




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Polylactic acid nano- and microchamber arrays for encapsulation of small hydrophilic molecules featuring drug release *via* high intensity focused ultrasound†

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Long term encapsulation combined with spatiotemporal release for a precisely defined quantity of small hydrophilic molecules on demand remains a challenge in various fields ranging from medical drug delivery, controlled release of catalysts to industrial anti-corrosion systems. Free-standing individually sealed polylactic acid (PLA) nano- and microchamber arrays were produced by one-step dip-coating a PDMS stamp into PLA solution for 5 s followed by drying under ambient conditions. The wall thickness of these hydrophobic nano-microchambers is tunable from 150 nm to 7 μm by varying the PLA solution concentration. Furthermore, small hydrophilic molecules were successfully *in situ* precipitated within individual microchambers in the course of solvent evaporation after sonicating the PLA@PDMS stamp to remove air-bubbles and to load the active substance containing solvent. The cargo capacity of single chambers was determined to be in the range of several picograms, while it amounts to several micrograms per cm². Two different methods for sealing chambers were compared: microcontact printing *versus* dip-coating whereby microcontact printing onto a flat PLA sheet allows for entrapment of micro-air-bubbles enabling microchambers with both ultrasound responsiveness and reduced permeability. Cargo release triggered by external high intensity focused ultrasound (HIFU) stimuli is demonstrated by experiment and compared with numerical simulations.

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1. Introduction

Encapsulation systems are urgently needed for active chemicals' delivery to encapsulate enzymes, cells, dyes, drugs or corrosion inhibitors for an extensive variety of advanced applications.¹ Micropacking especially is applicable for a wide

range of applications, including drug delivery,^{2,3} diagnostics,⁴ tissue engineering,⁵ lab-on-chip,⁶ corrosion inhibition,⁷ biosensors,⁸ membranes, templates for synthesis, optical systems,⁹ antifouling,^{10,11} self-healing materials¹ and coatings.⁷ Small molecule ($M_w < 1$ kDa) encapsulation remains a bottleneck, due to the commonly employed encapsulation matrix (*e.g.* polyelectrolyte multilayers, liposomes) being nanoporous.¹² Therefore, long term encapsulation remains a challenge especially in the field of small hydrophilic molecule delivery as well as in various microreactors and anti-corrosion applications.¹³

A recent approach to fabricate polyelectrolyte multilayer (PEM) microchambers based on layer-by-layer (LbL) self-assembly empowered individual PEM chambers to load microparticles with a size of 2.2 μm and oil-based solutions for light sensitive release-on-demand of the said cargo.¹⁴ Nevertheless, in this method, a minimum amount of PEM bilayers is a prerequisite to ensure mechanical stability,¹⁵ resulting in time consuming (3–4 days) lab work.¹⁶ Furthermore the inherent high permeability of LbL microchambers as well as microcapsules for small molecules is only slowly reduced with increasing

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bilayer number.¹⁷ Other approaches utilizing biodegradable layered films produced chambers only able to encapsulate hydrophobic polymers like polystyrene.¹⁸ Current attempts to seal PEM thin films demonstrated significantly lower permeability for hydrophobic compared to hydrophilic PEM films, due to higher chain packing density, which was achieved by altering deposition conditions.¹⁹ For microcapsules, increased hydrophobic barrier properties of the PEM wall were achieved either by incorporation or *in situ* synthesis of nanoparticles as well as by photochemistry.²⁰ Photochemical wall sealing relies on UV exposure for hours, whereby the *in situ* synthesis of carbon dots requires 20 hours at high pressure and temperature, preventing the utilization of biomolecules, yet even more their applications within biological systems.²¹

Except for encapsulation, triggerable drug delivery systems are being intensively studied, which enable controlled release on-demand profiles to enhance therapeutic effectiveness and reduce systemic toxicity.²² Until now, different external stimuli for drug release, such as pH, temperature, laser and microwave radiation have been utilized to alter the drug-carrier shell permeability and facilitate controlled release.²³ Of the aforementioned methods, the laser offers a low penetration depth (~1 cm), whereby changing the pH and temperature of the human torso causes undesirable side effects. Therefore these release stimuli are expected to offer limited application for biological and medical systems.²⁴ Microwaves offer higher penetration depths compared to the laser; however the detailed microwave–drug container interaction is not clear.²⁵ In contrast, ultrasound is already employed as a diagnostic and therapeutic method for many diseases (*i.e.*, prostate cancer, kidney stones). Particularly, high intensity focused ultrasound (HIFU), due to its long penetration depth and non-invasive nature has received extensive attention from biomedical scientists,²⁶ enabling HIFU to become one of the most promising external stimuli for controlled drug release.²⁷

