations.

Methods
A 500 KHz focused transducer (aperture = 100 mm; radius = 140 mm) was simulated with K-Wave Matlab toolbox, which is based on the k-space pseudospectral method. A 3-D fluid media was considered with CT-based acoustic maps of the skull. Geometrical algorithms were developed to select the position of the transducer normal to the skull from occipital to frontal areas and from temporal to parietal and several physical criteria have been applied to quantify the aberration level by comparing the focal area of the beam when it propagates with and without skull.

Results
Refraction has proven to be one of the most important effects in the propagation through the skull. Figure 117 shows the effect of the skull shape in the focus of the single element transducer simulated, both in a XY plane and in the YZ plane. We provide an evaluation of the aberration of the focus of the beam in each plane following a path from the occipital to the frontal areas. Aberration in each plane are different, and more important in the occipital area, were irregularities of the skull shape affects the propagation due to the refraction phenomenon. Here, also, the existence of muscle adhered to the bone increases the refraction of the beam.

Conclusions
The shift in the location of the focus respect to the propagation of the free ultrasonic beam changes when the beam targets different areas of the skull. It can be also affected by the presence of muscles attached to the bone, and of course, when not normal incidence is considered. The best target areas to minimize aberration in the beam have been selected and characterized.

Fig. 117 (abstract A110). Occipital incidence of the ultrasonic beam. The focus suffers a higher deviation in the XY plane than in the XZ one.
record the transducer and probe locations during pulse-echo acquisition. The stylus was mounted to a 3D translation stage to function as an echo target, and near-infrared reflectors with well-characterized geometry were affixed to the transducer with custom mounting hardware to make a rigid body for registration. We acquired locations of the rigid bodies as the probe was moved in 3-dimensions to maximize the echo signal, providing an estimate of the ultrasound focus position in optical coordinates. Next, multi-modal fiducials (visible via MRI) were attached to the transducer, and their positions measured with the Polaris system. Finally, the transducer and a sonication phantom were placed in the MR scanner. The thermal focus position was measured with MR thermometry, and the fiducial locations in MR coordinates were used to register the focus position to optical coordinates. The difference of MRI-derived thermal focus and the pulse-echo focus was the target registration error (TRE).

Results
MRI (green stars) and optically tracked fiducials (red triangles) were registered to coincident locations (Fig. 118b), yielding a fiducial registration error of 2.8 mm. After registration, the mean TRE between the pulse-echo focus and thermal focus across three phantoms was 5.7 +/- 1.5 mm (Fig. 118c). This error is comparable to published methods, but work is focus and thermal focus across three phantoms was 5.7 +/- 1.5 mm. After registration, the mean TRE between the pulse-echo spherical, 1024-element transducer. For thermal therapy, this design

Optical tracking allows prediction of FUS focal zone location. Future experiments will assess the target registration error by performing optically-guided transcranial ablation of a brain-equivalent phantom and assessing the lesion location in a post-procedural MRI.

A113
The feasibility of MR-guided FUS BBB disruption through an intact human skull with a clinical body transducer: Pre-clinical results in a rat model
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Objectives
MR-guided Focused Ultrasound (MRgFUS) has been used for many non-invasive mechanical and thermal therapies throughout the body. One of the most promising applications in the brain is the use of MRgFUS for safe, local, and transient Blood-Brain Barrier (BBB) disruption. Current transcranial MRgFUS therapy is performed using a hemispherical, 1024-element transducer. For thermal therapy, this design is warranted as a large aperture is required to obtain enough gain for focal heating without overheating the skull. For low duty cycle and low pressure applications like BBB disruption, it may be possible to use smaller-aperture transducer arrays like those found in clinical body MRgFUS systems. The goal of this study is to characterize a clinical body MRgFUS system in regards to ultrasound transmission through a human cranium and in an application to disrupt the BBB in vivo in rats.

Methods
A clean, degassed human cranium was mounted in a bath of degassed water over a 14 cm, 256-element clinical body transducer (Sonalleve V2, Philips). Short, 40 cycle ultrasound bursts at 1 MHz were transmitted through the skull, and acoustic pressures were measured using a hydrophone mounted to a positioning stage (Fig. 119). Three different skull orientations were evaluated at 3-4 different sonication depths, along with evaluating the sensitivity of ultrasound transmission to small transducer translations. Six anesthetized rats were shaved and positioned supine on an acoustic window with their brains at the location of prior pressure calibration. MR imaging was used for targeting. Microbubbles (0.02 mL/kg, Definity) were injected concurrently with sonifications (0.4-0.6 MPa in situ based on hydrophone calibration, 10 ms bursts at 1 Hz pulse repetition frequency for 120 s). Gadolinium was administered (0.2 mL/kg, Magnevist) and follow-up T1-weighted imaging was performed to evaluate BBB disruption. Three brains were further examined with histology to look for damage.

Results
Hydrophone measurements demonstrated a great deal of variation in ultrasound transmission with change in skull orientation, though the change was gradual. Attenuation ranged from ~ 3.7 dB to 9.3 dB (11 % to 43 % transmission). For a single skull orientation, the average variation in transmission was 3 % for a 5 mm transducer translation in-plane. Contrast enhancement in the brain indicated focal BBB disruption (Fig. 120). The error in the disruption location perpendicular to the ultrasound beam was 1.0 +/- 0.5 mm, (Fig. 121). The error in the dorso-ventral direction was 2 +/- 2 mm. The Lilliefors normality test did not reject the null hypothesis that these samples were drawn from a normal distribution (p < 0.05). In each animal, enhancement increased with in situ sonication pressure, though there was a large variation.

Conclusions
Transcranial MRgFUS is feasible with a smaller-aperture transducer for BBB disruption. The accuracy of sonication was 1 mm in plane and 2 mm out of plane without any corrections for the skull. For clinical feasibility and safety, however, the attenuation of the skull must be taken into account so as to apply appropriate pressures to disrupt the BBB without causing damage.
The desire to monitor the entire insonicated volume has led the field towards the development of rapid, 3D methods; however, acquiring fully sampled 3D volumetric data to monitor heating is time consuming, and so fast methods must be developed in order to meet the spatial and temporal requirements for adequate monitoring of thermal therapy. Spiral k-space trajectories have higher data acquisition efficiency than the traditional Cartesian scanning so that they are an attractive way to improve temporal resolution while maintaining spatial resolution in MR thermometry. Furthermore, temporal redundancy can be exploited with real-time reconstruction models such as the Kalman filter to further accelerate the scan without sacrificing the real-time tracking capabilities. The purpose of this study was to implement the 3D spiral thermometry sequence on a real-time platform to demonstrate its effectiveness.

**Methods**

A non-accelerated 3D retraced spiral-in/out (RIO) thermometry sequence was implemented on the RTHawk platform (HeartVista, Inc.) to enable real-time sequence control and monitoring of a FUS insonication. RTHawk interfaced with a GE Discovery MR750T 3 T scanner at the UVA Focused Ultrasound Center, where an Insightec ExAblate 650 focused ultrasound transducer was used to induce focal heating. Three Yorkshire pigs were used for initial sequence evaluation. Non-ablative power settings were used to compare the temperature measured by the 3D sequence to those measured by the standard 2D GRE thermometry sequence provided by Insightec. Maximum temperature, measured by the single hottest pixel, and mean temperature, measured as the average of the 9 pixels centered at the hottest pixel in a single plane, were recorded. After FUS, the animals were moved into a head coil array for high quality post-ablation assessment, which induced T2-, Diffusion, and T2*-weighted imaging using standard sequences.

To explore the acceleration technique, a Kalman filter model was developed given as:

\[
x(k) = x(k-1) + w(k-1) \\
z(k) = U(k)F(k) + v(k)
\]

in which \(x(k)\) is the target image, \(z(k)\) is the measurement from undersampling pattern \(U(k)\) and Fourier transform matrix \(F\). The model can give optimal estimation of \(x(k)\) based on undersampled measurement and previous images.

**Results**

Sequence parameters for 3D RIO thermometry were: \(\text{FA} = 5-10^\circ\), \(\text{TR}/\text{TE} = 22.0/12.8\) ms, readout length = 8 ms, interleaves = 24 over a FOV of 280 mm² for an in-plane resolution of 1.5 mm². 3D phase encodes = 12 with through-plane resolution of 3 mm so that the through-plane FOV was 36 mm. Total acquisition time per volume was 6.3 seconds. Sequence parameters for the 2D thermometry method were: \(\text{TR}/\text{TE} = 27.6/12.8\) ms, pixel bandwidth = 44 Hz/px, matrix size 256 x 128 over a FOV of 280 mm² for a resolution of 1.1 x 2.2 mm², slice width of 3 mm, and update interval 3.5 seconds. Real-time *in vivo* thermometry images using the non-accelerated 3D RIO thermometry sequence are shown in Fig. 122. The hot spot is resolved in volumetric space, albeit with lower SNR with the spiral 3D sequence, due in part to reduced voxel size. The maximum temperatures measured by the 3D sequence correlate well with those measured by the 2D sequence (Fig. 123). The mean temporal standard deviations of the 2D and 3D sequences were 0.8 °C and 1.3 °C, respectively.

**Conclusions:** The efficiency of spiral readouts supports rapid generation of 3D temperature maps. In *vivo*, we have successfully monitored the entire ablation with adequate spatial resolution to qualitatively compare the ablation area with temperature. The temporal resolution of the 3D sequence can be further improved with the application of the Kalman filter model to match that of the standard 2D Cartesian sequence. If this combination performs well *in vivo*, the result will be a self-training 3D real-time thermometry method with excellent temporal resolution and no focal spot shift.
A115
Water diffusivity changes with varying focused ultrasound exposure levels in rat brains
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Objectives
Focused ultrasound (FUS) for neuro-interventions is gaining popularity not only for ablating tissues as in the case of the treatment of essential tremors and Parkinson’s disease, but also to study the effects of neuromodulation, and temporary opening of Blood-Brain Barrier non-invasively. However, the biological effects of such low level exposure on the brain tissue are less understood especially at locations that are removed from the focal spot. In this study, we used in vivo diffusion kurtosis imaging (DKI) to measure brain water diffusion changes in vivo in rats at varying FUS exposure levels at the focal spot and away from the focal spot.

Methods
To systematically understand the rain structural, functional, and biochemical changes from focused ultrasound intervention, rats (n = 5; 300 ~ 400 g) were exposed to various FUS energies. Prior to FUS exposure, rats were imaged using an MR scanner (Bruker, Germany) for small animals. The rats were then subject to various levels of MR guided FUS exposure ranging from 1-3 W using an eight array transducer from IGT, France. Localization for the exposure was determined using ARFI technique. Once the focal spot was identified, each animal was subject to exposure at a specific energy level. Temperature monitoring was performed during the exposure to identify no over exposure or cavitation occurs in other areas of the brain. Following the exposure, high-resolution images similar to the pre-exposure were obtained within two hours after FUS. The axial temperature distribution of the focal spot was visualized with MRI during ultrasound exposure. To acquire the coronal temperature distribution of the focal spot, a k-space time-domain method was employed to simulate the acoustic field during the exposure by solving the full Westervelt equation. Temperature simulation was estimated by solving inhomogeneous Pennes equation of heat conduction.

Results
The T2 weighted and diffusion tensor images after a minute of ultrasound exposure with varying energy are shown in Fig. 124. As an example, recorded temperature with 3 W acoustic power is shown in Fig. 125, in which the maximum temperature rise was found to be 15 °C. This was also confirmed from the simulations using the bio-heat equation which also takes into consideration of the skull and tissue map from CT images. The tissue temperature distribution from the simulation over the exposure time was also found to be about 15 °C (acoustic power is 3 W; base temperature is 37 °C) as well (Fig. 126). With increasing acoustic power, increased mean kurtosis (MK) and reduced mean diffusion (MD) were observed both at the focal spot and away from the focal spot (Fig. 127). Increased MK typically means that there is increased heterogeneity in the tissue microstructure suggesting increased astroglial reactivity associated with the rise in temperature. These results suggest that at even low exposure rates and away from the focal spot, where the pressures reaches ~400-500 kPa, water diffusion changes occur that can be detected by diffusion tensor imaging in the absence of any visible damage observed on T1 or T2-weighted image.

Conclusions
We hypothesize that these water diffusivity changes induced by focused ultrasound are mainly due to shearing forces exerted on the brain tissue that have an effect on the tissue environment if only temporarily. Ultrasound simulations could provide information on the pressure distribution and temperature distribution throughout the brain to enable correlation with water diffusion changes. In the future we will be studying the long term effects of such low level exposure to understand if such changes are similar to those observed on animals with mild traumatic brain injury follow exposure to blast pressures.
A116
Cavitation localization using a modified trilateration method: Proof of concept
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Objectives
Since cavitation is a potential risk factor that must be monitored and limited, the power measurement and frequency response of the acquired signal through passive cavitation detectors (PCDs) are utilized to detect cavitation activity. Currently, the sonication system was designed to automatically shut down when the power of the cavitation signal exceeds a certain threshold. However, cavitation caused by the bubbles on the skin could also contribute to the appearance of large broad band signal resulting in unexpected stop of sonication. Localizing the cavitation source may improve the monitoring of the cavitation activity at the focal area and provide more reliable sonication with known cavitation location.

The purpose of this study is to localize cavitation near the focus using onboard PCD set without change of a current clinical system setup and to try to distinguish if the cavitation is activated in region of interest.

Methods
In order to generate the cavitation intentionally, high power sonication was targeted on the geometrical center in tap water with and without a human skull. Eight PCDs on a 220 kHz hemispherical transducer (ExAblate Neuro system, InSightec) were synchronized with the pulse sonication. The PCD signals were recorded each sonication and saved into a log file. One of the pulse signal (4096 sample points with 2 MHz sampling rate) acquired by eight the PCDs are shown in Fig. 128. The relatively large signals are from cavitation and we assume the first peak (marked with red arrow in Fig. 129) of this large signal train as initiation of the cavitation activity. The mean value of the first 200 samples and a constant value was used as a threshold to track the first arrived signal. A trilateration, which is normally utilized as a standard localization method on global positioning system (GPS), was modified to localize the cavitation spot. The cavitation signals from 4 PCDs were read in MATLAB software to implement the cavitation localization. The distance between tracked location and sonication location was calculated to verify the localization error.

Results
We found time shifting of the cavitated signals through a skull compared to the data without skull. As shown in Fig. 130, the red arrow denotes the arriving time on tap water data (without skull) and green arrow shows the shifted signals through the skull. And the strong signals pointed by yellow arrow is the reflected signals from skull. The simulation signals using a k-wave model result in cavitation localization with an initial errors of 3 ± 3 mm. Implementation of this localization algorithm to experimental data made the cavitation localization with larger errors of 49 ± 21 mm in tap water (without skull) compared to the water with skull (20 ± 4 mm shown in Fig. 131). It may be because of the signal patterns of the skull embedded data containing aforementioned discernible-vertical line (marked with white arrow in Fig. 130) and has higher SNR near the arriving time.

Conclusions
In conclusion, the cavitation localization could be possible with an error range of 20 ± 4 mm with skull with some limitations. So far, we verified the localization algorithm based on the experimental data targeted at the geometrical center only, and more off-centered data are required to confirm the robustness of this localization algorithm.
Objectives
Transcranial application of low-intensity focused ultrasound (FUS) for brain stimulation has proven to be a powerful tool to manipulate brain activity with a few millimeters of spatial resolution while keeping it completely non-invasive (Yoo et al., 2011). Until recently, however, transcranial FUS-mediated brain stimulation conceded some room for variability and inconsistency in exerting its neuromodulatory effects; it is unclear how different types of anesthetic drugs and varying anesthetic depth intervene in the process of ultrasonic neuromodulation. For these reasons, it is necessary to discover the effective time window for neuromodulation. The purpose of the present study is to monitor the anesthetic depth with electroencephalogram (EEG) to optimize the working parameters for neuromodulation in different anesthetic periods.

Methods
Sprague-Dawley rats (female, n = 8) were induced to anesthesia with 5 % of isoflurane, and sustained anesthesia by intraperitoneal injection of 0.1 ml/100 g ketamine/xylazine cocktail. Power spectra of two EEG channels were obtained from prefrontal cortex (Fp1, Fp2), and one ground electrode were placed at midline of the cerebellum (Fig. 132a). We adopted 350 kHz of fundamental frequency, 50 % of duty cycle, 1 kHz of pulse-repetition frequency, and 300 ms of sonication duration, and acoustic intensity of 3.3 W/cm2 Ispta (spatial-peak temporal-average intensity) as the sonication parameters for inducing the tail movement (Kim et al., 2014). EEG was recorded approximately for an hour until the animal wakes up, and each experimental recording was divided into 6 consecutive periods to calculate the depth of anesthesia. From EEG spectrum analysis, SEF 95 % (spectral edge frequency), MPF (median power frequency), and vigilance index, and other spectral parameters were used to estimate the depth of anesthesia.

Results
At the 3rd period, EEG delta and alpha activity were strongly suppressed while gamma activity was significantly enhanced during the time (Fig. 132b), which indicate rapid eyeball movement (REM) sleep like EEG activity (Ferri et al., 2001). At this stage of anesthesia, even at the intensity level above the previously confirmed excitation threshold did not elicit any tail movement as stimulating the motor cortex. This may be translated as the results of REM stage characteristics where the external stimuli are strongly inhibited. During the 5th and 6th periods, SEF 95 % and MPF repeated increasing tendency (Fig. 132c) from briefly deepened 4th anesthetic period after 3rd period (REM stage) implying lightened anesthetic depth (Kortelainen et al., 2012).

Conclusions
We observed voluntary whisker movements at 5th and 6th period which is a typical sign for superficial depth of anesthesia (Brecht et al., 2004), and the ultrasound induced tail movement success rates accompanying the whisker movements were higher at both periods than the rest of the anesthetic stages.
Objectives
Skull bone is heterogeneous and distorts ultrasound waves. This alters the location, shape, and intensity of the beam's focus. To compensate for skull distortions, phase corrections may be applied to each element - these corrections are estimated using simulations of the patient skull.

Traditionally, models of the patient skull are derived from high-resolution computed tomography (CT) images. However, there is a move towards ultra-short echo time (UTE) magnetic resonance imaging to reduce patient exposure to ionizing radiation.

UTE cannot achieve the same resolution as CT, therefore it is unclear whether simulations based on UTE-derived skull models will yield accurate phase corrections. To determine the feasibility of the lower resolution UTE-based simulations, we investigate the trade-offs between skull model resolution and characteristics of the corrected beam. In this study, we use CT images to create skull models of varying resolution.

Methods
We performed 3D simulations using the hybrid angular spectrum (HAS) method at 680 kHz. Skull models were generated by down-sampling high-resolution skull CT images with cubic convolution interpolation. Isotropic resolutions of 0.5 mm to 3 mm (in 0.5 mm steps) were used. To compensate for phase distortion introduced by the skull, the phase of each transducer element was estimated at the intended location. We applied the conjugate of this phase to achieve a phase correction. A different set of phases was calculated at each resolution, and the resulting beam was simulated using the full resolution CT model.

We define a global efficiency metric as the ratio of focal spot intensity to total intensity in the brain; this metric detects the presence of large sidelobes or multiple focal spots.

Results
With decreasing resolution, full width at half maximum and sidelobe level increase whereas focal intensity and global efficiency decrease (Fig. 133). There is minimal blurring of the focal spot for resolutions 0.5 mm to 1.5 mm. However, the focal spot quickly degrades as the voxel size exceeds 1.5 mm (Fig. 134).

Conclusions
Focal intensity, sidelobe level, and global efficiency worsen at resolutions 2 mm and below. Even though computation time is drastically shorter, the deterioration of the focal spot is unacceptable.

To avoid frequency aliasing in the HAS method, the minimum resolution should be 1.1 mm (assuming 680 kHz frequency and 1500 m/s speed of sound). The 1.5 mm and 2 mm resolutions will thus lead to aliasing. The 1.5 mm option, however, appears to yield similar results to the higher resolution options. This may be plausible considering that the acoustic wavelength in bone is approximately 3.4 mm (assuming 2300 m/s speed of sound). 1.5 mm would satisfy the resolution requirement in the skull, which is the dominant contributor to phase aberrations.

Therefore, a resolution cutoff is suggested at around 1.5 mm, with additional work recommended to determine the minimum effective resolution.

Objective
This study investigated the impact of oxygen versus medical air as a carrier gas for isoflurane anesthesia on Blood-Brain Barrier (BBB) disruption, a method under investigation that can enable and target the delivery of drugs to the brain.