Inspired by the fact that hydrophobic polymers possess efficient barrier properties for small hydrophilic molecules, a novel method to design and fabricate individually sealed polylactic acid (PLA) microchambers is reported in this work. Polylactic acid is a hydrophobic, biodegradable thermoplastic aliphatic polyester derived from naturally occurring organic acid (lactic acid).²⁸ Due to its biocompatibility, complete biodegradability and the non-toxic nature of degradation products, PLA based materials are ideal for polymer engineering, tissue engineering and various medical implants.²⁹ In these biomedical applications,²⁹ implants with restricted load like rods, plates and pins in bone, surgical sutures as well as dispersed drug delivery systems are made out of PLA.^{30–33}

We demonstrate the fabrication of sealed, biocompatible and biodegradable PLA nano- and microchamber arrays by one-step dip-coating and microcontact printing. In order to examine the feasibility of small molecule encapsulation *in situ* within individual microchambers and further investigate their release behaviour under HIFU treatment, the fluorescent dye Rhodamine B (RhB) ($M_w = 479$) and sodium chloride were chosen as model cargo substances. Furthermore, two different

methods for chamber sealing were employed: dip-coating and microcontact printing,^{34–36} respectively, whereby microcontact printing offers cargo sealing within entrapped micro-air bubbles. Air bubbles support encapsulation of the cargo up to 14 days under submerged conditions and enhance ultrasound responsiveness, allowing for spatiotemporal release upon HIFU exposure. Experimental demonstrations are compared with numerical defined finite element (DFE) simulations.

2. Experimental section

2.1 Materials

Poly(lactic acid) biopolymer ((PLA), 3 mm nominal granule size, weight 100 g, natural source) was purchased from Sigma-Aldrich. The poly(dimethylsiloxane) (PDMS) kit (Sylgard 184) was purchased from Dow-Corning, Midland, USA. In addition, PDMS (Elastosil, RT 602) from Wacker AG, München, Germany was used. Sodium chloride and chloroform were purchased from Sigma. Deionized (DI) water from a Milli-Q (Millipore) water purification system was utilized to make all solutions. All chemicals were used as received without further purification.

2.2 Fabrication of sealed free-standing PLA microchambers

Patterned PDMS stamps with microwells were prepared by casting the PDMS pre-polymer and curing agent (10 : 1 ratio) onto silicon masters with nano- or micro-pillars, degassing it for 30 minutes in vacuum and curing it at 70 °C (Sylgard 184 needed 3 hours for curing, Elastosil RT 602 needed 1 hour). The silicon master for micropattern creation exhibits round micropillars with a diameter and a height of 5 μm and 4 μm , respectively, featuring a centre to centre distance of 15 μm (ESIFig. S1†) and it was prepared by traditional photolithography and etching processes at Shenzhen Semiconductor, Shenzhen, China. The silicon master for nanochambers (500 nm diameter, 1 μm height, 1.5 μm centre to centre distance) was prepared *via* electron lithography at Tomsk Polytechnic University. The PDMS was cut with a sharp surgery knife and lifted off the silicon master.

PLA was dissolved in chloroform at different mass concentrations ($m\% = 1\%$, 2% and 5%) to determine the influence of PLA concentration on the resulting film thickness. The PDMS stamp was dip-coated for 5 s into the PLA solution, *via* a dip-coating robot³⁷ which featured an air pressure controlled robot arm performing pull-out at a speed of 8 cm s^{-1} . By the same method, a flat PLA sheet was coated with PLA by dipping a PDMS sheet without patterns for 5 s at the same pull-out speeds. The patterned PDMS stamp@PLA microchamber was sealed by printing it onto a flat PDMS sheet@PLA under pressure (2 kg cm^{-2}) for 10 s; subsequently the sealed PLA microchamber was collected after lifting off the PDMS stamp.