Methods
In experiments in mice (N = 7), four brain targets were sonicated in combination with administration of Optison microbubbles (100 μl/kg) under isoflurane anesthesia. A 690 kHz focused ultrasound transducer applied 10 ms bursts at 1 Hz for two minutes at applied peak pressure amplitudes of 0.51-0.54 MPa (N = 2) or 0.34-0.36 MPa (N = 5). After sonication of two locations in one hemisphere, the carrier gas for the anesthesia was changed from oxygen to medical air, or vice versa, and the sonications were repeated in the contralateral hemisphere. Acoustic emissions were recorded during each sonication, and the resulting BBB disruption was assessed using contrast-enhanced MRI.

Results
Example findings are shown in Fig. 135. BBB disruption, as measured by MRI signal enhancement after Gd-DTPA administration, was significantly greater (P < 0.001) in animals breathing medical air as compared with oxygen (Fig. 136b). MRI signal enhancement was detected in 13/14 locations sonicated in animals breathing medical air, but only in 7/14 locations with oxygen. Harmonic emissions were also greater with medical air (P < 0.001), and the decay rate of the harmonic emissions was 1.5 times higher with oxygen (Fig. 136b). A good correlation (R²: 0.46) was observed between MRI signal enhancement after Gd-DTPA administration and harmonic emissions (Fig. 136c). At 0.51-0.54 MPa, both the occurrence and strength of wideband emissions with greater with medical air. However, at 0.34-0.36 MPa, the strength and probability for such emissions was higher with oxygen. Little or no effects were observed in histology at 0.34-0.36 MPa.

Conclusions
The use of oxygen as a carrier gas has a profound impact on BBB disruption. These findings are consistent with earlier work comparing ketamine/xylazine versus isoflurane and oxygen anesthesia on BBB disruption as well as imaging studies showing increased clear- ance rates of ultrasound contrast agents in animals breathing oxygen. The increase in wideband emissions at the lower exposure levels with oxygen without significant BBTthis work was supported by NIH grants R01EB003268, P01CA174645B disruption is an interesting finding that should be pursued further. Overall these findings are important for comparing results from different research groups and should be taken into account in translation to clinical trials.
Using ray tracing to predict histotripsy focal shift in a transcranial setting
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Journal of Therapeutic Ultrasound 2016, 4(Suppl 1):A120

Objectives
Previous experiments with histotripsy has shown that it can be used to generate targeted lesions through the skullcap without aberration correction. However, before therapy is applied, one would like to ensure that the focal point of the transducer through the skullcap is within the confines of the treatment volume. The results from this study demonstrate how ray tracing can be used to predict the location of the focal point in a transcranial setting.

Methods
Histotripsy pulses for the experiments were delivered through human skullcaps using a 256-element 500 kHz hemispherical phased array transducer. Three excised human skullcaps were used for the experiments. During experiments, measurements of pressure profiles and focal shifts of the transducer through the skull were acquired using a needle hydrophone. The pressure profiles were measured by scanning the hydrophone along 2 cm long lines across the axial and transverse directions with respect to the origin of array. The experiments were performed without aberration correction. 3D surface scans of the excised skullcaps were acquired for the simulations. During simulations, straight line rays from each transducer element of the array were propagated through the skull. Calculations of ray trajectories based on Snell’s law, the time of flight of the rays, and losses due to attenuation as the rays propagated through the skull were made during simulations. The rays were then simulated as plane waves that propagated across the origin of the array to calculate the pressure field about origin. The results from the simulations were then compared to the results from the experiments.

Results
Experimental results revealed that the focal point of the transducer remained confined to less than 1 mm of the geometric origin of the array. The experiments also revealed that the propagation direction of acoustic pulses from each element of the transducer were not significantly altered when traveling through the skull. The simulations using ray tracing were able to accurately predict the overall shape of the pressure profile measured experimentally and the focal position of transducer through the skullcap.

Conclusions
The focal location of the transducer was not significantly affected by the skull even when the skullcap was moved within the array. Ray tracing was able to accurately predict the major features of the pressure profile and focal locations through the skull.

Bilateral anterior capsulotomy with MR-guided Focused Ultrasound for refractory obsessive-compulsive disorder: 1 year follow-up results
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Objectives
Obsessive-compulsive disorder (OCD) is a chronic and debilitating psychiatric condition. Traditionally, anterior capsulotomy was an established procedure for treatment of patients with refractory OCD. However, despite the potential benefits of these surgical procedures, patients show significant surgery-related complications. Previously, we demonstrated that bilateral thermal capsulotomy using magnetic
resonance-guided focused ultrasound (MRgFUS), a non-invasive technique with surgical precision for thermal ablation, is a safe and effective therapy for patients with refractory OCD. In the present study, we investigated the long-term outcome of the patients who underwent MRgFUS for refractory OCD.

**Methods**

Eleven medically intractable OCD patients who underwent MRgFUS bilateral thermal anterior capsulotomy from February 2013 to June 2014 were included in this study. Pre- and post-operative (1 week, 1 month, 3 months, 6 months, 12 months) Yale-Brown Obsessive-Compulsive Scale (Y-BOCS), Hamilton Depression Scale (HAMD) and Hamilton Anxiety Scale (HAMA) scores were recorded. Brain MRI was performed pre-operatively, 1 week, 1 month, 3 months and 6 months after the operation to assess the effect of the operation.

**Results**

There was a 36 % improvement in Y-BOCS at 12 months after MRgFUS. The HAMD and the HAMA were showed 54 % and 62 % improvement compared to pre-operative score. The improvement according to time had shown statistical significance (all p value was less than 0.001, repeated ANOVA test). There was a marked difference in MRI findings before and 6 months after the operation. Moreover, there were no neurological, psychological complications with the exception of mild headache and nausea.

**Conclusions**

The optimal treatment for refractory OCD has been unclear; however, in this study, MRgFUS was an effective and safe treatment for refractory OCD. Thus, MRgFUS may be an effective treatment modality for refractory OCD in high-risk surgery patients.

**A122**

**Prediction of shear-mode conversion for transcranial ultrasound using a viscoelastic model for sound propagation: First experimental results**

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**Objectives**

The skull barrier represents the main limitation for many therapeutic applications of High Intensity Focused Ultrasound (HIFU) in the brain tissue. A better understanding and modelling of this barrier facilitates an optimal design of transcranial HIFU applications. An important aspect not yet fully explored in transcranial ultrasound is the use of shear-mode conversion, which in principle can result in less defocusing effects caused by the skull barrier. However, there is not yet a well-established modeling method that can take into account both longitudinal and shear waves in the skull bone. In this study, we present our first results on the validation of a sound transmission model based on the viscoelastic wave propagation adapted for the case of transcranial ultrasound. We compared the predictions obtained with this model with measurements in human calvaria where shear mode conversion was present in the transcranial sound transmission.

**Methods**

Characterization of transcranial acoustic transmission. Human skull caps were immersed in degassed water and put under vacuum at –0.7 MPa for 8 h prior to measurements. A transducer (diameter of 5 cm and a focal length of 10 cm) was used to send 15 ultrasound pulses with a frequency of 270 kHz. A stereoscopic fixation system was used to ensure an incident angle of ultrasound of 40°. The skull was located 12 cm from the transducer. Measurements were performed in the presence of the skull and in water-only conditions using a 2 mm-diameter hydrophone over a scan plan of 35 mm x 35 mm with a spatial step of half-wavelength and located 15 cm from the transducer.

Viscoelastic wave propagation modeling for transcranial ultrasound transmission. The propagation model is based on a staggered grid scheme (Geophysics, 1986; 51(4): 889-90). We included refinements to the original model to consider transmission between fluids and solids (Computers & Geosciences, 2002; 28(8): 887-99) and attenuation effects (Geophysics, 1995; 60(1): 176-184). The equations were solved using a Finite Difference Time Difference (FDTD) solution using approximations of second order in time and fourth order in space.

**Results**

Figure 137 shows a scheme for the experimental setup and the relationships in function of the skull density of the longitudinal and shear speed of sound, and longitudinal and shear attenuation, which were used for this feasibility study. The simulated delay DS on the scan plane between water-only and skull measurements was compared to the corresponding experimental delay DE. Negative values of delay indicated that signals in the presence of the skull arrived before their respective water-only counterparts. Comparison was done in ten measurement locations from three different skull samples.

Figures 138 and 139 shows an example of experimental and simulated results where shear-mode conversion was clearly observed. Each figure shows the delay and normalized pressure maps. Also, plots of the acoustic signals at the center of scan plane are shown in water conditions, in the presence of the skull and in the presence of the skull plus the calculated delay. All measurements combined, seven of ten measurements showed shear-mode conversion (negative delay) while three others (all from skull #3) still showed that longitudinal transmission dominated (positive delay). The linear regression analysis (Fig. 140) of DS versus DE indicated that viscoelastic model predicted accurately the delay for all cases with R2 = 0.97 and a significance of p < 1e-5.

**Conclusions**

The proposed model represents a significant progress for a more accurate prediction of transcranial ultrasound transmission. The comparison with experimental results indicate that the model is adequate to predict sound propagation when either shear or longitudinal transmission dominates. In our tests, two of three skull samples clearly showed shear-mode conversion while the other did not. Closer inspection of skull #3 indicated a less differentiated cortical bone layer and a density 20 % lower compared to the other two specimens. As a result, the average longitudinal speed of sound of skull #3 was slower compared to the other two specimens, implying that its critical angle for shear-mode conversion should be higher than 40°. These results indicated how important can be the degree of differentiation among different subjects. An adequate modelling that considers the skull bone as a viscoelastic medium will be an important asset to better design and execute transcranial HIFU therapy.
A123
Comprehensive dissection of ultrasonic neuromodulation biophysics predicts cell-type-selective stimulation
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Objectives
US can noninvasively suppress or excite central nervous system (CNS) activity using different combinations of stimulation parameters. While applications are already emerging, the underlying biophysics remains unclear regarding the relative contribution of possible mechanisms: extracellular bubble cavitation, thermal effects, acoustic radiation pressure and US-induced intramembrane cavitation within the bilayer membrane (the bilayer sonophore or BLS model). Interestingly, temperature transients, radiation pressure and intramembrane cavitation can induce plasma membrane capacitance changes. Here, we use detailed predictive modelling and find that only intramembrane cavitation can explain all the observed aspects of ultrasonic neuromodulation.

Methods
We analyzed the relevant experimental literature using modified Rayleigh–Plesset intramembrane cavitation BLS biomechanics and acoustic radiation pressure gradients (RPG) - induced membrane dynamics. By coupling these biomechanical models to biophysical membrane models we predict dynamical biophysical responses of artificial bilayer membranes, and of three common neocortical single cell Hodgkin-Huxley type models: i) Regular Spiking (RS) cortical pyramidal neuron, ii) Fast Spiking (FS) cortical inhibitory neuron and iii) Low Threshold Spiking (LTS) cortical inhibitory neuron. Rs-FS-LTS Hodgkin- Huxley based network model and CNS axon model. In addition, live brain tissue RPG subjected areal strains were evaluated in a viscoelastic brain model. To explore high intensity US-related thermal effects on membranes’ biophysics a Gouy-Chapman-Stern-based model was developed.

Results
Only the Neuronal Intramembrane Cavitation Excitation (NICE) models were able to explain US-induced action potential generation through BLS-type pulsating nano-bubbles inside the bilayer plasma membrane: the leaflets’ periodic vibrations induce US-frequency membrane capacitance and potential oscillations, leading to slow charge accumulation across the membrane (on a time scale of tens of milliseconds), until action potentials are generated. In contrast, the analysis of RPG-induced membrane capacitance variations associated with membrane area changes explain artificial membrane results, but were found to be highly unlikely sources for neural excitation, when considering the areal strains expected to form in brain tissue during normal sonication. Further, the NICE-LTS inhibitory neurons show a much higher relative sensitivity to sparse ultrasonic stimulation compared to the other neurons, resulting from their T-type voltage gated calcium channels. This model-based prediction was found to explain the results of a significant body of suppression and excitation experimental studies, including in humans. Finally, a rigorous analysis of thermal effects using Gouy-Chapman-Stern theory reveals that an elevation in membrane capacitance coupled with surface charge-related thermal processes can be associated with high intensity US’s temperature transients.

Conclusions
This study provides a unified theoretical framework for a large body of experiments in multiple preparations across the field of US neuromodulation, lending further support to the hypothesis that intramembrane cavitation is responsible for ultrasonic neuromodulation. They could thus pave the way towards new CNS therapeutic protocols, using the only method that currently allows targeted non-invasive neuromodulation with millimeter spatial resolution essentially anywhere in the brain.

A124
Proportional integral temperature control during Focused Ultrasound treatment with clinical systems in the brain
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Objectives
US can noninvasively suppress or excite central nervous system (CNS) activity using different combinations of stimulation parameters. While applications are already emerging, the underlying biophysics remains unclear regarding the relative contribution of possible mechanisms: extracellular bubble cavitation, thermal effects, acoustic radiation pressure and US-induced intramembrane cavitation within the bilayer membrane (the bilayer sonophore or BLS model). Interestingly, temperature transients, radiation pressure and intramembrane cavitation can induce plasma membrane capacitance changes. Here, we use detailed predictive modelling and find that only intramembrane cavitation can explain all the observed aspects of ultrasonic neuromodulation.

Methods
We analyzed the relevant experimental literature using modified Rayleigh–Plesset intramembrane cavitation BLS biomechanics and acoustic radiation pressure gradients (RPG) - induced membrane dynamics. By coupling these biomechanical models to biophysical membrane models we predict dynamical biophysical responses of artificial bilayer membranes, and of three common neocortical single cell Hodgkin-Huxley type models: i) Regular Spiking (RS) cortical pyramidal neuron, ii) Fast Spiking (FS) cortical inhibitory neuron and iii) Low Threshold Spiking (LTS) cortical inhibitory neuron. Rs-FS-LTS Hodgkin- Huxley based network model and CNS axon model. In addition, live brain tissue RPG subjected areal strains were evaluated in a viscoelastic brain model. To explore high intensity US-related thermal effects on membranes’ biophysics a Gouy-Chapman-Stern-based model was developed.

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Conclusions
This study provides a unified theoretical framework for a large body of experiments in multiple preparations across the field of US neuromodulation, lending further support to the hypothesis that intramembrane cavitation is responsible for ultrasonic neuromodulation. They could thus pave the way towards new CNS therapeutic protocols, using the only method that currently allows targeted non-invasive neuromodulation with millimeter spatial resolution essentially anywhere in the brain.
Objective

Current clinical systems for Focused Ultrasound (FUS) brain surgeries are not designed to precisely or accurately deliver a desired thermal dose to a target in brain tissue. This study attempts to modulate the power applied during sonications to maintain a target temperature in order to accumulate thermal dose in a predictable and controllable fashion, allowing researchers and clinicians to prescribe a delivered thermal dose.

Methods

The regular gating and control of an MRI-guided FUS brain system (ExAblate 4000 Neuro 650 kHz system, InSightec) was bypassed and fed into a proportional-integral-derivative (PID) control loop, and empirically tuned during sonications of gel phantoms with a proportional gain of 5 and an integral gain of 1.5. The system was then used for an animal trial with pigs in vivo, performing thalamus ablation following a craniectomy.

Results

Trials in gel phantoms demonstrated the effectiveness of PID temperature control with the brain system, with different settings for proportional and integral gain affecting the time to reach the target temperature and the magnitude of the temperature overshoot. When used in vivo, the system was able to deliver a desired thermal dose within 10 CEM, with lower temperatures demonstrating a lower rate of accumulation of dose.

Conclusions

In conclusion, PID temperature control is possible to implement in current FUS brain systems, and can be used to deliver a desired thermal dose with the principle limitations of uncertainty in starting temperature and the low signal to noise ratio of the MRI-thermometry.

A125

Evaluation of nonlinear effects in the acoustic field of a hemispherical focusing transducer for ultrasound applications in brain

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Objectives

Ultrasound treatment of the brain is typically performed using large-aperture submegahertz-range sources of a hemispherical shape to achieve better energy concentration. Heat deposition at the focus depends on the frequency content of the ultrasound wave that can be different from the source single-frequency signal due to nonlinear generation of harmonics on the way to the focus. Several theoretical models exist for simulating generation of higher harmonics for weakly or moderately focused waves. In this case the acoustic field can be considered as a directed beam. In the case of the hemispherical sources this assumption is no longer valid, so different strategies are needed.

Methods

Theoretical methods are proposed for evaluating the amplitude level of the second harmonic generated in hemispherically focused fields. Two approaches are used: 1) a simplified nonlinear 1D model; and 2) a more accurate model that accounts for the combined effect of diffraction and nonlinearity. In the first approach, the nonlinear propagation is studied by considering a spherically converging nonlinear wave for which an exact solution of the generalized Burgers equation can be obtained [R.V. Khokhlov et al., Acustica 14 (1964)]. The solution is used to predict the degree of nonlinear effects when the wave arrives to the focal region. In the second approach, the calculation consists of several steps and employs the successive approximations method. First, the linear acoustic problem is solved for a hemispherical source using a spherical harmonics series expansion [O.A. Sapozhnikov and T.V. Sinilu, Acoust. Phys. 48 (2002)]. Then the volume sources for the second harmonic generation are calculated based on the solution to the linear problem. Finally, these sources are used for obtaining an analytic solution for the second harmonic amplitude at the focus.

A126

Research platform for rodent studies of wavefront engineered ultrasonic neuromodulation

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Objectives

Recent years have seen intensifying scientific and clinical interest in US neuromodulation of CNS circuits, and relevant empirical data is rapidly accumulating. To date, studies in this field, and in particular rodent studies, employed single-element transducers. The introduction of wavefront engineering techniques based on phased ultrasonic arrays could open up new vistas for this field, including the generation of multi-focal patterned stimulation.

Methods

We adapted an Insightec prostate phased array (986 phase elements over 2.5x4 cm², 2.3 MHz center frequency) to interface with several custom holders for either in vitro and in vivo work. For in vitro recording we coupled the transducer to a 256-channel planar multi-electrode array (Multichannel systems) and recording system (Ripple Technologies). For in vivo work we designed custom couplers and calibration procedures for operating the system over head-fixed rodents, allowing high-accuracy targeting of the sub-mm focal spot using either marker-based or MR-guidance in a 9.4 T small-animal MRI (Bruker). For generating accurate distributed acoustic phase holograms we applied our recent mathematical framework, based on adaptations from optical computer-generated holography.

Results

The system generates steerable foci whose dimensions are 0.6 x 0.8 x 3 mm (FWHM) in an open water bath, and 0.7 x 0.9 x 4.5 mm through the top portion of a mouse skull, with a sub-mm targeting accuracy. Excitation foci and holograms can be flexibly generated and switched every 100 ms. We observed and characterized stimulus-locked neuronal responses in extended-duration US pulses to different neural systems. In isolated mouse retinas, we observed highly structured off-responses from retinal ganglion cell single-units. In live rats and mice we observed motor responses of the animal’s limbs and distributed BOLD responses to cortical stimulation.

Conclusions

Our early results and the system’s versatile ability to steer and pattern acoustic fields on a sub-mm scale may open new vistas in the study of causal US neuromodulation in relevant small animal models. These abilities can possibly help advance multiple aspects of US neuromodulation, including target optimization and dissecting the underlying biophysical mechanisms.
Efficacy and treatment envelope of transcranial histotripsy therapy without using aberration correction

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Objectives
Histotripsy is an ultrasound therapy that generates cavitation bubble clouds to fractionate soft tissue using short duration, high-amplitude ultrasound pulses. Previous studies have shown that histotripsy is capable of producing lesions through the skullcap using aberration correction. The goals of this study are to demonstrate the ability of histotripsy to generate targeted, precise single lesions through the skull without using aberration correction and to show that larger lesions of arbitrary shapes and sizes can be generated through the skull using either mechanical or phased array steering without aberration correction. We also plan to establish the treatment location envelope, e.g., proximity to the skull surface, over which lesion generation may be accomplished using histotripsy through the skull without aberration correction.

Methods
Histotripsy therapy was delivered using a 500 kHz, 256-element hemispherical phased array transducer with an aperture diameter of 30 cm and a focal distance of 15 cm fabricated in our lab. This transducer is capable of producing estimated peak rarefractional pressures in the free field in excess of 200 MPa with pulse durations of <2 acoustic cycles. For the single lesion generation study, targeted single lesions were generated in red-blood cell (RBC) tissue phantoms through three excised human skulls using histotripsy pulses delivered at a PRF of 1 Hz, with estimated peak rarefractional pressures in the focal zone through the skull of 31-38 MPa. Lesions were examined by optical imaging during treatment. During the treatment envelope and large lesion studies, histotripsy pulses were delivered at a PRF of 200 Hz using a combination of electrical focal steering and mechanical steering to target volumes with characteristic dimensions up to 50 mm, extending up to the skull's interior surface, at locations near the front, back, top, side, and base of the skull. During these experiments, the skull's volume was filled with RBC phantom, and lesion generation was evaluated following treatment via MR imaging. No aberration correction was used.

Results
Histotripsy was able to generate precise, single lesions with radii ≤1 mm in the RBC phantoms without using aberration correction through all skull samples tested. Lesion generation was consistently observed at the geometric focus of the transducer even as the position of the skullcap with respect to the transducer was varied. During large lesion studies, lesions with characteristic dimensions up to 5 cm were generated. Treatment envelope studies revealed that lesion generation was consistently possible at depths of 5-10 mm from the interior skull surface at all locations targeted.

Conclusions
These results show that histotripsy may be used to generate targeted, arbitrarily shaped lesions through intact human skulls without using aberration correction. They demonstrate that both mechanical and phased array steering strategies may be used for lesion generation, and that the treatment envelope for lesion generations extends to within as little as 5 mm from the skull's interior surface. Such capability has the potential to greatly simplify and expand the target window of transcranial ultrasound therapies for non-invasive transcranial applications.

Non-invasive targeted delivery of GABA for neuromodulation: evaluation by resting state functional MRI

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Objectives
We aim to develop novel methodology to non-invasively deliver neurotransmitter chemicals to spatially localized brain regions for modulation of neuronal activity. In this approach, focused ultrasound in conjunction with microbubble injection is used to transiently open the Blood-Brain Barrier (BBB) in a localized region of the rat brain. Neurotransmitter chemicals, such as GABA or glutamate, are then systemically injected such that they leak out and have an effect on neuronal activity only at the site of BBB disruption. We have previously used electrophysiology measurements to demonstrate functional blockade via BBB disruption and GABA administration. The current study will use resting state functional MRI (rs-fMRI) to evaluate the effects of GABA delivery to the rodent thalamus on resting state networks.

Methods
5 - 10 Sprague-Dawley rats will be imaged under resting state conditions using a 2x2 factorial design with conditions of BBB Open vs BBB Closed and GABA Injected vs No GABA Injected. The case of BBB Closed and No GABA Injected will be considered the baseline state. The other three cases will be compared against this baseline for changes in activation patterns.

BBB opening: Right thalamus targeted. Microbubbles injected (Optison, 200 μg/kg), 690 kHz transcranial FUS with 10 ms bursts applied at 1 Hz for 60 seconds.

GABA delivery: Systemic tail vein injection of 20 mg/kg bolus followed by 5 mg/kg/min infusion.

rs-fMRI: Images acquired on a Bruker 7 T scanner with a single shot EPI sequence (TR = 1.5 s, TE = 18 ms, 18 slices, 280 images). Continuous infusion of Dexametomidine used for anesthesia to ensure light anesthetic state. rs-fMRI temporal correlation analysis performed as described by Pawela et al (MRM 2008).

Results
This is a prospective study to be carried out by Jessica Cahill during her summer internship at the Focused Ultrasound Laboratory. We hypothesize the following: 1) that we will see robust resting state network activation in the baseline case; 2) that the resting state networks seen for the cases of BBB Open and No GABA Injected, and BBB Closed and No GABA Injected will show little to no differences relative to the baseline; 3) that the resting state networks seen for the case of BBB Open and GABA Injected will be significantly different than the baseline case. We will not speculate on how these differences will manifest.

Conclusions
Resting state fMRI consistently shows that BOLD signal fluctuations are temporally correlated in brain regions that make up functional networks, such as the visual or sensorimotor systems. Demonstrating that non-invasive targeted delivery of GABA via FUS-temediated BBB opening can alter the activation patterns seen in these networks will be convincing evidence that the approach can modulate brain function.

Acoustic characterization of two pediatric skull phantoms

Elodie Constanciel Colas1, Adrian Wydra1, Adam Waspe1, Thomas Lool1, Roman Maeve2, Samuel Pichardo4, James Drake1
1Centre for Image Guided Innovation and Therapeutic Intervention, Toronto, Ontario, Canada; 2The Institute for Diagnostic Imaging Research, Windsor, Ontario, Canada; 3Hospital for Sick Children, Toronto, Ontario, Canada; 4Thunder Bay Regional Research Institute, Thunder Bay, Ontario, Canada

Objectives
Clinical trials for neurological disorders are ongoing using MR guided Focused Ultrasound (MRgFUS) on adult patients. Pediatric patients have a thinner skull and could highly benefit from this new technique. However, the acoustic properties of their skull have not been reported extensively in the literature. Patient growth affects the structure and shape of the skull. Designing an MRgFUS system dedicated to pediatric patients requires a full understanding of the evolution of the skull with the age of the patient. Acoustic characterization of ex vivo pediatric skulls is ongoing but specimens are rare and cannot represent the entire population. Creating a library of skull phantoms based on CT or...
MRI scans could offset the lack of ex vivo specimens. The aim of this study was to compare two pediatric skull phantoms to their corresponding ex vivo skulls to validate the material properties and the manufacturing process.

Methods
A neonate ex vivo skull and an 8-year-old (8-y.o.) ex vivo skull were scanned using a micro-CT scanner and a clinical CT scanner, respectively. Skull phantoms were manufactured, based on these CT datasets, using a bone-mimicking material previously described (Wydra and Maev, 2013), and in turn CT scanned. Variations in the composition of the material were used to mimic purely cortical bone but also mixe cortical and trabecular bone structures. The 2 skulls and their phantoms were acoustically characterized using a clinical MRgFUS system (Philips Sonalleve V1) and a needle hydrophone at 1.2 MHz. Using the 256 elements of the transducer, a mapping of the Time-Of-Flight (ToF) delays after insertion of the skull was obtained. The thickness of the skulls and the phantoms was estimated using the CT datasets. The average speeds of sound (SoS) of the phantoms were compared to the SoS of the real skulls.

Results
Visually, phantoms and ex vivo skulls were similar. The neonate phantom even featured a fontanelle made of an ultrasound-transparent material. CT dataset comparisons showed a discrepancy between the thickness of the real skulls and the phantoms. The neonate phantom was on average 100% thicker than the ex vivo skull. The 8-y.o. skull phantom was 50% thicker than the ex vivo skull. The ToF delays obtained reflected these differences with an average of -0.4 μs for the neonate phantom vs -0.15 μs for the ex vivo skull and an average of -1.1 μs for the 8-y.o. skull phantom vs -0.7 μs for the ex vivo skull. The average SoS of the neonate phantom was 2250 m/s vs 2080 m/s for the ex vivo skull. The average SoS of the 8-y.o. skull phantom was 2500 m/s vs 2300 m/s for the ex vivo skull.

Conclusions
The bone-mimicking phantoms are a good alternative to rarely available ex vivo pediatric skulls. The thickness of the phantoms will be further adjusted in order to better mimic the real skulls. Future work will focus on the attenuation properties of these phantoms.

A130
Long-term transgene expression in rat brain after intranasal administration of hGDNF DNA nanoparticles and enhancement by Focused Ultrasound (FUS)
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Objectives
The therapeutic potential of glial cell-line derived neurotrophic factor (GDNF) for treating Parkinson’s disease (PD) has been limited thus far by its inability to cross the Blood-Brain Barrier (BBB). We have previously shown that intranasal administration of PEGylated lysine 30-mer (CK30PEG10K) DNA nanoparticles (NPs) encoding hGDNF, developed by Copernicus Therapeutics, Inc., can transfect brain cells in vivo, induce transgene expression, and provide neuroprotection of substantia nigra (SN) dopamine neurons in the rat 6-hydroxydopamine model of PD. We have also shown using double-label immunohistochemistry that transgene expression in the rat brain occurs primarily in cells lining the vasculature, most likely pericytes. Although intranasal administration allows large biomolecules to bypass the BBB, it is a low efficiency route and results in widespread distribution with no means of targeting specific brain regions. Here we investigated whether focused ultrasound (FUS) combined with circulating microbubbles can enhance intranasal delivery of pGDNF NPs to the brain.

Methods
The first goal of our current study was to assess the duration of transgene expression after intranasal administration of DNA NPs. We used a reporter plasmid (pUGG), which produces an eGFP-GDNF fusion protein. All rats received 90 μg of pUGG NPs, a dose that we previously showed yields significant eGFP expression throughout the rat brain 1 week after intranasal administration. Groups of 8 rats were sacrificed at 1 week, 3 months and 6 months after intranasal dosing. Separate groups received intranasal saline and were sacrificed at the same time points.

Our second goal was to evaluate whether FUS can be combined with intranasal administration of pUGG NPs, as well as the naked plasmid, to yield greater overall delivery, improved tissue penetration, and enrichment of transgene expression in the desired location(s) in brain. To date, we have conducted a pilot study in 3 rats given 180 μg of naked pUGG DNA intranasally. FUS bursts (32 msec bursts at 4 Hz for 100 sec) at 274 kHz were applied during infusion of Optison microbubbles to one brain hemisphere before intranasal delivery, and repeated 1 hour later at a caudal location on the same side after intranasal delivery. The rats were sacrificed 1 week later.

Expression of eGFP was assessed throughout the brain by eGFP-ELISA.

Results
In the time course study, ELISA revealed that whole brain eGFP expression (above background in saline-treated controls) was highest at one week, and persisted at ~30% of maximal expression at both the 3 and 6 month time points (p<0.05 for 1 week versus 3 and 6 months). These results provided evidence that intranasal delivery of Copernicus’ pDNA NPs results in significant and prolonged transgene expression in rat brain, with highest levels at 1 week and continued expression over at least 6 months.

Results from the pilot study in rats which received naked plasmid intranasally, in conjunction with FUS and microbubbles, indicated that the amount of transgene expression was increased on the sonicated versus the unsonicated side of the brain, and overall whole brain eGFP weighted averages were significantly increased in the sonicated rats. These results suggest that FUS with microbubbles may increase the overall delivery of DNA NPs to the sonicated brain regions.

Conclusions
In conclusion, FUS may enable agents with poor capabilities of crossing the BBB, e.g. neurotrophic factors, and viral and non-viral vectors encoding them, to become disease-altering therapies by a non-invasive route of administration. Our ongoing studies are examining the effect of FUS on transgene expression in the sonicated regions, and will assess tissue penetration and cell types transfected after intranasal administration of our DNA NPs.

A131
The comparison of different FUS parameters mediated therapeutic responses in a subcutaneous model of melanoma
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Objectives
Melanoma has a 5-year survival rate of <10%, primarily due to therapeutic resistance and immune tolerance. Durable responses to therapy of metastatic melanoma are enhanced when the anti-tumor immune response is activated. FUS is an emerging non-invasive treatment modality for localized treatment of cancers. Damaged tumor cells have been shown to release endogenous danger signals, which could stimulate an immune response, implying that the patient’s dying cancer cells serve as a therapeutic vaccine that stimulates an anti-tumor immune response. Specifically, we hypothesize that FUS will augment tumor antigen release and danger signals to mediate rejection of both FUS treated and untreated tumors. We believe that this approach will increase the percentage of patients who
will respond to immunotherapies by stimulating a melanoma-specific immune response. However the optimum parameters to simulate this response is mostly unknown. Therefore we conducted a preliminary experiment to compare different FUS parameters with biological responses.

**Methods**

Studies using C57BL6 mice inoculated subcutaneously with B16 melanoma cells were approved by the institutional animal care and use committee. When tumors became palpable, mice received FUS treatments (two sites per tumor) using a 1.14 MHz transducer (FUS Instruments, Canada). FUS treatment was applied either in pulsed (group-1: 44 W, 10 ms pulse duration, 1 s intervals, 60 s total duration at 7 MPa or group-2: 44 W, 14 ms pulse duration, 0.5 s intervals, 120 s total duration at 7 MPa or in continuous mode (group-3: 15 W, 10 s, 4 MPa or group-4: 44 W, 3.5 s, 7 MPa). The temperature maps were generated for treated tumors. Following FUS treatment, necrosis, and apoptosis were measured. Standard H&E staining was performed to evaluate histological changes that may have occurred as a result of the FUS exposures. The detection of apoptotic cells was performed using staining for Caspase-3.

**Results**

We observed that the temperature rise for tumors treated in continuous mode (group-3) were in the range of 55-65 °C and temperature of pulsed tumors (group-1 and 2) were approximately ~40 °C. We were unable to measure the temperature of group-4, due to the tumor movement as result of FUS treatment. We observed increase in both necrosis and apoptosis with FUS treatment (group-1, 2 and 3) when compared to untreated tumors.

**Conclusions**

Our data shows that FUS parameter used in the group-4 cannot be adopted due to technical difficulties that arise due to movement of the tumor. However our data shows that group-3 conditions give rise to increase in temperature when compared groups-1 and 2. Additional mouse studies are currently underway to confirm these results and to characterize and compare the immune response elicited in the group-1, 2 and 3.

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**A132**

**Evaluation of bone ablation size as a function of sonication time during MRgFUS**

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**Objectives**

One challenge in MRgFUS ablation of bone tumors is extending the depth of bone ablation beyond the cortical surface, which rapidly attenuates sound. Anecdotally, providers will lower ultrasound frequency or increase sonication duration, in order to achieve better penetration. The purpose of this study was to evaluate the effect of longer versus shorter sonication durations on the size of the post-ablation appearance in a swine model of MR guided Focused Ultrasound ablation of bone, for a given total energy.

**Methods**

Experimental procedures received approval from the institutional committee on animal research. MRgFUS was used to create two ablation foci (distal and proximal) in the left femoral diaphysis of 6 pigs. The spacing of sonications was equivalent between the two foci. Both targets were subjected to six sonications with 400 J of energy each; the distal targets were dosed with 20 W for 20 seconds (standard time) and the proximal targets were dosed with 10 W for 40 seconds (long duration). The hypoenhanced ablation zone was then measured on post-contrast MRI sequences in three dimensions.

**Results**

w7-MRGFUS created focal hypoenhanced lesions at the distal and proximal targets. Interestingly, the use of the conventional 20 second duration sonication resulted in the largest depth of the transverse intramedullary hypoenhanced zone and the craniocaudal dimension of the ablations, which measured up to an average of 7.3 mm and 26.7 mm, respectively. By comparison, the mean ablation measurements for the 40 second long duration sonication group were 4.5 mm and 21.0 mm: these differences reached statistical significance (paired t-test, p = 0.026 and 0.006). There was no significant difference in the anteroposterior measurements between the two groups.

**Conclusions**

While different techniques can and should be used to maximize the size of the ablation zone in MRgFUS of bone lesions, these results suggest that increasing the sonication duration, while concomitantly decreasing the acoustic power to maintain a given total energy, is not an effective technique and may be counterproductive. These results can be used to achieve more complete MRgFUS ablations of bone lesions.
Conclusions
Establishing whether ThermoDox combined with local HIFU-induced hyperthermia is safe and tolerable, is the first step towards potential future replacement of surgery in a selected group of high-risk, primary breast cancer patients.

Retrospective study of patient suitability for MR-HIFU liver therapy
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Objectives
Liver cancer treatments range from radical surgery to minimally-invasive treatments like radiofrequency ablation. Severe side effects of liver surgery limit its use to possibly curable patients, while inadequate precision and monitoring of image-guided minimally-invasive methods limit their effectiveness. These methods also carry the risks of ionizing radiation, infection, and inadvertent off-target treatment. Shortcomings of current therapies may be addressed with MR-HIFU, which permits exacting, non-invasive tumor ablation without ionizing radiation. Following recent advancements in organ motion tracking and treatment approaches to MR-HIFU treatment of highly perfused organs, a need exists to define eligibility criteria for liver MR-HIFU. However, patient suitability for liver MR-HIFU is currently not well defined. Factors including the degree of underlying liver dysfunction, tumor extent, and anatomic accessibility may influence MR-HIFU inclusion and exclusion criteria. This study seeks to identify suitable candidates for MR-HIFU and its possible combination therapies based solely on HIFU beam access to tumor.

Methods
We assess the theoretical suitability of patients for MR-HIFU by retrospectively evaluating the medical and treatment-planning records of patients with liver tumor. All patients were enrolled at the Center for Interventional Oncology at the Clinical Center in National Institutes of Health between January 2015 and January 2016 and have had known, non-MR-HIFU treatments. The institutional review board approved this retrospective chart review study and informed consent was waived. All medical records were anonymized and de-identified. A blinded clinician segmented computed tomography (CT) scans with thin slices using 3DSlicer. Segmented volumes were exported to MatLab where a semi-automated algorithm evaluated whether the tumor(s) would be accessible by an MR-HIFU platform. Treatment feasibility was evaluated on both a clinical tabletop HIFU system (Sonalleve V2, Philips, Vantaa, Finland) and a freely positionable extracorporeal mock HIFU transducer. Patients were categorized into the following treatment groups: 1) full ablation of the tumor; 2) combination of HIFU and trans-arterial chemoembolization (TACE); 3) full or partial sonication of tumor for immune response evaluation; and 4) HIFU mediated targeted drug delivery. The categorization of patients into these therapeutic arms was based upon size, volume, number, and distribution of disease.

Results
Data for 250 patients with liver mass were analyzed, and suitability for HIFU therapy evaluated. We aim to report patient and tumor characteristics, percentage of patients eligible for MR-HIFU, and the distribution of non-HIFU treatments the patients received. Data analysis is pending.

Conclusions
MR-HIFU may improve existing treatment algorithms by offering patients a precise, non-invasive choice in both curative and palliative settings. Randomized control studies are required to evaluate the true potential of HIFU as a therapy modality. To better inform patient enrollment strategies, our retrospective study aims to define characteristics of a suitable candidate for MR-HIFU liver therapy.

Experimental assessment of phase aberration in MRgFUS breast treatments
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Objectives
MR-guided Focused Ultrasound (MRgFUS) has the potential to be used as a non-invasive breast cancer therapy. Clinical feasibility has been demonstrated in multiple studies and hardware development has progressed in addressing the challenges of treating the breast. It has been demonstrated in a simulation study that phase aberration caused by the patient-specific heterogeneous anatomy of the breast may decrease the efficiency of MRgFUS breast treatments by causing unwanted heating when using a small phased-array transducer (256-elem, 1-MHz, 14.4 x 9.8-cm aperture). The purpose of this work is to develop tissue-mimicking heterogeneous breast phantoms in which the simulated effects of phase aberration and phase aberration correction techniques can be experimentally verified.

Methods
Phantom: A heterogeneous phantom was constructed that mimics the breast anatomy and adequately replicates the acoustic properties of breast fibroglandular and fat tissue. The phantom consists of 250-bloom gelatin with a 1 to 1 mixture of water and evaporated milk. Six latex balloons (diameter = 1.5-2.0 cm) filled with canola oil were dispersed throughout the phantom. The phantom was mounted in our breast MRgFUS system and 3D MR images were obtained to create a numerical model of the phantom (3D 2-point Dixon). MRgFUS heating schemes were tested: a single-point sonication (23 W, 30 s) and a 9-point raster-scanned trajectory (2-mm spacing, 32 W, 20 s/point, no cooling). Both heating schemes were performed with occlusions in the near field. To evaluate different acoustic windows, these sonications were repeated at two different transducer locations. MR thermometry was used for real-time monitoring (3D seg-EPI GRE). Simulations: Using the hybrid angular spectrum method applied to the generated numerical model, simulated pressure patterns were obtained both with and without applying a phase correction technique. Temperature response was modeled with a finite-difference implementation of the Penns bioheat equation.

Results
Figures 141, 142 and 143 show the comparison of experimental and simulated results for the single sonication case. In all cases the temperature map is overlaid on the phantom’s magnitude image or the numerical model, and the transducer (not shown) is located to the left for all images. The experimental temperature response is shown in an oblique slice in Fig. 141. The focal spot is clearly aberrated and resulted in low temperatures (peak = 9.5 °C) at the desired focus. Figures 142 and 143 show the simulated temperature maps for a single sonication at the same location without and with phase aberration correction, respectively. The simulation of the focal spot after phase correction is applied shows a dramatic improvement in energy delivery. The aberrated focal spot resulted in a much lower temperature rise (scaled to match experimental) when compared to the focal spot with phase correction applied (peak = 27.6 °C). Also, the peak temperature was 3 mm in front of the desired focal location for the single sonication without phase correction and at the desired location with the correction applied. A similar trend for the peak temperatures and focal location accuracy was seen for the 9-point trajectory.
Conclusions
This work presents a heterogeneous anatomy-mimicking breast phantom that produced aberrations both experimentally and in simulation. The correction of these aberrations in simulation demonstrated an increased temperature rise and more accurate focal location. Work is ongoing to apply the phase correction technique experimentally. Additional heterogeneous aberrating breast phantoms are under construction to further validate this technique.

Fig. 141 (abstract A122). A coronal oblique slice of the heterogeneous aberrating breast phantom is shown with the experimental MR temperature map (PRF-shift method) overlaid on a T1-weighted image. The white scale bar represents 5 cm

Fig. 142 (abstract A135). The simulation on the segmented numerical model obtained from the same aberrating phantom as shown in Fig. 141 is shown with temperature maps overlaid for uncorrected phases. The white scale bar represents 5 cm

Patient preparation influence on respiratory compensated PRFS based MR thermometry performance in the pancreas
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Objectives
Magnetic Resonance Imaging-guided High Intensity Focused Ultrasound (MR-HIFU) has recently been suggested as an alternative treatment modality, for pancreatic cancer patients in cases where surgery is not an option. MR-guidance allows to monitor temperature for feedback control, however using proton resonance frequency shift (PRFS) based thermometry is challenging, because of unpredictable motion (peristaltic) and the presence of air in the digestive tract near the head of the pancreas. This can lead to changes in the magnetic volume susceptibility distribution resulting in artifacts in the temperature maps. Solutions have been proposed to this end, and even if it is tempting to transfer methods from one organ to another it is not necessarily straightforward.

In this study, we propose an approach aimed at reducing susceptibility heterogeneity by filling the stomach and duodenum with fluid, and evaluate its effect on PRFS based temperature mapping using different known methods for respiratory compensation.

Methods
Experiments were performed on a 1.5-T MRI scanner (Achieva, Philips Healthcare, The Netherlands) equipped with a clinical MR-HIFU system. Experiments were executed twice on healthy volunteers (n = 7), first, without any special preparation. Then during a different exam, volunteers were asked to ingest 2 * 500 mL of fluid with favorable relaxation times (pineapple juice) to fill the tract, 5 hours and 2 to 5 minutes prior to scan, respectively. A breathhold 3D fat suppressed T1-weighted scan was acquired for planning. For temperature mapping, a dynamic gradient echo series consisting of 3 coronal slices covering the pancreas was acquired. Scan parameters: TR = 100 ms, TE = 19 ms.
FA = 21, reconstructed voxel size = 2.5x2.5x8 mm³, dynamic scan time = 400 ms, 500 dynamics.

Statistical analysis: Datasets were analyzed offline using three techniques, viz. gating2, multibaseline4, and referenceless5. The temperature standard deviation (TSD) over time was calculated voxel-by-voxel. As reference the whole pancreas was semi automatically segmented on the T1 image and used as ROI for statistical analysis. For each technique the volume of pancreas under the threshold value of 2 °C precision was compared to the total pancreas volume in our images leading to a percentage of coverage at 2 °C.

Results
Figure 144 shows an example of the juice filling effect in one of the volunteer. Without juice the TSD varied from 2 to 10 °C, within our three methods. After juice intake, the TSD range went down to values between 1 to 2 °C and the spatial distribution became clearly more homogeneous.

In Figure 145, preparation without liquid is compared to preparation with juice, for every technique the coverage of the pancreas with a precision under 2 °C is increased by 25 % in average. It allows to reach values, up to 80 % with multibaseline and gating.

Conclusions
In all volunteers, oral ingestion of pineapple juice for patient preparation reduced the range of values in the TSD leading to an increase of the precision coverage in the pancreas. Filling the digestive tract in this way could be an attractive solution for more precise temperature monitoring during MR-HIFU therapy in the pancreas.

Quantitative magnetic resonance imaging of ultrasound induced transient shear waves
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Objectives
Tissue mechanical properties are altered both in disease states and following ablative, or other, therapy. Changes in mechanical tissue properties are manifested in changes to the speed of a shear wave propagating through the tissue. Transient shear waves can be induced in tissue via application of short bursts of focused ultrasound (US). Imaging the propagating shear wave with MR confers the benefits of that modality’s excellent soft tissue contrast, 3D visualization, and the ability to assess tissue immediately following US ablation. Here, we quantify tissue elasticity using constructive multi-pulse transmission under MR and US guidance and validate with mechanical testing.

Methods
MR measurement of transient shear wave (SW) propagation was performed using a 7 T MRI (Bruker Biospin, Germany) and a 3 MHz 16-element annular array (IMASONIC SAS, France, 48 mm diameter, 35 mm radius of curvature, 0.5 x 0.5 x 2 mm³ focal volume at -6 dB), controlled by an embedded MR-compatible 2D positioning and US system (Image-Guided Therapy, France). Three foci (3 MHz, 450 ms pulses with 4.5 mm spacing between transmit foci, 12.5 MPa PNP) were sequentially excited along the beam axis creating a planar SW which was imaged with MRI (Fig. 146a, TR/TE/FA = 500 ms/26.1 ms/180°). A series of images was acquired with different propagation times, varying from 0.45 ms to 4.05 ms and the shear modulus was estimated using the time-of-flight (TOF) method. The US-based SW measurement was performed with a Vantage 256 (Verasonics, WA, USA) and a Phillips L7-4 probe. To generate the SW, 4.3 MHz, 10 MPa PNP, 100 μs pulses were transmitted with 5 mm spacing between transmit foci. US images were obtained with a 5.2 MHz center frequency and 10 kHz PRF and the local inversion method (LIM) was used to estimate the shear modulus. Phantom cylinders (9.7 mm in diameter and 8.5 mm)

Results
The propagating SW was visualized with both MRI (Fig. 146a) and US (Fig. 146b) and quantitative elasticity maps (Fig. 146c) were generated. Using multi-pulse transmission, the shear modulus in a 1.5 % agar and 1.5 % gelatin phantom was estimated as 27.3 ± 2.3 kPa with MR and 28.2 ± 2.3 kPa with US, which agreed with 32.3 ± 1.8 kPa yielded by compressive testing (Fig. 1D).
Conclusions
In conclusion, the shear modulus in tissue-mimicking phantoms was accurately estimated using the combination of an MR shear imaging sequence and multi-pulse transmission.

Fig. 146 (abstract A137). Elasticity maps from a MR and b ultrasound. c Strain-stress results based on mechanical testing. e The estimated shear moduli were 28.2 ± 2.3, 27.3 ± 2.3 and 32.3 ± 1.8 kPa in US, MR and mechanical testing, respectively

A138

In vivo biodistribution studies of drug-loaded nanodroplets for prostate cancer management using ultrasound-mediated drug delivery under MR-guidance
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Objectives
Our ultimate goal to initiate the present study is to develop a novel approach for treating prostate cancer with improved efficiency while reducing normal tissue toxicities. In this treatment approach, the drug is delivered into the tumor via encapsulation inside an ultrasound-sensitive, drug delivery system (nanodroplets). Once the system reaches tumor, MR-guided pulsed High Intensity Focused Ultrasound (MRg-pHIFU) is applied to the tumor to help activate the release of the drug from nanodroplets inside the tumor, thus increasing the bioavailability of the drug.

The purpose of the present work is to study the biodistribution of paclitaxel-loaded nanodroplets in vivo in nude mice. We consider our present study to be of high impact because it tries to answer specifically for the first time an important question of appropriate timing to apply tumor-targeted focused-ultrasound for activating release of drug from these drug carriers for effective prostate cancer treatment.

Methods
Poly (ethylene oxide)-co-poly (D, L-lactide) (PDLA) nanodroplets loaded with fluorescently labeled-paclitaxel (F-PTX) were synthesized using solid dispersion technique. Human prostate cancer, LNCaP cells were implanted orthotopically in prostates of male nude mice. Tumor-bearing mice (n = 3) were injected with 0.1 % F-PTX, 2 % PDLA nanodroplets using tail vein. At chosen time points (30 min; 2 h; 4 h; 6 h; 12 h; 24 h) animals were anesthetized, blood was collected by heart puncture, liver, spleen, kidney lung and heart were excised, cut and weighed. The lysates were read using fluorescence spectrophotometer (Excitation- 496 nm, Emission- 524 nm). Fluorescence readings were compared with values from the standard calibration curve. Tumor-bearing mice (n = 3) were also injected with 0.1 % F-PTX solution (fluorescent drug in free form).

Results
Quantitative analysis showed 20 % of the injected drug-loaded nanodroplets in the tumor, 30 min after systemic injection; which increased to 30 % after 2 h; 35 % after 4 h; then decreased to 27 % after 6 h; 11 % after 12 h and 5 % after 24 h. When drug in free form was injected, drug accumulation in the tumor was 7 % within 30 min; 8 % within 2 h; 10 % within 4 h; 9 % within 6 h, 3 % within 12 h and 2 % within 24 h. Liver showed major drug accumulation for drug-loaded nanodroplets while heart and lung showed less. For drug in free form, liver and spleen showed maximum drug concentrations while heart showed concentrations similar to that in tumor.

Conclusions
Treating prostate cancer with MRg-pHIFU mediated drug-delivery using ultrasound-responsive, drug-delivery vehicle (nanodroplets) is innovative and aims at minimizing toxic effects to healthy tissues, as caused by chemotherapeutics drugs and maximizing better treatment efficacy. Present bio-distribution indicates that the optimum time point for applying focused ultrasound is after 4 h of systematically injecting drug-loaded nanodroplets. Using this optimum time point, in vivo experiments in nude mice are being carried out to test the efficacy of combined MRg-pHIFU exposures and chemotherapeutic agents encapsulated nanodroplets for prostate cancer therapy.

A139

From research to clinics: an example of clinical translation of MR guided FUS treatment of the liver
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Objectives
There are many groups around the world working on MR guided FUS research topics to enhance the treatment for patients and to advance the technologies. The main goals of the research is to provide faster and more accurate algorithms and tools while ensuring the clinical safety, efficiency and effectiveness. In the European funded TRANS-FUSIMO project (www.trans-fusimo.eu), the consortium is currently developing a quality assured treatment system for MR guided Focused Ultrasound in moving abdominal organs which will be validated in a pre-clinically tested. The target of this project is to translate a clinical study using the developed technologies. Here we will discuss the quality assurance that is needed for translation of a research tool towards a clinical study.

Methods
If newly developed features or technologies from research shall be translated to clinical use, there are many obstacles on the way. In particular if the research system shall be used in a clinical study it has to comply with high quality standards. It is also possible to perform clinical studies without a CE mark but the medical device must have a quality status that is CE mark ready. The most important norm is 93/42/EEC which is the European medical devices directive. This directive also asks for an underlying quality management system which can be EN ISO 13485. This means that the parties, which are responsible for this medical device have to be qualified according to EN ISO 13485. If a software tool is the critical part of the clinical device, its development and implementation has to conform to IEC 62304, which is the international standard for medical device software. Furthermore, a risk management for the medical device according to ISO 14971 has to be performed. Only if development and validation are conducted according to these standards authorities can approve a clinical investigation of the medical device.

Results
In the work presented here, we are translating software, which is able to communicate with an MR device as well as an HIFU device. The HIFU device and HIFU system have CE marks or are at least CE mark ready for their intended use, the new research communication software not, still it shall be used in a clinical trial. The software shall have the ability to follow the breathing motion of the patient and steer a focal spot in the liver accordingly. Since this software without CE mark shall be used in a clinical study we must follow the above
described quality assurance. In our case the software with its sub-components is developed according to the EN ISO 13485 norm and also the preclinical in vitro and ex vivo validation is performed according to this quality management system. Furthermore, since the software acts as the control system of the HIFU device it is a critical part of the system and thus the development and implementation has to conform to IEC 62304. Finally, a risk management for the medical device according to ISO 14971 was created.

**Conclusions**

If a clinical investigation is anticipated, CE mark status is necessary. The example presented in this work shows how to use a non CE marked software tool in conjunction with existing qualified hardware. Also a preclinical validation is mandatory in order to ensure the safety, efficiency and effectiveness for the patients during the treatment.

**A140**

Magnetic resonance-guided focused ultrasound in treating painful bone metastasis: preliminary results of a phase IV study

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**Objectives**

Bone metastasis represents one of the major causes resulting in cancer pain. Traditionally, conventional radiotherapy serves as the standard treatment for painful bone metastasis. Recently, magnetic resonance-guided focused ultrasound (MRgFUS) has been demonstrated as an alternative therapy for pain palliation of bone metastasis. Therefore, this phase IV study was conducted to evaluate the long-term safety and efficacy of MRgFUS treatment for patients with painful bone metastasis.

**Methods**

Current ongoing single-institute phase IV study has enrolled 20 patients at Taipei Cancer Center, Taipei Medical University in Taiwan since Nov 15th, 2014. The MRgFUS is delivered by ExAblate 2000 system (InSightec Ltd., Israel) which integrates a high-intensity focused ultrasound compartment for hyperthermia treatment with a magnetic resonance imaging scanner for real-time position and temperature monitoring. The treatment response is evaluated by the interval change of numerical rating scale (NRS) before and after treatment. Pain relief is defined as a reduction of 2 points or more on NRS pain scoring system without increasing analgesics use. Patient's quality of life is assessed by the Brief Pain Inventory (BPI-QoL) score. The adverse event is also documented.

**Results**

The median age of the 20 treated patients (12 male and 8 female) was 62 years old (range, 40–83). Each patient underwent MRgFUS treatment for single painful bone metastasis. Sixteen targeted lesions were located in the pelvic bone, whereas 3 at rib cage and 1 at scapula. At a median follow-up duration of six months, none of the patients experienced treatment related adverse events above grade 2 according to Common Terminology Criteria for Adverse Events version 4.0. Significant improvement of NRS pain score and BPI-QoL score is observed after treatment.

**Conclusions**

The initial results of current study demonstrate the safety and efficacy of MRgFUS in treating painful bone metastasis. MRgFUS has the potential to serve as a first-line treatment in selective patients with painful bone metastasis.

**A141**

Heating position accuracy assurance of a clinical HIFU treatment for bone metastasis via MR-guided Focused Ultrasound system

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**Objectives**

Utilizing magnetic Resonance-guided Focused Ultrasound Surgery (MRgFUS) to treat metastatic bone tumor is an advance therapy for palliative pain control. MRgFUS system performs High Intensity Focused Ultrasound (HIFU) with real-time MRI monitoring and results in local high temperature hyperthermia. In this study, we assessed quality assurance concepts to endorse accuracy of thermal dose and position stability. Furthermore, this research tried to share experience of calibration protocol, and to provide permissible criteria according to cases treated so far. Moreover, the goal is to establish standard operating procedure of examining accuracy of focal spots, in order to provide patients a more qualified treatment.

**Methods**

In this research, we measured focusing error distances between focal spots and real heating spots from 31 cases in Taipei Medical University Hospital to identify the effectiveness of Daily Quality Assurance (DQA) for preventing focusing error in MRgFUS. MRgFUS applies Magnetic Resonance Imaging (MRI) to locate metastatic bone tumors, uses heating technique of focused ultrasound to convey ultrasound energy to painful bone area. The energy would heat the area up to 60 degrees Celsius and ablative nerve fibers on the surface of bone to achieve the purpose of palliative pain control. Advantages of this treatment include non-invasive, non-irradiative, and long-term pain relief in single treatment.

**Results**

Comparing the differences before and after quality assurance, focusing errors in DQA group were observed consistent diminutions, which reached statistical significance (p-value < 0.01). Nevertheless, DQA procedures also benefited heating accuracy in practice. The 95 % confident interval of accuracy were reported from 1.73 ± 0.43 mm (Pre-DQA group) to 0.43 ± 0.12 mm (DQA group).

**Conclusions**

In summary, a comprehensive DQA standard guarantees treatment accuracy and patient safety. As a suitable calibration procedure, DQA was evaluated its advantages to reduce focusing errors and unexpected heating spots. Therefore, DQA procedures were highly recommended to adjust each distance between focal spot and real heating spot within 1 mm before treatment.

**A142**

Simulation and evaluation of enhanced thermal necrosis distribution via high intensity focused ultrasound hyperthermia therapy

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**Objectives**

High intensity focused ultrasound (HIFU) has been reported for its non-invasive heating capability in tumour treatment. As an advance application, magnetic Resonance guided focused ultrasound (MRgFUS) system performs High Intensity Focused Ultrasound (HIFU) with real-time MRI monitoring and results in local high temperature hyperthermia. However, despite correct delivery of thermal dose to treatment area within a short period, to protect nearby critical organs real time temperature should also be monitored. Since local temperature over 56 °C heat shock may lead to thermal tissue necrosis. In this study, gold based micro particles were designed as an enhanced for ultrasound energy absorption in tumor lesions. Through our thermal dose simulation, we found that intra-artery micro enhancer is able to improve thermal dose response during routine HIFU treatment. Furthermore, we tried to share experience of thermal necrosis, and provided a feasible strategy to establish novel treatment.

**Methods**

ExAblate 2000 system (Insightec, Israel) installed in Taipei Medical University and was used to treat bone metastasis for severe pain patients. This system contented a 1.5 T GE magnetic resonance system and a phase array transducer with 208 independent HIFU sonication
elements. We simultaneously used an insulated thermocouple wire (Thermoway co., Taiwan) to detect true temperature and compared with PRF temperature. Based on this HIIFU system, SAS human oral cancer cell xenograft in nude mouse (National Laboratory Animal Center, NLAC, Taiwan) served as the animal model in this experiment. 6 weeks old nude mice were injected SAS cancer cells in both hind legs and formed about 1 cm tumor. Micro particles conjugated polyethylene glycol (PEG) were inject via intraportal injection and conducted higher affinity with tumor. The ablation target was located in the center of pathology tumor volume (PTV) and sonicated by 300 J energy. Statistical analysis distinguished significant differences for various dose of particles and PRF temperature. Moreover, multiple regression and Matlab models validated correlation of dose bias.

Results
It is observed that micro particles promote both MR image resolution and focused ultrasound absorption in target soft tissue. As was shown by man MR signal, that photon intensity (PD) calculated within pathology tumor volume (PTV) in particle injected group (PD = 71.2) was higher than that of sham group (PD = 47.7) and blank group (PD = 67.8). After HIIFU sonication, via implementation of PRF algorithm, temperature distribution at target treatment area was observed to increase form 58.1 ± 3.6 °C (control group) to 63.5 ± 6.3 °C (particle injected group), with statistical significance from Wilcoxon rank-sum test (p = 0.03) between control group and particle group. In addition, from imagine evaluation, post sonicated tissue also shown a dark area in the center of tumor which indicates thermal necrosis of ablation, implying thermal damages that block blood circulation in tumor vessel.

Conclusions
In summary, a comprehensive simulation has demonstrated that enhanced tumor necrosis in local lesions generated benefit effects for deep tumor controlling. Nevertheless, we also discovered that micro enhancer is able to improve thermal energy absorption during routine HIIFU treatment. This finding indicated that enhanced thermal dose distribution within solid tumor is feasible for first line tumor treatment.

A143
Quantitative T1 mapping of breast cancer xenografts during HIIFU ablation
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Objectives
A current interest in the field of clinical MR-guided HIIFU tumor treatments is the ability to acutely assess the extent of tumor necrosis. Previous studies have investigated MR-derived multi-parametrics such as the apparent diffusion coefficient and contrast-enhanced T1 and T2 maps as indicators of cell death within minutes of ablation. In the field of radio-frequency ablation, investigators have used the non-contrast T1 mapping technique, modified look-locker inversion recovery (MOLLI) to predict areas of complete thermal ablation in cardiac tissue. The following study investigates the use of this quantitative T1 imaging technique as a single-parametric predictor of thermal dose achieved in human breast cancer tumors injected in mice.

Methods
T1 maps were acquired using the MOLLI method pre- and post- partial thermal HIIFU ablation of human breast cancer tumors grown in NOD-SCID mice. Cell suspensions (1 x 106 cells) of one of two human-derived triple negative breast cancer cell lines, HC002 and, were surgically injected into the mammary fat pads of 16 female NOD-SCID mice (n = 8/ cell line), and grown to an average volume of 1609 mm2 or 558 mm2 (HC002, HC019, respectively). After localization, a single slice pre-ablation T1 map was acquired (MOLLI). For HIIFU treatment, four to nine ablation points (8.3-14.9 W, 30 s) were applied to the center region of each tumor with a small animal MR-compatible HIIFU system (Image Guided Therapy, f = 3.0 MHz, FWHM = 1x3 mm), while temperature rise was measured with MR temperature imaging (segmented EPI GRE). Finally a post-ablation T1 map was acquired post-ablation at 20 or 120 s post-ablation (HC002, HC019, respectively). Thermal dose of the total treatment was calculated using the CEM@43 °C metric from the MR temperature data, assuming a tumor baseline temperature of 30 °C. ROIs for T1 maps were selected via manual registration with temperature imaging DICOMs (Osirix).

Results
As shown by Fig. 147, there is a consistent trend of increased T1 values in the heated region of the tumors post-ablation. In contrast, the background (control) signal remained relatively constant. Additionally, a small ROI in a region of each tumor not directly heated with HIIFU was selected for comparison to the heated regions (Fig. 148). Calculating the percent change in T1 values resulted in P-values of 0.0709 and 0.0136 between the heated and non-heated tumor and control ROIs, respectively. The less significant difference between the heated and non-heated ROIs may be due to the large variance in local tumor changes for both groups, as well as a limited ability to select non-heated regions distant (>3 mm) from the heated regions, since the diffusion of HIIFU heating likely contributed to local T1 changes. Finally, the correlation between thermal dose and T1 change is shown (Fig. 149). While there is a correlation between thermal dose and T1 value for the HC002 cohort, which accumulated a lower thermal dose levels and also at a later time post-ablation, the HC019 cohort, which accumulated a lower thermal dose, shows no correlation.

Conclusions
Preliminary results of this breast cancer xenograft study shows that localized changes in quantitative T1 maps can be observed as a result of HIIFU ablation within two minutes of treatment completion. Additionally, T1 values were not shown to significantly decrease with increased cooling time, indicating presence of a T1 effect independent of temperature change (data not shown). Finally, the observed T1 change may be correlated with achieved thermal dose, particularly if time after treatment is increased to two minutes or at high thermal doses. Histology for directly determining cell viability in these tumors is ongoing.

Fig. 147 (abstract A143). Average T1 values from the heated tumor ROI and a background (control) ROI in a homogenous medium surrounding the tumor (either agar gel or pork loin) pre- and post-ablation (n = 16)
Improve immunity by high intensity focused ultrasound

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Objectives
HIFU is a method which cures local tumor by thermal effect of ultrasound. But Cancer is systemic disease. Even the local cancer is removed, the rest of the cancer cells will still be transferred to other parts. How to further improve the treatment effect is still a problem.

This paper discusses how to improve immunity by HIFU, to further kill the metastasis of cancer cells by immune function. The specific method is that controls the temperature 46 °C for half an hour at the tumor edge.

Methods
The method of this paper is based on the production of magnetic resonance imaging guided HIFU. The MRI guided HIFU device use MRI as positioning system and HIFU to treat cancer. This is a new generation of HIFU device, MRI image is much clearer than the ultrasound scanner as the positioning system. At the same time, because the MRI can measure temperature. So the device can control temperature by MRI and generate heat by HIFU, in order to achieve the treatment method of this program.

The HIFU is focused on the center of the tumor. When temperature is near 46 °C, the dose is reduced to maintain the temperature. This paper uses improved PID arithmetic to control temperature and discusses a lot of experiments in vitro, to verify the specific method.

Results
The device has achieved the HIFU immunotherapy of control the temperature of tumor edge which is in line with the intended purpose of the design. For irregular margin tumor, the error of the temperature will be increased. For the position of the vessel, the error of temperature will be further increased.

Conclusions
The immunotherapy can increase the effect of HIFU. So we should research this method further. Next step is to do clinical trials to verify the efficacy and safety of HIFU immunotherapy.

First steps of pre-clinical validation of Transfusimo MRgFUS treatment system for liver tumors

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Objectives
Trans-fusimo Treatment System (TTS) (Fraunhofer MEVIS) aims at applying Magnetic Resonance guided Focused Ultra Sound (MRgFUS) therapy to ablate liver tumours under respiratory motion through the rib cage. Sonication of a moving target is a very challenging task. TTS controls the Conformal Bone System (CBS) transducer of (ExAblate 2100, Insightec, Ltd, Tirat Carmel, Israel) for ablation of human tissue while monitoring using the 1.5 T GE MR Scanner. TTS is classified as high risk software as it controls high energy levels emitted by the CBS transducer.
into human body. For safe clinical use of the system, thorough pre-clinical validation of the software is mandatory. Preclinical validation is divided into two parts; first static case and as next step based on the working evidence of the first part, moving case scenarios. Specs were set with tolerance levels for delivered acoustic power, sonication duration, thermometry and sonicating to a planned position.

**Methods**

To validate TTS, novel protocols were designed and feasibility studies were completed for each spec. Acoustic power measurements were completed using the radiation force balance; following the standard IEC 61161(Fig. 151). Results were cross validated against the measured acoustic power delivery of ExAblate 2100 (Insightec, Ltd, Tirat Carmel, Israel). Sonication duration measurements were completed using the hydrophone for measuring the duration of actual sonication deviation from the planned, and the delay after the emergency stop (Fig. 152). Thermometry measured by TTS was validated against measurements of ExAblate 2100. Sonication positioning experiments were completed using polyacrylamide (PAA) egg white phantom by measuring the position of the scar after grid surface sonication’s. Measurements were completed both using external calliper (Fig. 153) and grey scale DICOM images. Each of these described protocols were repeated by two independent operators for 30 times for repeatability purposes. The Quality Management System ISO13485 has been applied to provide the required documentation for the approval for labelling the new system.

**Results**

For safe usage of the system, delivery of acoustic power should be within the tolerance level of efficiency (70-90%). TTS can deliver the acoustic power reliably in a controlled way within this specs (75 and 80%) as well as ExAblate 2100 (75 and 83%). Actual sonication deviation from the planned sonication duration was maximum -0.17 (Std 0.33) seconds meeting the spec being less than 1 seconds. Maximum sonication stop delay was 0.14 ms (0.16). Measured temperature values by TTS were relatively lower than ExAblate but still efficient enough to provide a scar for coagulation process to measure the deviation. Grey scale images were able to provide better resolution for distance measurement when compared to external calliper, maximum deviation being less than 1 mm. Results show that for static validations TTS successfully meets the criteria for the defined specs.

**Conclusions**

Due to regulatory requirements, it is mandatory to provide pre-clinical validation before any tests on animals take place. Designed protocols were able to provide traceability of the input and the output in a controllable way. Repeatability studies provide statistical evidence for the reliability of the static application of TTS software in pre-clinical settings. Based on these results, motion validation protocols are planned as a next step.
(Insightec, Ltd, Tirat Carmel, Israel) software. TTS has successfully met the set of specifications for delivering acoustic power, sonication duration, thermometry and sonicating to a planned location by using the designed protocols for validation in pre-clinical settings. The next step after Methods

A novel phantom, providing both temperature information and detecting sufficient amount of landmarks for tracking algorithm was developed. For measuring thermometry and observing coagulation procedure, the middle part of the phantom consisted of polyacrylamide (PAA) egg white material surrounded by 2 % agar and samphire to replicate the vein structure of liver. Echo Planar Imaging (EPI) was used during sonications to provide real time thermometry information using 1.5 T GE Scanner. To replicate breathing motion, phantom was moved within 20 -30 mm range by using the INNOMOTION Robotic arm (IBSMM, Prague). A special phantom holder was designed to provide the linear motion with the robotic arm while sonicating using the transducer of CBS. The experiment setup is as shown in Fig. 154. With this designed phantom holder, dual flex MR coils were positioned at a closer distance to each other for better image quality. Sonication was planned for 100 W and 30 seconds to observe the sonication scar. The Quality Management System ISO13485 has been applied to provide the required documentation for the approval for labelling the new system.

Results

Initial results show that samphire can replicate the vein structure for tracking algorithm in EPI scanning, without using any contrast agents. TTS can detect the vein structure as landmarks (Fig. 155) and sonicate to a planned location, by leaving a permanent scar to observe coagulation. The distance of coils to each other play important role in pre-clinical validation, due to noise problem caused by air in between the coils, which is not supposed be a problem in clinical settings. Designed phantom holder to place coils at a near location (16 cm in between distance) led into the successful application of tracking algorithm for sonication to a planned location. TTS can monitor the prescribed range of motion provided by the robotic arm during sonication, replicating the breathing motion for abdominal organs.

Conclusions

Pre-clinical validation is the mandatory step in qualifying a software controller TTS of MRI and the FUS system a class 3 device. Initial results show that TTS can function to track landmarks and provide thermometry information in real-time in pre-clinical settings. As a next step, repeatability experiments are planned in more detail to provide statistical evidence and prepare the first in human trial.

A147

Effect of transducer ground plane configuration on Magnetic Resonance radio-frequency coils and corresponding significance on image SNR during MRgFUS treatments

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Objectives

In MRI, the highest signal-to-noise ratio (SNR) is achieved by placing the radio-frequency (RF) coils as close to the region of interest as possible. In MRgFUS treatments, the transducer is often within the volume that would contain the ideal RF coil placement. In addition, if the transducer has a single, continuous ground plane, eddy currents can reduce the SNR in the region of interest. This study compliments the previous work that has been done to reduce gradient induced eddy currents. Using a similar approach this work investigates effects of ground plane eddy currents on RF coils by quantifying the resulting reduction of SNR. Finally, design strategies are outlined that are required to reinstate the diminished SNR.
Methods
A rectangular radio-frequency coil (20 cm x 13 cm) was built on a homogeneous cylindrical phantom (12 cm x 11 cm). The transducer ground plane was simulated by placing copper tape on the face of a 3-D printed plastic transducer (aperture: 13 x 10 cm, 10 cm radius of curvature). The ground plane was then broken into multiple copper patch elements by removing different numbers of thin copper strips (1 mm) in the short and long axis directions (Table 6 and Fig. 156). For each ground plane configuration, the RF coil loop was tuned and matched with the ground plane in position at 123 MHz with an insertion loss better than -35 dB. Active and preamp detuning were better than -35 dB and -20 dB, respectively. Using a Siemens 3 T MRI scanner (PrismaFit), SNR measurements were made using standard gradient echo sequences (TR/TE/FA/FOV = 500 ms/10 ms/90°/256 mm, 256 x 256 mm matrix). The homogenous cylindrical phantom was imaged with the single loop using each of the variations of the ground plane. SNR was calculated in the center of the phantom as well as near the transducer face using a sum of squares calculation.

Results
SNR results of various ground plane conditions are shown in Figs. 157 and 158. Quantitative SNR values measured in the center of the phantom and at the transducer's face are plotted in Fig. 159. SNR was reduced by 57 % at the transducer face when comparing the single element ground plane case to the no ground plane case. By segmenting the ground plane, the SNR was nearly recovered (95 %) while still maintaining a ground plane.

Conclusions
In general, increasing the segments in the ground plane increased the normalized SNR both at the center of the phantom and at the transducer face, as seen in Fig. 159. By incorporating this design principle into transducer ground plane design, eddy current effects could be reduced allowing for increase image SNR that would positively enhance both temperature and anatomic imaging during MRgFUS treatments.

Table 6 (abstract A147) Description of ground plane conditions

<table>
<thead>
<tr>
<th>Ground plane element configuration</th>
<th>No ground plane</th>
<th>4x1</th>
<th>1x4</th>
<th>3x3</th>
<th>3x2</th>
<th>2x3</th>
<th>3x1</th>
<th>1x3</th>
<th>2x2</th>
<th>2x1</th>
<th>1x2</th>
<th>1x1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total number of ground plane segments</td>
<td>No segments</td>
<td>4</td>
<td>4</td>
<td>9</td>
<td>6</td>
<td>6</td>
<td>3</td>
<td>3</td>
<td>4</td>
<td>2</td>
<td>2</td>
<td>1</td>
</tr>
</tbody>
</table>

Fig. 156 (abstract A147). Left image showing imaging setup with mock transducer (white) radio-frequency coils (copper) phantom holder (gray) on the right is a mock transducer with 1x3 ground plane configuration

Fig. 157 (abstract A147). Coronal SNR plots of a homogenous phantom under different transducer ground plane configurations. No transducer present (Left) and a Solid transducer 1x1 ground plane configuration (Right)

Fig. 158 (abstract A147). Coronal SNR plots of a homogenous phantom with a 3x3 transducer ground plane configuration (Left) and 1x2 transducer ground plane configuration (Right)

Fig. 159 (abstract A147) Plot of number of ground plane segments versus the normalized SNR in the phantom (center Xs and Transducer face Os)
A148
Improving intercostal HIFU through a novel transducer design: a numerical evaluation
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Objectives
Not released for publication

Methods
Not released for publication

Results
Not released for publication

Conclusions
Not released for publication

A149
High-Intensity Focused Ultrasound ablation in the treatment of hepatocellular carcinoma
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Objectives
Nonalcoholic Fatty Liver Disease (NAFLD) is a chronic liver disease which is manifested by steatosis, steatohepatitis, cirrhosis, and hepatocellular carcinoma. Patients with Nonalcoholic Steatohepatitis (NASH), which is a subset of NAFLD, have high risks of progressive liver disease and are of particular concern.

Methods
High-intensity Focused Ultrasound also known as HIFU can be delivered with precision resulting in necrosis of hepatocellular carcinoma (HCC) without a dysfunction liver. HIFU has some important advantages such as non-invasion and non-ionization. The heating effect produced by a focused ultrasound transducer should be checked by using a positioning system. SonoeIastography is a way to quantify fibrosis and it can compensate for.

Results
The results of this simulation indicated that with thermal analysis, according to tissue acoustic parameters in different frequencies, the desired temperature can be achieved for the destruction of cancer tissues and also the destruction of tumors.

Conclusions
The aim of this research was to evaluate the safety and efficacy of High-Intensity Focused Ultrasound therapeutic ablation of HCC as primary tumors in liver lesion that is in difficult location s such as main blood vessels, the heart, the gallbladder and bile ducts, or the stomach.

A150
A real-time data processing framework for steered FUS
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Objectives
FUS in moving organs can be realized by updating the focal spot position to follow the target motion (steered FUS). A key feature of such a system is the ability to predict the target motion for the treatment time. Real-time 2D imaging enables capturing some of the organ motion, which is quantified by consecutive tracking of anatomical features in the image stream (observation). All occurring delays need to be compensated for by introducing a spatio-temporal prediction of the potentially unobserved target region and surrounding structures. The final step in the processing pipeline is the update of the FUS control parameters derived from the prediction (control). A common problem is the tight coupling between observation and control loop.

Methods
While the observation rate is restricted by the update frequency of the imaging device, it is desirable to update the FUS focal spot position as often as possible to achieve the best possible approximation of the target trajectory given the hardware constraints. We propose an approach for decoupling observation and control loop by inserting an asynchronous motion model between the two loops. The motion model is continuously updated using tracking information while an independently running control loop queries the model for target position predictions and controls the FUS device. The system continuously measures the delays caused by all components (imaging, processing, FUS hardware upload) of both loops to calculate the time between the observed target motion and the actual arrival of FUS energy at the target (time of action). A spatio-temporal prediction is then calculated using the motion model for the time of action to compensate the delays.

Results
We propose a concept for decoupling observation and control in a steered FUS system. This enables the use of multiple imaging modalities with varying update rates to observe target motion. Further, the update rate of the focus position is thereby limited only by the FUS hardware and not by the imaging update rate.

Conclusions
Our System may enable a better compensation of the target motion by the steered FUS system. All delays between observation and control are measured continuously by the system and are being compensated for.

A151
Library sorting for real-time multi-baseline thermometry
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Objectives
MRFUS is typically monitored by MR thermometry, where temperature differences are calculated from phase differences with respect to a baseline-scan. For periodically moving scenarios, such as FUS in liver treatments, a popular method is the so-called multi-baseline method, where a set of baseline images covering the whole motion cycle (library) is acquired prior to heating. For a given monitoring image, temperature differences are then calculated with respect to the most similar phase-image within the library. Sparse sampling of the motion cycle will thus lead to errors in temperature estimation, dense sampling on the other hand leads to another problem: Searching through hundreds of library-images, acquired over several motion cycles, to find the most similar baseline image is computationally expensive and impedes real-time applicability. We propose a library sorting method that reorganizes the library-images acquired over several motion cycles to represent a single, but temporally densely sampled motion cycle.

Methods
In our method baseline library-images I_i, acquired over several motion cycles, are sorted into a single motion cycle by a three step pipeline: First, a motion cycles extreme position I_e is identified by computing an image similarity (S) between the first library image I_0...
and all other images and choosing the most dissimilar \( I_e = \text{ARGMIN}_I(S(I_0, I_e)) \). In the second pass all library-images are labelled with their similarity to \( I_e \): \( S_e(i) = S(I_e, I_i) \). A peak detection is used to find peaks and valleys in \( S_e \). Library-images between a valley and a peak are tagged ascending, images between peak and valley are tagged descending. All ascending and all descending images can then be grouped and sorted according to \( S_e \). Appending the sorted ascending to the sorted descending images yields the sorted library. With a sorted library the search for the best fitting library image can be narrowed down, e.g. by taking the time between successive monitoring images into account.

Results

The proposed method is work-in-progress and currently evaluated on data of several FUS experiments in moving phantoms and an animal experiment. We expect to considerably lower time to find the most similar baseline image as compared to search over an unsorted library.

Conclusions

The method accounts for hysteresis in the motion cycle and may increase robustness of multi baseline thermometry. It may enable the real time use of big libraries that densely sample the motion cycle.

A152

Training and learning software in the field of Focused Ultrasound therapy of the liver

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Objectives

Currently, non-invasive treatment using MR guided Focused Ultrasound (FUS) is an emerging technology. Especially in the field of moving abdominal organs there are ongoing developments to compensate for breathing motion while targeting a certain location in a liver tumor. Furthermore, up to our knowledge there is currently no product on the market which could perform such treatments. Within an EU FP7-funded project (TRANS-FUSIMO), a treatment software was developed capable of compensating for breathing motion during FUS. The presented work serves as a possibility to train the usage of the software without being dependent on MR or FUS hardware. In addition, the training and learning software enables the operator to review previously stored cases from a case database which was also implemented in the course of these developments. Furthermore, also trainees can learn from previous procedures to perform safe, efficient and effective treatments in the future.

Methods

A modified version of the TRANS-FUSIMO treatment software is presented that acts as training and learning system. Since the TRANS-FUSIMO treatment system has a modular software architecture with respect to several parts like motion compensation, MR image acquisition, and FUS beam steering, components may easily be exchanged without changing neither core functionality nor user interface of the software. For the training and learning system, MR image acquisition and FUS beam steering components were exchanged by emulating software pieces. Images are then loaded from a database instead of being acquired by a real MRI whereas FUS beam steering commands are either just logged or used for a real-time simulation of temperature. This way operators can train to use the software in a safe and effective way. If also trainees will perform such FUS treatments it is important to learn from previous procedures about, i.e., cooling effects by the vessels. A secure case database with encrypted communication has been developed and access to it is possible with the training and learning system. That enables querying of cases which can then be reviewed or trained on. This approach also enables storing data which was not processed with the TRANS-FUSIMO treatment software.

Results

A training and learning system is a necessary part when introducing new and complex software tools to be used in clinical studies. It enables the operator to work with the software before treating a patient while using the same interface and functionality like in the real world scenario. To guarantee these requirements, our software implementation is based on the same code as the actual TRANS-FUSIMO treatment software. However, only the connection to the hardware components are replaced by emulated hardware components. In the future, this training and learning system can be extended in such a way that the system itself learns from the previous procedures and provides treatment proposals for the actual cases. However, from the quality assurance point of view, there should be no direct connection between the training and learning system and the TRANS-FUSIMO treatment system.

Conclusions

In the course of the TRANS-FUSIMO project a training and learning system was developed to train the clinical personnel performing the anticipated clinical studies using a database of previously performed interventions. The main goal is to provide training and knowledge about the used TRANS-FUSIMO treatment software to ensure a safe, efficient and effective focused ultrasound treatment for the patient.

A153

Optical flow based motion tracking in the abdomen: Ultrasound vs. MRI; A direct comparison

Jeremy Tan, Cornel Zachiu, Pascal Ramaekers, Chrit Moonen, Mario Ries


Objectives

High-intensity focused ultrasound (HIFU) therapies in the abdomen usually employ motion tracking to ensure that sufficient thermal dose is delivered while minimizing collateral damage. Previous work proposed optical flow based tracking schemes using either ultrasound (US) or magnetic resonance (MR) imaging for this purpose. While MRI has thereby the advantage of superior soft tissue contrast, ultrasound imaging can achieve higher frame-rates. However, HIFU guidance with US penalizes the image quality by the requirement to perform standoff-imaging. This study directly compares the accuracy and precision of motion tracking of real-time MRI and standoff-US imaging using a hybrid imaging setup that can observe with both modalities simultaneously.

Methods

Simultaneous US-MR imaging was performed using an MR-compatible 128-element phased-array US imager installed adjacent to the HIFU transducer of a Philips Sonalleve MR-HIFU system. Two data sets were acquired: one on a phantom undergoing a known motion pattern and a second one on the abdomen of a healthy volunteer. Tracking was performed separately on the US and MR images using an optical flow algorithm. The precision and accuracy of the displacements estimated on the US data were then evaluated relative to the injected motion pattern for the phantom dataset and relative to the displacements estimated on the MR data for the in vivo case. US acquisition was performed at 4 MHz with tissue harmonic imaging while the MR Echo Planar Imaging (EPI) sequence employed the following parameters: TE = 9.0 ms, TR = 33 ms, matrix size = 176x176, FA = 25°, voxel size = 2x2x7mm³, EPI-Factor = 13. A total of 500 images were acquired for both phantom and in vivo studies. US-MR synchronization was achieved via a TTL pulse generated by the MR-scanner.
Results
Figures 160 and 161 respectively illustrate, for the healthy volunteer dataset, an example of raw and overlapped images of both modalities. Precision and accuracy evaluation of the US and MR motion estimates were restricted to a region of interest (ROI), for both the in vivo and phantom scenarios. Figure 162 displays the average displacements estimated for the phantom dataset. Relative to the injected motion pattern, an accuracy and precision of 0.71 mm and 0.32 mm respectively were achieved for the MR data, and 1.26 mm and 0.85 mm respectively for the US images. Figure 163 shows the average motion in a ROI defined on the in vivo images. Relative to the MR estimates, the accuracy and precision of the US displacements are 1.57 mm and 0.42 mm respectively.

Conclusions
This investigation validates the accuracy and precision of optical flow in standoff US imaging. Phantom and in vivo experiments show that sub-pixel performance can be achieved (<2 mm). As such, standoff US imaging is a viable alternative to MR tracking and can allow more flexibility in MR protocol design for additional scans or optimization. It can also be a powerful tool for triggering dynamic switching in MR scans. Further work includes optimization of US parameters and real-time applications of this dual modality system.

Fig. 160 (abstract A153). Sagittal images of the liver and the upper intestinal tract of a volunteer with T2*-weighted MRI on the left and tissue harmonic imaging ultrasound on the right.

Fig. 161 (abstract A153). MR (green) and US (red) images overlapped in the same coordinate system. The blue ROI outlines prominent features tracked by the optical flow algorithm.

Fig. 162 (abstract A153). Average displacement of phantom within an ROI, estimated on the MR (green) and US (red) datasets. The blue curve corresponds to the injected motion pattern.

Fig. 163 (abstract A153). Average displacement, for the healthy volunteer dataset, within an ROI (see blue outline in Fig. 161), estimated on the MR (green) and US (red) images.

A154
Trans-pulmonary FUS ablation of liver using lung flooding, aspects of lung penetration and diaphragm motion
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Objectives
Several patient studies investigated HIFU treatment of liver tumours. Its location and respiratory induced motion affects the ability to treat. The surrounding lung inhibits extracostal HIFU penetration and respiration induced motion requires compensational guiding and thermometry techniques. It has been shown that One-Lung Flooding (OLF) is safe and originates a superior acoustic path for FUS ablation of lung cancer. FUS treatment in liver could benefit from a direct trans-pulmonary HIFU path and a stable target fixation. Therefore the effects of OLF, enabling a direct transpulmonary HIFU liver ablation and reducing diaphragm motion were investigated.

Methods
One-Lung flooding was performed using a porcine model (female, 30-60 kg). Ultrasound guided HIFU was applied transthoracic, transpulmonary into liver on three animals. HIFU exposure (SU102, Sonic Concepts)
for 5 sec (3 MHz, 75 W) was sonographically targeted at each animal five times into the upper liver dome. During autopsy, examination of induced ablation zones in liver was performed as well as on lung and diaphragm tissue out of the beam path.

For diaphragm motion, MR imaging on five animals was performed during OLF under mechanical ventilation at the ex-inspirational endpoints. Images based on T2w HASTE (TE84ms/TR900ms) sequence was merged after edge detection and the diaphragmatic deviation in cranial direction was extracted.

**Results**

Fifteen of fifteen (100 %) thermal lesions were generated by transpulmonary HIFU sonication in liver. After HIFU exposure the focal zone appeared bright in B mode (Fig. 164). Macroscopically the lesions appeared elliptical with a mean length of 11.3 mm and 3.8 mm width. Histology showed no signs of diaphragm and lung tissue damage. On MRI examinations the largest diaphragm movement (35 mm) was measured in the latero-posterior diaphragm on the ventilated side. The movement wanes towards the flooded side down to 15 mm in the central area. At the lateral edge of the flooded side diaphragmatic movement becomes deterred (Fig. 165).

**Conclusions**

OLF provides a direct transpulmonary acoustic pathway for HIFU ablation in liver. Therefore the therapeutic window for transcostal FUS can be enlarged, which increases the ablation speed and enables treatment of liver tumors in the proximity to lung. Further, the reduced diaphragm and therefore liver motion simplifies targeting strategies. Alternatively pleural effusion has been reported for acoustic coupling of the upper liver dome during FUS treatment. But OLF might comprise several advantages to pleural effusion. So a lower invasivity, a position independent diaphragma location and the mentioned reduced target motion. It needs to be discussed whether the advantages for a transcostal - transpulmonary FUS ablation of liver would justify the increased procedural efforts of OLF.

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**A155**

Feasibility study of boiling histotripsy for cardiac applications using an MR-guided clinical HIFU system

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**Objectives**

Hypertrophic Obstructive Cardiomyopathy (HOCM) is characterized by abnormal thickening of the ventricular walls leading to dynamic outflow obstruction. Existing intervention techniques are invasive and present risks, such as complete heart block or arrhythmias due to scarring. Histotripsy is used to perform mechanical fractionation of tissue and has been proven effective in the creation of atrial septum defect on animals. When applied in bulk tissue, the lesions obtained are in a liquid state and can stay inside the lesion to be reabsorbed by the surrounding tissue. Thus, the creation of several, relatively small histotriptic lesions is hypothesized to potentially contribute to the reduction of myocardial volume non-invasively and with minimal scarring and risk of thrombus formation, relieving HOCM.
Methods
A boiling histotripsy feasibility study was conducted on ex vivo tissue using a clinical MR-guided HIFU system (Sonalleve V1, Philips Healthcare) to determine the optimal sonication parameters and to study the guidance and monitoring capabilities of MRI for this application. Sonifications of beef steaks and heart and pig heart were performed at 1.2 MHz using duty cycles (DC) varying from 0.3 % - 2 %, power from 200 W to 850 W and sonication duration from 20s to 70s.

Results
The MR-guided HIFU system was effective at delivering shocks sufficient to create lesions of 3 - 5 mm in diameter without causing visible thermal injury (see Fig. 166a). Optimal results were reached at 0.5 - 0.7 % DC, 50 seconds of sonication duration and power output between 550 - 750 W. MRI was proven efficient for guidance and could be used for lesion assessment for DC more than 0.5 % and power output more than 300 W (see Fig. 166b).

Conclusions
Boiling histotripsy was proven feasible in ex vivo cardiac tissue under MRI guidance and MRI monitoring. Preliminary results also showed that thermometry data could be of value in predicting lesion aspect (see Fig. 166c). Further research is needed to evaluate the feasibility of in vivo ablation through transthoracic application of HIFU.

Objectives
Pulsed focused ultrasound (pFUS) has been shown to alter molecular microenvironment of kidney and muscle and has the potential to enhance homing permeability and retention of intravenously (IV) administered mesenchymal stem cells (MSCs) to target tissue. In this study, we examined the mechanotransductive induced proteomic and histological changes following MRI-guided pFUS on rat myocardium.

Methods
Female Sprague Dawley rats (8-10 weeks old) were imaged on a 3 T clinical MR scanner (Philips) and T2-weighted MR images were acquired with 8.9 ms repetition and 4.5 ms echo time in 1 mm slice thickness. The images were used as guidance for pFUS to target heart apex (1 MHz; 0 to 8 MPa; 10 ms bursts; 1Hz PRF; RK-100, FUS Instruments). Post-pFUS changes were evaluated ex vivo using 7 T MR microimaging system (Bruker) with 7,000 ms TR and 50 ms TE in 250 μm slice thickness followed by fluorescence immunohistochemical (fIHC) analysis of albumin, Evans-blue dye (EBD) extravasation and haematoxylin and eosin (H&E). Proteomic analysis was performed using ELISA kits (Rat Cytokine 24-plex, Bio-Rad, US) and cardiac injury markers (cardiac troponin I (cTnI), pro b-type natriuretic peptide (pro-BNP) and heart fatty acid-binding proteins (hBAFP)).

Results
Targeted pFUS resulted in interstitial edema in the myocardium based on 3D reconstructed T2w MRI. Significant EBD extravasation and fIHC staining for rat albumin was observed in the left ventricular apex. H&E staining showed no morphological changes in myocardium in pFUS targeted region (Fig. 167). Temporal proteomic analysis revealed that pFUS treatment significantly elevated pro-inflammatory cytokines consistent with sterile inflammatory response with the levels returning to baseline after 24 hours post pFUS. Cardiac injury markers, cTnI, pro-BNP and hBAFP, demonstrated no significant difference compared to non-pFUS treated SHAM.

Conclusions
Noninvasive attempts to the repair of heart diseases as part of a regenerative medicine strategy has been slow. Recent studies suggest that targeted pFUS in combination with IV administered MSCs has the potential to modulate inflammatory response and improve clinical outcomes. In this study, the mechanotransductive effects of pFUS treatment triggered transient increases in pro-inflammatory and anti-inflammatory cytokines (TNF-a, IL1α, IL1β, IL10 and IL13) consistent with sterile inflammation without histological damage to myocardium or changes in cardiac injury markers. Histology reveals leakage of plasma albumin and edema into pFUS targeted myocardium in absence of microhemorrhages or muscle damage. pFUS in combination with MSCs maybe used as non-invasive technique for regenerative medicine for delivering stem cells in myocardial diseases.
A157
A retrospective simulation study of renal denervation using magnetic resonance guided focused ultrasound in a porcine model
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Objectives
Approximately 30% of adults in America are affected by hypertension. Because of the established role the sympathetic renal system has in hypertension, renal sympathetic denervation (RSD), disruption of the renal nerves found in the arterial wall and perivascular soft tissue, has emerged as a potential treatment for hypertension. The feasibility of using magnetic resonance guided focused ultrasound (MRgFUS) to non-invasively perform RSD has been demonstrated in both pre-clinical and clinical studies. A porcine animal study was recently done to test the efficacy of MRgFUS in performing RSD. The efficacy of the procedure was determined through assessing kidney medulla norepinephrine in 24 pigs. Using the MRI data obtained during these procedures, 3D models of the treated animals were created and acoustic simulations are underway to correlate the simulated pressure patterns to the experimental efficacy results and determine the effects of phase aberration.

Methods
Segmentation: For each of the 24 animals, a T1-weighted VIBE (T1w Volumetric Interpolated Breath hold Examination) image series is used to perform the segmentation (Fig. 168a). Segmentation is done using Seg3D (Fig. 168b), a volume segmentation and processing tool (Center for Integrative Biomedical Computing at the University of Utah). Twelve tissue types are segmented and a correction algorithm is used to prepare the resulting models for acoustic simulations.

Acoustic Simulations: Each tissue type is assigned an attenuation, speed of sound, and density value consistent with the literature. Using the Hybrid Angular Spectrum (HAS) method, pressure patterns for each experimental sonication treatment location are obtained. The global maximum pressure as well as the maximum pressure achieved within 5 cm of the intended focal point is identified. Additionally, a phase spread metric (PSM) is calculated to quantify the heterogeneity of each model. A PSM value of 0 indicates no phase spread while a value of 1 denotes full phase spread. For comparison, simulations are also performed for a homogenous model where all tissue types are assumed to be uniform (an average value of all 12 tissue types).

Results
At the time of submittal, simulations have been completed for 7/24 animals. Initial results demonstrate that phase aberration due to the heterogeneous anatomical structure of the pig decreased the maximum pressure achieved at the focal location by 36% when comparing the heterogeneous to artificially homogeneous segmented models. Additionally there is an average 10% increase in the PSM, indicating that the heterogeneity of the segmented models increases the phase spread. The distance between the intended focal point and maximum pressure location was also affected by the heterogeneity of the model. The mean average distance between intended focal point and maximum pressure location was found to be significantly larger (p < 0.001) when comparing the heterogenous (18.4 mm) and artificially homogeneous models (5.6 mm).

Conclusions
These initial results indicate it would be beneficial to perform phase aberration correction when performing RSD with MRgFUS in order to increase the accuracy and efficacy of the procedure.
predict treatment outcome. We present a preliminary computational study towards a better understanding of the stone fragmentation process. Specifically, we apply a novel multiphase fluid-solid coupled computational framework to investigate the interaction of a kidney stone immersed in liquid with a lithotripter-generated shock wave and a gas bubble near the stone. The main objective is to elucidate the effects of a bubble in the shock path to stone fracture.

Methods
The computational framework used in this work couples a three-dimensional (3D) finite volume two-phase computational fluid dynamics (CFD) solver with a three-dimensional finite element (FE) computational solid dynamics (CSD) solver. The surface of the stone is represented as a dynamic embedded boundary in the CFD solver. The evolution of the bubble surface is captured by solving the level set equation. The interface conditions at the surfaces of the stone and the bubble are enforced through the construction and solution of local fluid-solid and two-fluid Riemann problems. The computational analysis reported in this presentation are performed on the BlueRidge supercomputer at Virginia Tech, using 200 to 400 processor cores.

Results
The first step in this study is to verify the computational framework. To this end, we present a 1D multi-material Riemann problem, and two 3D problems on shock-stone and shock-bubble interactions. For all the three problems, our numerical solution is in close agreement with the reference (Fig. 169). Next, we present a series of 3D shock-bubble-stone coupled simulations (Figs. 170, 171 and 172).

Conclusions
Our computational analysis suggests that the dynamic response of a bubble to an LSW, as well as its impact to the stone, heavily depends on the initial size of the bubble. Bubbles smaller than a certain threshold implode within 1 s after the passage of LSW; whereas larger bubbles do not. For a specific (and realistic) setting, we find this threshold, in terms of bubble radius, to be 0.12 ± 0.02 mm. Moreover, our results suggest that a non-imploding bubble imposes a negative effect on stone fracture, as it shields part of the LSW from the stone. The effect of an imploding bubble is a bit more complex: it may promote fracture on the proximal surface of the stone, yet hinder fracture from stone interior.
A perfused breathing kidney phantom

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Objectives
One of the main challenges during FUS treatment of abdominal organs is organ motion due to respiration. If motion is ignored, the thermal dose spreads over a larger area, leading to insufficient heat for ablative necrosis. The current FUS treatment systems lack compensation of breathing motion. Research requires ex vivo moving organ models with appropriate perfusion. The work reports preparation of a ‘perfused, breathing kidney phantom’ using fresh and preserved porcine kidneys.

Methods
Fresh ex vivo porcine kidneys are prepared carefully to remove excess fat and to make the ureters, artery and vein clearly visible. The vessels are then flushed out repeatedly using saline water to remove blood. The process is then continued using ‘Thiel’ preserving fluid and then the prepared tissue is immersed in the preservation fluid for up to one week. This process will leave the tissue re-usable for months and thus making it suitable for repeated sonication and testing. The vessels in the kidneys are then attached to connectors to allow perfusion. The preserved kidneys with the connected connectors are embedded between two layers of agar phantom. The agar block with the embedded kidneys were attached to balloons to mimic breathing motion as seen in Fig. 173. Heart-lung machine is used for perfusion. The inflow from the heart-lung machine is connected to the renal artery and the outflow from the kidney is taken back to the heart-lung machine from the renal vein. Two balloons each side of the phantom are filled with water and air respectively and one of the balloons connected to the ventilator to mimic the breathing motion.

Results
The perfusion of the kidneys was imaged using x-rays (Fig. 174) and it showed very good circulation in the kidneys. The blood flow volumes can be controlled from 100 ~ 300 ml/minutes. The phantom was scanned with real time MRI under respiratory motion. The respiratory motion range along superior-inferior direction could be adjusted within 10 ~ 40 mm. Meantime, the motion speed was adjusted according to different applications as well.

Conclusions
A perusable, breathing kidney phantom was prepared using ex vivo porcine kidneys. The circulation was imaged using x-rays and the breathing motion was accomplished using the ventilator. This phantom uses Thiel preserved ex vivo kidneys accounting for its increased longevity.

Breath-hold MR-HIFU hyperthermia: Phantom and in vivo feasibility

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Objectives
The use of MR-HIFU to deliver mild hyperthermia (40-45 °C for 10-60 minutes) requires stable MR temperature mapping for long durations. While this can be achieved in the extremities, for targets in the pelvis, abdomen, chest, and neck, respiratory motion causes magnetic field variations that induce unacceptably large temperature mapping errors. Pediatric MRI examinations are routinely performed under general anesthesia with mechanical ventilation, where forced breath holds of 1-2 minutes interrupted by 20-30 seconds of hyperventilation. Our goal is to investigate the use of forced breath holds to deliver mild hyperthermia to pediatric solid tumors. In this study we evaluate the precision of MR thermometry acquired in phantoms near a moving structure and in pig back muscle near the lungs.

Methods
Experiments were performed using a 3 T MRI (Ingenia, Philips Healthcare) equipped with a clinical MR-HIFU system (Sonalleve V2, Philips Healthcare). Temperature maps were acquired over 10-20 minutes and optical temperature sensors (Neoptix T1) were embedded to validate MR thermometry. In phantom studies, breathing motion was simulated using an MRI-compatible motion platform programmed to move an aluminum block along a sinusoidal one-dimensional trajectory. Gating was simulated by halting the motion at the furthest distance away from the phantom.

In vivo experiments were performed in ventilated, anesthetized pigs lying with their back muscles above the acoustic window of the MR-HIFU system. The lungs and diaphragm were outside of the imaging field of view, but close enough to trigger the motion artifacts. Sterilized temperature sensors were embedded in the muscle adjacent to the target region. Breathing was controlled at 20-24 breaths per minute. End expiration breath holds of 40-60 seconds were implemented by switching off the ventilator, during which mild hyperthermia with a target
temperature of 42 °C was performed in an 18 mm diameter region. A recovery period of 20-40 seconds with HIFU disabled was performed after each breath hold. This cycle continued for 10 minutes hyperthermia and another 10 minutes of cooling.

Results
In phantoms next to a moving structure with a large relative susceptibility, MR temperature measurements acquired during a series of motion holds demonstrated a precision of 0.8 °C, compared to measurements acquired while the structure was moving had a precision of 1.6 °C (Fig. 175). In pig back muscle with no HIFU heating, precision during breath holds was 0.9 °C compared to 2.0 °C during ventilation (Fig. 176). MR-HIFU hyperthermia delivered during intermittent breath holds over a 10 minute duration heated the 18 mm diameter target region above 41 °C for 11.6 minutes, demonstrating the feasibility of breathhold MR-HIFU hyperthermia (Fig. 177).

Conclusions
The use of forced breath holds enables precise control of MR-HIFU hyperthermia in targets near moving structures, and may allow MR-HIFU hyperthermia to be extended to pediatric solid tumors of the pelvis, abdomen, chest, and neck.

A162
Non-invasive growth plate ablation treatment in rodent models using MR-guided High-Intensity Focused Ultrasound
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Objectives
Leg length discrepancy is an orthopaedic problem that usually appears in childhood. Without treatment, the patients will have noticeable limp, back pain and other symptoms. Traditional treatment includes surgically injuring the growth plate to shorten the longer leg. While effective, this procedure is invasive and associated with the risks of surgery. MR-guided high-intensity focused ultrasound (MR-HIFU) could provide a non-invasive approach for growth plate ablation. The hypothesis of this study was that exposure of the growth plate to HIFU would achieve a reduction in limb growth, as an alternative to surgical destruction. To test this hypothesis, we adapted a pre-clinical small animal MR-HIFU system to perform growth plate ablation treatment in rodent models.

Methods
Experiment were performed with a 3 T MRI (Ingenia, Philips Healthcare) with a preclinical HIFU system (RK 100, FUS Instruments). Growth plate ablation treatment was performed on young animals in the growth phase of their limbs (Sprague Dawley rats, male, 4-6 weeks old, n = 8). All in vivo experiments were approved by UT Southwestern Institutional Animal Care and Use Committee. Anaesthetised animals were stabilized on the HIFU platform, with one side of the thigh facing the acoustic window. The target region was selected as the growth plate of the proximal femur, with the contralateral leg acting as a control. Approximately 3 W of acoustic power was delivered to the bone for 20-30 seconds for the ablation. Temperature was measured with MR thermometry within the target region during the treatment. The animals were allowed to recover and were observed for 8 weeks. Contrast-enhanced MR images were acquired on the day of sacrifice. Femurs on both sides were collected and micro-CT images were obtained to evaluate the treatment effect on bone growth.
Results
A localized temperature increase was observed in the targeted region and appeared to cover the entire region of the growth plates (Fig. 178). All animals recovered quickly and did not exhibit any deficits in gait or function during the 8-week observation period. An average length difference of -6.4 ± 6.2 % (p = 0.02, paired t test, Fig. 179a) between treated and control femur was observed from the micro-CT images. Post-treatment MR contrast imaging (n = 8 animals) showed a signal increase of 16.7 ± 12.1 % (p < 0.01, paired t-test, Fig. 179b) in the exposed region, suggesting residual scar tissue in the region of exposure.

Conclusions
In this study, we have confirmed that growth plate ablation treatment in rodent models can be achieved with a pre-clinical MR-HIFU system. The procedure was safe and resulted in a reduction in the growth rate of the exposed leg. These results support further investigation of HIFU as a non-invasive treatment for pediatric leg length discrepancy.

Methods
MRgFUS was used to perform sonications on a tissue mimicking gel phantom (ATS Labs Inc, Bridgport, CT) in two experimental scenarios. In the first scenario, the phantom was sonicated directly without a coupling device attached to the far field of the phantom. In the second scenario, a room temperature water bag was attached to the far field of the phantom.

In both scenarios, two fiber-optic temperature probes were placed approximately 1 cm below the far field air-phantom interface, 5 cm above and distal to the planned level of sonication, with the tip of both probes at approximately the center of the phantom’s cross-sectional surface area (Fig. 180). The probes provided continuous thermal readings. A mock flat circular target of 5 cm was prescribed 4 cm from the base of the phantom on planning images. A small amount of degassed water was placed at the near field interface in both scenarios. Each phantom was sonicated a minimum of 15 times, and additional sonications were performed if necessary to produce a 10 °C temperature rise. Maximum temperature rise following each sonication was recorded.

Results
In the first scenario (no water bag coupling device), the temperature began at 19.6 °C and rose to a maximum of 32.4 °C after 15 sonications (Fig. 181). An increase of 10 °C was seen after 12 sonications. Intermittent temperature spikes were also noted, most strikingly after sonications 5, 9, and 12. In the second scenario (with water bag coupling device in the far field) the temperature began at 18.4 °C and rose to a maximum of 26 °C after 15 sonications. Additional sonications were performed and an increase of 10 °C was seen only after 18 sonications, rising to a temperature of 28.4 °C. No temperature spikes were seen in the second scenario with a relatively smooth/gradual increase in temperature noted.

Conclusions
This experiment demonstrates that a water bag coupling device in the far field significantly delays the rate of heating and maximum heating in this region during MRgFUS. These results suggest that this type of device can both allow for more complete treatments and help minimize risk of far field skin injury.
A164
Low Intensity Pulsed Ultrasound (LIPUS) for the treatment of spinal disc degeneration — ultrasound exposimetry system and in vivo implementation in a rat tail model

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Objectives
Discogenic back pain presents a major health issue, with current therapeutic interventions limited to short term symptom relief without providing regenerative remedies for diseased intervertebral discs (IVD). LIPUS or non-thermal pulsed focused ultrasound (p-FUS) offers a potential treatment option, where ultrasound mechanical energy targeted to a damaged IVD may produce a favorable biological response to modify, stall and even reverse IVD degeneration in a non-invasive manner. LIPUS has been shown clinically to be an effective noninvasive method for the stimulation of bioactivity and favorable tissue remodeling, such as wound and fracture healing. Recent in vitro studies have shown that LIPUS stimulation can induce desirable effects within cultured intervertebral disc cells such as an increase in extracellular matrix metabolism, upregulation of proteoglycan synthesis, stimulated collagen synthesis, and induction of anti-inflammatory factors. The objective of our “work in progress” presented herein is to develop a LIPUS exposimetry system specific to targeting degenerated tissue.

Methods
Ultrasound exposimetry systems were devised and fabricated specific to delivering LIPUS to damaged caudal discs in the rat tail (Fig. 182). The two configurations consisted of 2.5 cm diameter spherically focused PZT4 transducers (1.0 MHz, f = 1; 1.6 MHz, f = 3.8), integrated within a water-filled 3D printed plastic housing with a mylar window w/ cross-hairs, designed as an acoustic standoff to place the focus within the targeted rat tail disc. The applicators/apparatus were evaluated with comparative beam plot measurements with/without insertion of sectioned rat tails ex vivo (Fig. 183a, b). In vivo studies using the disc damage model are performed following standard procedures approved by UCSF IACUC, with stab incisions applied within 16 Sprague-Dawley rats to generate damage/inflammation in tail discs. Five daily p-FUS exposures (ISP2A120 mW cm-2) are to be applied to stab discs and compared to stab only and normal controls. Histology and microarray gene analysis are performed at 5 days and 28 days after injury, also assessing changes to cartilage, fibrous tissue, and bone.

Results
Intensity measurements and radiation force observations with and without the tail sections (Fig. 182c, 183c) demonstrate the focused systems can deliver LIPUS (at 30-50 % of mylar surface intensity) through the skin and connective tissue, to the center of the narrow ~1.5 mm x 5 mm rat tail disc, while reducing exposure to significant portions of the adjacent bone and negligible temperature elevation. The LIPUS exposimetry to damaged rat tail discs, and subsequent histological and microarray analysis are currently ongoing and will be presented.

Conclusions
This study demonstrated the technical feasibility of delivering ultrasound isolated to a targeted rat tail disc for studies of LIPUS exposure. In summary, this LIPUS exposimetry apparatus and disc damage model can be implemented for further study in vivo to demonstrate potential of ultrasound to increase cellularity, reduce inflammation, and improve remodeling to acute or degenerative disc-related back injury.

A165
Experimental evaluation of MR-guided HIFU for ablation of sacroiliac joint using a model of human sacrum

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Objectives
Chronic low back pain (CLBP) is a leading cause of work disability. In 15-30 % of CLBP patients, the source is the dysfunction of the Sacroiliac Joint (SIJ). [1] One method providing therapeutic effectiveness is radiofrequency ablation (RFA), however this invasive and laborious procedure has shown slow clinical adoption, limiting effective treatments for SIJ pain.[2-3] HIFU could provide a non-invasive alternative by creating a continuous lesion lateral to the sacral foramina (Fig. 184). The main risk of using HIFU for SI joint ablation is damaging adjacent neural structures. A pilot study in swine demonstrated that MR-guided HIFU (MRgHIFU) ablation of SIJ is feasible, and lateral branch nerves can be ablated without damaging adjacent nerve roots. [4] The anatomy of human sacral foramina, however, is not identical to that of swine (Fig. 185). Hence, translating preclinical safety results into human patients may be prone to risk and further evaluation is needed.

Methods
Phantom: Images from an adult pelvic CT scan will be used to segment the sacrum bone from adjacent muscle tissue. The segmented model will be manufactured in a 3-D printer using (Acrylonitrile Butadiene Styrene) ABS plastic.[5] The muscle tissue will be mimicked by a high-temperature hydrogel matrix (gellan gum) combined with different sizes of aluminum oxide particles and other chemicals.[6] Experiment: MR-HIFU experiment will be performed using a clinical HIFU system (InSightec, Ltd., Israel) installed in a 1.5 Tesla MRI scanner (GE Healthcare, USA). HIFU beam will be applied as shown in Fig. 184. Temperature will be monitored using MR thermometry and four fiber-optic temperature sensors (Luxtron, USA) inserted into a foramen.

Results
For L5 to S3 vertebral levels, we will determine the range of the acoustic parameters (focal spot size and position) and HIFU beam...
orientations within which thermal ablation of the SJ can be achieved without causing thermal damage inside the sacral foramina.

Conclusions
Detailed investigation of the safety of HIFU of SJ using human-sacrum-mimicking phantom will inform future studies aimed at applying MR-HIFU clinically to treat sacroiliac low back pain.

References

Methods
A 3D spiral UTE sequence was used for rapid UTE imaging. Briefly, a 3D stack-of-spirals acquisition was modified to achieve very short echo times by beginning each spiral readout immediately after the through-plane phase-encoding gradient waveform is complete. This results in a variable TE in the through-plane direction. For the center of k-space where the PE gradients are small (or nonexistent), the minimum TE achievable is approximately 50 μs (Fig. 186). A room-temperature bone sample was first placed into the scanner, and the spiral UTE sequence was used to acquire volumetric images every 75 seconds for 12.6 minutes. Following this acquisition, the bone was placed into a 55 °C water bath for 5 minutes, then imaged again with the same protocol. After imaging, cortical bone was manually segmented and the mean signal intensity recorded at each time point.

Objectives
Currently, there is no feasible way to measure bone heating in vivo. Tissue close to bone can be monitored and temperature models can be used to determine when it is safe to apply another burst of ultrasound energy. However, a noninvasive MR-based bone thermometry method would be useful to directly monitor bone heating during FUS treatment. Because of its short T2/T2*, the popular PRF shift thermometry technique cannot be performed in bone. A method proposed by Miller and further demonstrated by Han uses UTE imaging to detect signal changes in bone caused by changes in the relaxation rates due to heating. Here, we have used a 3D spiral-based UTE sequence to rapidly collect UTE images and detect signal changes in bone as it cools down inside the bore of the scanner. Furthermore, we have used this sequence in a proof-of-concept UTE skull imaging experiment using a human volunteer.

Results
In Fig. 187, the early and late TE images from the bovine tibia experiment are shown, along with the subtraction image of the two. Cortical bone signal is only visualized with the early TE configuration. Figure 188 shows the mean signal of the heated sample increasing as the bone cools in the bore of the magnet over a 12-minute time-span. In contrast, the sample that remained at room temperature shows no change in signal. Figure 189 shows whole-head spiral UTE images alongside late-echo images to illustrate the difference in contrast achievable with this sequence. Direct subtractions as well as scaled subtractions are shown, highlighting the bone signal. SNR, measured in a region of the frontal bone, is 54 in the minimum-TE image, 23 in the direct-subtraction image, and 43 in the scaled-subtraction image.

Conclusions
This work shows the feasibility of temperature monitoring for bone across potentially large fields of view; however, the small phantom size and number of experiments presented here call for further study. Volumetric UTE sequences have typically required scan times of about 10 minutes, which may preclude their use in focused ultrasound procedures; by comparison, the 3D spiral UTE sequence can acquire large volumetric data in about one minute. The ultimate goal is to monitor the entire skull and skull base between therapeutic insonifications, which are typically separated by a few minutes.
A versatile ultrasound system for in vitro experiments on cells
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Objectives
One of the most difficult tasks to achieve with available instrumentation used to study the interaction between ultrasound (US) and cellular model systems is to design an experiment, where the effects of only one physical parameter is evaluated, while all the others are kept constant. The set-ups are usually custom made, often use instruments intended for a different purpose. The outcome are results not always well reproducible, and rarely comparable with those obtained with techniques considered standard in molecular and cellular biology. Sterility, as well as temperature, are not well controlled. All these considerations translate into poor reproducibility of US in vitro data, leading to standard deviations which are often larger than the values themselves. We present US experiments performed on cell samples with our designed bench-top US apparatus versatile to be adapted for several in vitro experiments and show easy and robust reproducibility and low Standard Deviations data, using standard set-ups for the cell samples.

Methods
One main feature of our bench-top US system is that it has been designed in order to minimize interferences and degradation of the acoustic energy using the standard plasticware that are commonly used in molecular biology labs, ensure temperature controls and sterility conditions needed in the field. We present a set-up where the simultaneous use of a set of transducers operating at different frequencies on the same plate, allows the comparison of the deposition of the same acoustic pressure, while evaluating the effect of frequency alone on the readout of the cell experiments. The apparatus modular design allows also to use a set of transducers operating at the same frequency, in experiments where the throughput is a relevant factor. We show that it is possible to define the position of the target within all the achievable areas of the acoustic field with sub-millimetric accuracy.
Tests for several applications based on biologic effects by ultrasound have been carried out by varying the acoustic parameters such as the power, the frequency range, the sonication time and the duty cycle, all controlled within robust protocols executed in automation.

**Results**

We show data that demonstrate that it is possible to perform in vitro experiments for different purpose (i.e. drug or gene delivery via cellular sonoporation, tissue regeneration, neuronal cell stimulation, bacterial growth etc.) keeping constant the relevant physical parameters of sonication, for instance acoustic pressure, but varying one at the time the others parameters (i.e. frequency, pulse length or duty cycle etc.).

**Conclusions**

We show that with our apparatus it is possible to obtain robust and reproducible results on experimental ultrasounds using all the standard devices that are commonly available in biological labs. The improvement on the side of reproducibility and portability of the experiments allows a straightforward comparison between our results and those obtained with other techniques.

**A168**

**Stability of a low-cost thermochromic quality assurance phantom for FUS devices**

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**Objectives**

A simple phantom was designed, constructed and tested for daily quality assurance (DQA) of ultrasound therapy transducers. Regular checking of the therapeutic transducer is indeed a key safety test as differences in power output can either result in harm to the patient or ineffective treatment. We introduced recently a DQA phantom which allows the user to determine if the power output of a transducer has changed after calibration through visual inspection. We present here repeated experiments over 6 months on a thermo-chromic material-based phantom, with two identical transducers, in order to investigate the repeatability and the stability of the DQA phantom. The phantom is made of an attenuating ultrasound absorber (AptFlex F28, Precision Acoustics Ltd, Dorchester, UK) with a surface layer of thermochromic liquid crystals (TLCs) (R35SCB, LCR Hallcrest LLC, Glenview, USA). The TLC changes color when temperature exceeds 35 °C.

**Methods**

Two phantoms made from the same sample of AptFlex F28 were tested respectively with two high-power 1.1 MHz transducers (Sonic Concepts, Bothell, USA). A waveform generator (DG1022, Rigol, Beaverton, USA) supplied a 1.1 MHz sinusoid to a power amplifier (100A100, Amplifier Research Corp., Souderton, PA, USA). The amplified signal was monitored by an oscilloscope (TDS2001C, Tektronix, Beaverton, OR, USA) and connected to the input of a Sonic Concepts matching network and ultrasound transducer. Measurements of the time until first TLC material color change were recorded to across a 0.4-11 W range of power in ten sessions spanning over a six-month period.

In addition, the pressure field of the transducers was measured in a water tank (AIMS III, Onda Corp., Sunnyvale, CA, USA) and used as input to a time-domain finite difference simulation of the bio heat equation to simulate the thermal rise in the absorbing material. Simulations were performed with an in house Matlab code (Mathworks, Natwick, USA) operated on a laptop. Table 7 provides material properties used in the simulation. The time required to achieve a color change temperature of 35 °C is recorded, and the simulation is run across electrical intensities ranging between 0.5 and 12 W, based on 85 % efficiency.

**Results**

Transverse and longitudinal maps of the pressure fields are displayed on Fig. 190. All measurements of time to see a thermal change on both DQA phantoms are plotted in Fig. 191 as a function of electrical power, as well as the corresponding simulated curve. Good agreement is observed between simulation and experimental results. For optimal sensitivity, it is recommended to select an electrical power of 1.5 – 2.0 W. Here, DQA assessments are complete within 20 seconds, and the typical user (with reaction time of 0.5 s) can detect a 10 % deviation in acoustic power output.

**Conclusions**

Long term (6-month) testing of two DQA phantoms on two identical transducers proved to be stable enough to compile all measurements on a single graph (Fig. 190). This system is a cheap and effective way to produce DQA phantoms, both because of the ease of use, and because the use of TLC makes visual changes in the phantom reversible and reproducible over a long period of time. Next steps involve refining the simulation methods used to better validate the DQA design and deploying custom-mounted DQA phantoms to end users and iterate upon the design based on their scenario-specific experiences.

**Table 7 (abstract A168). Simulation material properties**

| Density (kg/m3) | 1010 | 1050 | 3800 | 1.21 |
| Speed of sound (m/s) | 1500 | 1500 | 9900 | 3.30 |
| Attenuation (Np/m) | 3.30 | 2.5x10^-3 | 2.5x10^-3 | 1.89 |
| Specific heat (J/kg*K) | 1800 | 4200 | 775 | 0.001 |
| Thermal diffusivity (mm2/s) | 0.091 | 0.086 | 0.13 | 0.21 |
| Thermal conductivity (W/m*K) | 0.165 | 0.38 | 39 | 0.026 |

**Fig. 190 (abstract A168).** Measured pressure in x-z plane and computed x-y plane pressure

**Fig. 191 (abstract A168).** Time required for observable DQA color change at different driving electrical power levels
A169 Quantification of cavitation histotripsy lesion volumes on a clinical MR-HIFU system

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Objective
High intensity focused ultrasound (HIFU) is a non-invasive therapeutic technique currently used in the clinic to thermally ablate tissues. However, there exists a limitation for significant collateral tissue damage and/or incomplete treatment. Alternate HIFU regimes such as cavitation histotripsy (CH) utilize microsecond-long pulses, low duty cycles, and high acoustic pressures to mechanically fractionate tissue. This approach could help disrupt tissues while sparing nearby critical structures from thermal effects and increase the clinical applicability of HIFU.

Method
To understand the effect of CH parameters on lesion volumes, we explored a wide range of sonication parameters, such as acoustic power, total sonication time, number of cycles/pulse (pulse width), and pulse repetition frequency (PRF). The goal of this work was to quantify lesion volumes in tissue-mimicking phantoms for various CH sonication parameters.

Result
CH sonifications produced varying lesion volumes in the tissue-mimicking phantoms. Increasing the peak acoustic power from 500 to 800 W linearly increased the lesion volumes from 790 to 2780 mm3 (R2 = 0.83). On the other hand, 300 W did not produce a visible lesion and 400 W produced a lesion that was visible to the naked eye, but not spatially resolvable using a 6 MHz ultrasound imaging transducer. A linear increase in lesion volume from 600 to 2711 mm3 (R2 = 0.76) was also observed for the tested range of number of cycles/pulse. When increasing total sonication time from 271 to 1630 s, lesion volume increased from 1392 to 1500 mm3. Varying PRF from 10 to 60 Hz also increased lesion volumes (1688-2182 mm3). Both 0.17 and 0.33 % duty cycle sonifications produced small lesion volumes (<0.5 cm3), while sonifications at higher duty cycle produced lesion volumes greater than 2 cm3.

Conclusion
Cavitation histotripsy was performed on a clinical MR-HIFU system and resulting lesion volumes in tissue-mimicking phantoms were quantified across a broad range of sonication parameters. Results indicate that acoustic power and number of cycles/pulse strongly affect lesion volume, while total sonication time and PRF do not. This study informs future CH work in tissue-mimicking phantoms as well as ex vivo and in vivo studies. The use of a commercial, clinical HIFU system may speed up clinical translation of histotripsy compared to custom-built preclinical HIFU systems.

A170 Tissue-mimicking thermochromic rib phantom for characterization of intercostal HIFU sonifications and beam-shaping methods

Navid Farr1, Satya V.V.N. Kothapalli2, Avinash Eranki3, Ayele Negussie1, Emmanuel Wilson1, Reza Seifabadi1, Peter Kim1, Hong Chen2, Bradford Wood1, Ari Partanen3
1National Institutes of Health, Bethesda, Maryland, USA; 2Washington University in Saint Louis, Saint Louis, Missouri, USA; 3Children’s National Health System, Washington, DC, USA

Objective
High intensity focused ultrasound (HIFU) is a non-invasive therapy modality, offering a unique method of tumor thermoablation. HIFU treatment of hepatic tumors requires intercostal transmission of sufficient energy while minimizing rib and adjacent tissue heating. Assessment and experimental validation of beam-shaping methods and numerical models may aid in clinical translation of HIFU for hepatic tumor therapy. The objective of this study was to develop and produce a tissue-mimicking thermochromic phantom (TMTCP) with embedded rib-mimics to characterize sonifications and evaluate beam-shaping methods for transcostal HIFU therapy.

Methods
3D-printable ribcage was based on rib renderings from computed tomography. Rib-mimics were manufactured using 3D-printed UV-curing photopolymer with acoustic properties comparable to human ribs. Phantoms were produced by fixing a rib-mimic within a plastic container and then filling it with TMTCP gel. Before HIFU, rib-mimic was visualized on T2-weighted MRI and manually segmented. During sonication, MR images were acquired and temperature was calculated using the PRFS method. Beam-shaping was performed by projecting the ribs onto the transducer surface by ray tracing from the focal point. Transducer elements shadowed by ribs were disabled, while power for active elements was compensated accordingly. Locations 3 cm behind the ribs were targeted with all elements enabled or by using beam-shaping. Temperatures within the targeted region and around rib-mimic were assessed. Phantoms were sonicated by electronic steering (target = 8 mm) at 1.2 MHz frequency and 80 W power for 60 s under MRI guidance using a clinical MR-HIFU system (Sonalleve V2, Philips, Finland). Post-HIFU, TMTCP enabled evaluation of temperature distribution within the targeted location and at the rib-mimic. Moreover, acoustic pressure measurements were performed in water by scanning a hydrophone within: 1) focal plane behind rib-mimics, 2) same location with beam-shaping, and 3) plane at the rib-mimic level.

Result
TMTCP material with embedded rib mimics was developed and produced. Real-time MRI temperature monitoring at the targeted location and around the rib-mimic was successful. Qualitative and quantitative assessment of temperature was also obtained by phantom color change. Temperatures over the rib-mimics were reduced using beam-shaping, i.e., selectively disabling transducer elements whilst maintaining equivalent acoustic power. Acoustic pressure fields were evaluated using a hydrophone (results pending).

Conclusion
TMTCP rib phantom was developed and applied in characterization of intercostal HIFU sonifications. The utility of beam-shaping techniques to minimize temperature rise on the ribs while maintaining focus quality during transcostal sonifications was demonstrated by monitoring heating patterns. This phantom may be useful in characterization and validation of intercostal HIFU sonication methods and numerical models.
A171

Multifunctional theranostic contrast agent for ultrasound and photoacoustic image guided tumor therapy with Focused Ultrasound triggered local delivery

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Objectives
Not released for publication

Methods
Not released for publication

Results
Not released for publication

Conclusions
Not released for publication

A172

Low intensity ultrasound and microbubbles increase cellular cisplatin and enhances cisplatin effect in a bladder cancer mimicking culture model

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Objectives
Multi-modality treatment for muscle invasive bladder cancer aims selective organ preservation with favorable outcome. Cisplatin-based chemotherapy has a survival benefit in combination with radiotherapy. Local delivery of cisplatin may improve local disease control and survival rates. Ultrasound triggered microbubble cavitation enhances chemotherapeutic agents including cisplatin both in vitro and in vivo. However, it is not fully elucidated whether the enhancement is related to cisplatin concentration in tumor cells. The purpose of this work is to clarify the relation between enhanced cisplatin effect and cellular cisplatin concentration in a 3D culture system.

Methods
Human urinary bladder cancer cells were mixed with collagen type-I and were seeded into a culture dish at the volume of 50, 100, and 200 microl (Fig. 192). The gel mixture was exposed to non-focused ultrasound (center frequency 1 MHz, duty factor 50 %, intensity 0.9 mW/cm2, time 1 min) in the presence of cisplatin and microbubbles. For platinum measurement, cells were harvested from the gel right after the sonication and were separated from the dissolved gel. Cisplatin concentration was determined by measuring ICP-MS with an inductively coupled plasma mass spectrometer. For assessing cell viability, cells were culture for 4 days in the gel after the sonication; thereafter cells were harvested and counted manually.

Results
The combination of ultrasound and microbubbles increased the cellular platinum concentration at all tested gel volume. Ultrasound alone did not significantly increase the cellular platinum concentration. Sonication did not increase the platinum concentration of the dissolved gel (i.e. without cells). The platinum concentration of the dissolved gel increased with the gel volume. The enhanced cytotoxic effect was observed at the gel volumes of 50 and 100 microl.

Conclusions
This study shows the feasibility of our culture model for assessing the relation of the intracellular delivery and the cytotoxic effect. With our ultrasound parameter, ultrasound-triggered microbubble cavitation has a potential to deliver cisplatin to cancer cells. Meanwhile, the results of this study suggest cisplatin in the extracellular matrix gradually enters cells and shows the cytotoxic effect even without ultrasound. Because cisplatin is a small molecular hydrophilic drug, it rapidly diffuses and may remain in the collagen. The thicker gel retained the higher platinum concentration. Further studies on ultrasound parameter may improve efficiency of the intracellular delivery and contribute to the local delivery of cisplatin in bladder cancer treatment.
the algorithm, using the model in its current implementation, will have a tendency to under-treat rather than to cause immediate thermal damage. It also reveals that the algorithm would benefit from a refined mechanism for online-adjustment of the heating estimation.

Conclusions
We have implemented the basic functionalities of a MPC-based algorithm for hyperthermia using HIFU. Given a treatment region and a safety margin, it computes a HIFU treatment sequence which meets the minimum required treatment temperature while respecting a given maximum temperature to avoid thermal damage. Using the model in its current implementation, the algorithm will likely have a tendency to under-treat rather than causing thermal damage as temperature estimation errors are more extreme in over-estimations. We have yet to prove the viability of the algorithm in an experiment which uses the algorithm to steer the transducer. The good agreement between a real temperature development and the used model's prediction, however, gives us reason to believe that we will be able to present the working prototype of an MPC-based HIFU hyperthermia system in the near future.

A174
Image-based spatial HIFU transducer calibration for MRgFUS
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Objectives
In MRgFUS applications with movable transducers it is a common problem to establish a connection between scanner- and transducer-coordinate systems (spatial calibration). Vendor calibration tools are often only available in proprietary software and not accessible in research applications. Thus, transducer position and orientation are often prescribed in a tedious and error prone manual process. The current work presents an image-based spatial transducer calibration method that transforms a transducer model automatically into a given setup. Our method is based on the idea, that the water systems tubes are typically well visible in MR imaging and attached to one side of the transducer, giving information on the transducers position and orientation. An MR image of the transducer and its periphery can thus be used for an automatic calibration method.

Methods
Inputs for the method are an MR image of the transducer in a given setup and a transducer model. A transducer model consists of all relevant transducer information (such as e.g. transducer center, transducer orientation, or relative element positions) and a corresponding MR image. Note that this transducer model has to be established only once and can afterwards be used on any given setup. Both model- and setup-image should contain the transducer and its direct periphery, slice orientation should be roughly parallel to the transducer surface.

The transducer calibration is implemented as two-step process: An initialization is used to coarsely align setup and model image, a subsequent image registration aligns the images precisely. The accumulated transformation can then be used to map from transducer to MR coordinates and vice versa.

The initialization is a series of image registrations that subsequently align transducer to skin coupling, transducer center, and cable orientation.
The initial estimate is used as a starting point for an NCC-based, rigid image registration.

**Results**

After setting up a transducer model once, the proposed method can be used to automatically determine transducer position and orientation from a single MRI scan within 2-4 seconds. The method was used in several validation experiments as a suggestion to the user. In most cases the proposed calibration was accepted, only a few times minor adjustments were necessary. We currently evaluate the method on the basis of manual ground-truth and with different transducer models.

**Conclusions**

A hardware- and vendor-independent, image-based transducer calibration may be useful in a variety of research applications. Our method was implemented with publicly available modules in MeVisLab and can be provided for research purposes.

**A175**

Functionalised human Thiel embalmed cadavers – a clinical model for focused ultrasound development & testing

Helen McLeod
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**Journal of Therapeutic Ultrasound 2016, 4(Suppl 1):A175**

**Objectives**

Human Thiel embalmed cadavers have a number of advantages over traditional formalin embalmed cadavers. Properties such as flexibility, tissue tone and a patent vascular system provide the basis for an anatomically accurate model. Researchers at University of Dundee have developed techniques to provide extracorporeal perfusion to the cadaver vascular system, facilitating multimodal imaging techniques such as CT, Xray, Ultrasound and MR. Model complexity is further improved by the addition of respiratory motion, produced by mechanical ventilation. Applications for perfused cadaveric models in the development and validation of Focused Ultrasound Technologies have been demonstrated.

**Methods**

Input and output (if required) connections are placed into the cadaveric target vessels and secured. Connections are made to a heart bypass machine to provide extracorporeal flow into the vessels, with flow conditions input to mimic the native flow expected. The fluid selected during perfusion is dependent upon the imaging modality, for example vascular ultrasound requires contrast therefore, a specific blood mimic solution is used. X ray, CT and MRI can successfully use a variety of fluids, with contrast being delivered to target vessel as bolus dose, mimicking clinical practice. Respiratory motion is created by intubating the cadaver with ET tube and ventilating with a mechanical ventilator.

**Results**

Exemplar images from all imaging modalities demonstrate the application of multi modal imaging techniques within Thiel embalmed human cadavers. The perfused cadaveric model has been used within the preclinical development and validation of Focused Ultrasound therapies targeting liver and kidneys.

**Conclusions**

The perfused Thiel embalmed human cadaver can be imaged in multiple modalities, CT, fluoroscopy, angiography, MRI and Ultrasound. It offers a robust and versatile model, ideal for development and evaluation of advanced technologies such as focused ultrasound therapy in moving organs.

**A176**

Magnetic resonance guided high intensity focussed ultrasound ergonomic animal experiment platform

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**Objectives**

Magnetic Resonance Guided High Intensity Focused Ultrasound (MR-HIFU) is a minimally-invasive treatment modality that is gaining clinical use. It is used to non-invasively ablate tissue or disrupt the Blood-Brain Barrier. Translation of this technique can be accelerated by facilitating animal experiments, and the development of ergonomic platforms can achieve this purpose. MR-HIFU exposure is performed under water requiring unique engineering solutions to waterproof and tune RF coils. Custom radiofrequency (RF) coils are necessary to obtain optimal signal-to-noise ratio (SNR) in this setting. Without careful tuning and matching of RF coils, targeting, temperature monitoring and thermal dose assessment may not be accurate enough.

The purpose of this study is to provide an engineering solution to treat mice using MR-HIFU using custom RF coils submerged in water with sufficient SNR that allowed for accurate thermal mapping.

**Methods**

An MR-HIFU animal treatment system was designed so that:

- SNR was increased compared to commercial coils
- Tissues were acoustically coupled while the animal remained appropriately anesthetized and warm
- HIFU beam could pass-through window to target

The developed system was compared to the Small Flex Coil (Philips, USA). An ex vivo mouse was imaged to compare systems. Two parameters were used to determine quality of new system: SNR of targeting images and temperature stability from thermal mapping. The system was tested at two different locations, TBRHSC (Thunder Bay, Ontario, Canada) and Sickkids (Toronto, Ontario, Canada) to confirm reproducibility.

**Results**

An ergonomic treatment bed was designed with a waterproofed coil embedded. After first tuning, RF coils experienced significant loading from the surrounding water, with a frequency shift as high as 7.3 MHz when submerged in water. A tunable matching circuit was added on the circuit to achieve further fine tuning.

Figure 196 shows a comparison between the Flex coil and the developed coil. SNR of the in-house developed coil was 28 and 23.5 compared to an SNR of 6 from the Small Flex coil during TBRHSC and Sickkids tests respectively. (Gradient Echo, TE = 4 ms, TR = 8 ms, FOV = 50 mm x 50 mm, 2.5 mm slice thickness NEX = 4, FA = 10o).

Figure 197 shows a 1D thermal calculation over time for both coils. The custom coil showed a stable temperature where the Flex coil recorded a decrease over time for the temperature.

Figure 198 shows the device used to ergonomically hold mice for focused ultrasound exposure.

**Conclusions**

We successfully designed an ergonomic treatment bed designed for HIFU exposure that can be used underwater while providing sufficient SNR for targeting and thermal mapping. This treatment bed will further facilitate research for focused ultrasound small animal experiments.
A177
Numerical and experimental validation of a novel therapeutic phased array transducer aimed at sparing of the near-field: the Voronoi-tessellated Fermat’s Spiral
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1University Medical Center – Utrecht, Utrecht, Netherlands; 2IMASONIC SAS, Voray sur l’Ognon, France

Objectives
Magnetic resonance-guided focused ultrasound (MRgFUS) has repeatedly been demonstrated as a safe, non-invasive therapy for uterine fibroids. However, the treatment response of fibroids with high T2-weighted signal intensity (T2wSI) on pretreatment MRI is poor. While blood flow and cellularity have been hypothesized to play a role, there is a lack of research investigating the cause of poor outcomes in high T2wSI fibroids. This abstract presents an innovative excised, perfused human uterine fibroid model designed to address this clinical challenge. Implementation of this model will allow for characterization of the MR, physiological, thermal, and acoustic properties of uterine fibroids and clarify the reason for variable efficacy in MRgFUS fibroid treatments.

Methods
In this study, women undergoing hysterectomy due to symptomatic fibroids receive a preoperative MRI (Siemens 3 T Trio). Immediately following surgical removal, both uterine arteries are cannulated, and the specimen is perfused and immersed in a Modified Krebs Henseleit Buffer maintained at a temperature of 37 °C. Tissue pH, temperature, and interstitial pressure are monitored throughout. This setup keeps tissue viable up to 8 hours post surgery, during which time, the preoperative MRI protocol is repeated on the ex vivo uterus. Comparing in vivo and ex vivo images reveals how/if MR tissue properties and organ perfusion have changed. Additional property measurements include thermal diffusivity and conductivity, acoustic properties of density, speed of sound, and attenuation, and histological examination of vessel density, cellularity, and fibrosis.

Results
Figure 199a shows an in vivo coronal MRI identifying two fibroids with hypointense T2wSI. The ex vivo model (Fig. 199b) includes heated water baths and fiberoptic probes for temperature control, perfusion-line manometers, and bubble traps; it is MR-compatible and integrated with the MRgFUS system. Photos of the excised uterus before and after perfusion (Fig. 199c) demonstrate effective clearing of blood. While the fibroids’ T2 values increase slightly ex vivo (Fig. 199d, TrueFISP MRI), they remain hypointense relative to the myometrium.

Figure 199e shows an ex vivo T1 TWIST MRI where contrast agent does not extravasate into the fibroids, indicating the low vascularity of these fibroids. Thermal diffusivity and thermal conductivity of the fibroid tissue were 0.173 mm²/s and 0.431 W/m/°C, respectively. Participant recruitment and data collection are ongoing.

Conclusions
This unique setup for the characterization of fibroid properties has great potential for explaining the variable treatment response of fibroids, for improving MRgFUS planning, monitoring, and control, and for identifying the patient population most likely to benefit from MRgFUS therapy.

A178
A unique tissue preparation for characterizing uterine fibroid tissue properties for MRgFUS thermal therapies
Christopher Dillon, Margit Janát-Amsbury, Allison Payne
University of Utah, Salt Lake City, Utah, USA

Objectives
Not released for publication

Methods
Not released for publication

Results
Not released for publication

Conclusions
Not released for publication

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Results
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Conclusions
Not released for publication
A179
Acoustically transparent surface coil arrays for MR guided HIFU
Joseph Corea1, Patrick Peiyong Ye1, Ana Claudia Arias1, Kim Butts Pauly2, Michael Lustig1
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Objectives
In MR imaging, a surface coil array is most sensitive to tissue close to it. So for MR guided HIFU, placing the array between the transducer and the patient gives the highest signal-to-noise ratio (SNR), and in turn the best quality images to track the heating. During therapy, the transducer is moved in the water bath, which would pass acoustic energy directly through different parts of an array between patient and transducer. Ultrasonic energy easily scatters and attenuates in the materials commonly used in array construction (Fig. 200a). As a result, lower SNR body coil in the center of the image where heating occurred. Furthermore it showed up to 5 times the SNR at the surface of the phantom (Fig. 201). The array displayed 84% transmission of acoustic energy and no measureable beam distortion, allowing it to be arbitrarily placed in front of the transducer (Fig. 202).

The 4-channel array was easily able to track the heating inside the skull phantom, imparting no significant attenuation or distortion (Fig. 203).

Conclusions
Utilizing the materials and processes described here it is possible to create high SNR arrays that minimally interact with heating in MR guided HIFU, giving clinicians excellent images to guide therapy.

Fig. 200 (abstract A179). A) Acoustic energy measured at the focal point with (ii) no obstruction, (iii) typical tuning capacitor and conductor, and (iv) thin printed tuning capacitor. B) Fabrication of printed coil. C) Photograph of acoustically transparent array.

Fig. 201 (abstract A179). A Printed array wrapped around gel phantom submerged in water inside head transducer. B) Scan of phantom with (i) body coil and (ii) 4-channel array. C) SNR comparison of printed array and body coil.

A180
Multi-echo pseudo-golden angle stack of stars thermometry with high spatial and temporal resolution
Bryant Svedin, Allison Payne, Dennis Parker
University of Utah, Salt Lake City, Utah, USA

Objectives
Radial acquisitions offer several unique advantages for proton resonance frequency (PRF) shift thermometry for guiding focused ultrasound therapy in the breast. Frequently sampling the k-space center provides motion robust images, as well as the ability to correct for respiration induced off resonance. It has been shown that arbitrary high spatial and temporal resolution can be achieved in dynamic

Fig. 202 (abstract A179). A Scan of phantom on a fibroid transducer. B) Heating point inside of phantom (i) without and (ii) with the array present. C) Phantom inside head transducer. D) i) Coronal and (ii) axial scans of heating point in phantom tracked with array.

Fig. 203. (abstract A179). A Location of array wrapped around 3D printed skull containing bovine brain/gel submerged in water in head transducer. B) Sagittal of brain phantom showing heating tracked with array. C) Axial scan of skull and brain phantom from 4-channel array.
MRI by acquiring successive radial spokes separated by the golden angle and applying a sliding k-space weighted image contrast (KWIC) filter, to the reconstruction. This work investigates a pseudo-golden angle 3D multi-echo stack of stars acquisition to simultaneously measure PRF shift temperature, T2* and ρ (signal magnitude at TE = 0), correct respiration induced off resonance, and provide water/fat separation with high spatial and temporal resolution.

Methods
A 3D multi-echo stack of stars spoiled gradient echo (GRE) sequence was modified to use a pseudo-golden angle increment to repeat the k-space trajectory after 377 views, ~137.56°. Experiments were performed in a normal breast volunteer on a Siemens 3 T Trio scanner to assess the effectiveness of this sequence and reconstruction technique in coronal and sagittal slabs (1.3x1.3x3mm, FOV = 166 mm, Matrix Size = 128x128x8, Flip Angle = 10°, TR = 20 ms, 13 Echoes, TE's = 2.46/3.69/4.92/6.15/7.38/8.61/8.84/11.07/12.3/13.53/14.76/15.99/17.22 ms). Using the same imaging parameters and Ambu bag movement to simulate respiration, a pork phantom was heated with FUS. Reconstruction: After accurately locating the k-space center, the slope of the phase with TE at the k-space center was used for respiration correction. Data was then reconstructed using a sliding filter with 13 lines in the innermost ring (1.56 s temporal resolution), with each successive ring using enough lines to meet the Nyquist criteria. The KWIC filtered k-space distribution repeats after 29 reconstructed time points. Echo phase was combined using a weighted least squares estimate. PRF temperature was also calculated using the combined echo phase. T2*/ρ maps were calculated using linear regression of the log of the magnitude images along the echo dimension.

Results
Figure 204 shows the water and fat images produced from the sequence using the three point Dixon method. Figure 205 shows the standard deviation through time images of the PRF temperature measurements for four cases: PRF temperatures were calculated using both the first time point and the time point with the same k-space sampling pattern as the reference phase for the 13th echo phase and the combined echo phase. Table 8 lists the spatially averaged precision of all tissue voxels in the breast and pork for the four cases. Figure 206 shows the PRF temperature, and percent change in ρ and T2* values vs. time of the hottest voxel while heating with FUS in pork.

Conclusions
All measurements display a structured artifact based on the k-space sampling pattern used for reconstruction. For this reason, a pseudo and not pure golden angle increment was chosen to cause the artifact to repeat and thus be removable in temperature difference measurements. The method shown here provides promising results for this sequence and reconstruction method for use in free breathing interventional treatments. The respiration correction also corrects main field drift. The high bandwidth multi-echo readout offers several advantages as it removes the need for fat saturation, provides fat/water images, and T2*/ρ measurements which could be used as another possible measure of temperature change, especially in adipose tissues which do not exhibit a PRF shift with temperature. Echo combination significantly improves PRF temperature precision.

References
2. Song HK, et al. MRM, 2000;44(6), 825-832.

Table 8 (abstract A180). Spatially Averaged PRF Standard Deviation °C

<table>
<thead>
<tr>
<th></th>
<th>1st Image Phase Reference</th>
<th>Library Phase Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breast Coronal</td>
<td>13th Echo Phase, TE = 17.22 ms</td>
<td>1.18</td>
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<td></td>
<td>Combined Echo Phase</td>
<td>0.56</td>
</tr>
<tr>
<td>Breast Sagittal</td>
<td>13th Echo Phase, TE = 17.22 ms</td>
<td>1.9</td>
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<tr>
<td></td>
<td>Combined Echo Phase</td>
<td>0.71</td>
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<tr>
<td>Pork Phantom</td>
<td>13th Echo Phase, TE = 17.22 ms</td>
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<tr>
<td></td>
<td>Combined Echo Phase</td>
<td>0.41</td>
</tr>
</tbody>
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Fig. 204 (abstract A180). Three point Dixon images produced by the sequence and reconstruction. Top: Coronal breast. Middle: Sagittal breast. Bottom: Pork phantom.
Fig. 205 (abstract A180). Standard deviation of temperature through time maps for Top: Coronal and sagittal breast. Bottom: Pork phantom. Maps are derived for four cases. Using just the phase information from the 13th echo or from the combined echo phase, and using the 1st time point as the reference phase or a sampling pattern based library of phase references.

Fig. 206 (abstract A180). Percent change in $\rho$ (rho - red) and $T_2^*$ (green) and calculated PRF temperature change (blue) for the hottest voxel during heating in tissue with FUS in the pork phantom. A sampling pattern based library was used for all 3.