2.3 NaCl and RhB encapsulation

To examine the feasibility of small molecule encapsulation within individual polylactic acid (PLA) microchambers and to



further investigate their release behaviour under High Intensity Focused Ultrasound (HIFU) radiation, sodium chloride ($M_w = 58.44$) and a fluorescent small molecular dye (Rhodamine B) RhB ($M_w = 479$) were chosen as model drug substances. Both NaCl (3 M) and RhB (0.2 mg mL^{-1}) were dissolved in DI water and stored in a 25 ml polymer tube. Afterwards the PLA microchambers@PDMS stamp was dipped into the model drug solution under ultrasound sonication at 37 kHz for 30 s to remove air bubbles and to load the model drug solution into the microchambers. The sample was wiped with cotton swabs to remove the solution not filling the chambers followed by drying the chamber filling solution under ambient conditions. Such a treatment facilitates drug precipitation within the chambers. Each individual microchamber loaded 8–17 pg of RhB as well as NaCl inside.

2.4 PLA microchamber drug release triggered by HIFU

The sealed PLA@RhB microchambers were inserted into a water filled quartz tube and exposed to HIFU. At specific time intervals (0 s, 5 s, 10 s, 20 s, 60 s, 100 s, 120 s, 150 s), the PLA@RhB microchambers were taken out for measurements. It is noted that PLA chambers exposed to HIFU contain no sputtered gold, as it was added later for electron microscopic observations only. The amount of released RhB was calculated by measuring the solution fluorescence intensity corresponding to the HIFU treatment duration (shown in ESI, Fig. S2†). To ensure the release was not induced by diffusion, the release of the PLA@RhB microchambers (PLA $m\% = 2\%$) kept in the dark (to prevent photobleaching of RhB) without any treatment was also monitored.

2.5 Characterization

Scanning electron microscopy (SEM) (FEI, Inspect-F) was used to measure PLA microchamber morphologies after 2 min of gold sputter coating ($\sim 10 \text{ nm}$ of gold) to ensure conductive samples. SEM observation was carried out using an acceleration voltage of 10 kV, a spot size of 3.5, and a working distance of approximately 10 mm. Fluorescence spectrometry (LS 55, PerkinElmer) was used to analyse fluorescence from the sample within quartz cuvettes (S10C, Sigma). Confocal Laser Scanning Microscopy (CLSM) images were obtained with a Leica TS confocal scanning system (Leica, Heidelberg, Germany) equipped with a $20\times$ objective. Chamber loading was determined *via* binary micrographs and analysis of the chamber content was performed *via* the ImageJ³⁸ “analyse particles” plugin.

For determining the effects of the PLA dipping solution's withdrawal speed, the aforementioned dipping robot with a defined withdrawal speed of 80 mm s^{-1} and a home build robot arm with controllable withdrawal speeds from 0 to 4.5 mm s^{-1} were used. The home build robot comprised of a USB-PIO digital IO card, an OR-8 relay card (both BMCM, Maisach, Germany), a ZD-6560-V3C (Shenzhen TOBY Technology Co. Ltd, Shenzhen, China) stepping motor controller, as well as a 90° angular plate on a linear optical stage which is driven by a stepping motor (TSA-150, Zolix

Instruments Co., Ltd, Beijing, China). This linear optical stage was mounted *via* a 90° right-angle bracket to a $30 \times 30 \text{ cm}$ breadboard (PHOB-3 and OTBB33-1 from Zolix Instruments Co., Ltd, Beijing China). The control software of both robots was customized in LabView 2007 and 2011.

Release experiments were performed by a home built HIFU system, operating at 2.5 MHz with an acoustical density in the focal area of $200\text{--}300 \text{ W cm}^{-2}$ displayed in ESI, Fig. S3.† The HIFU system consists of two units: the first device is a generator for radiofrequency signals that feeds the piezoelectric transducer generating the ultrasonic beam. The generator is electrically matched with the piezoelectric transducer and is obliged to generate the electrical signal with the power tunable from zero up to several hundred watts per cm^2 , whereby an energy density of $200\text{--}300 \text{ W cm}^{-2}$ was adjusted. The generation frequency of the radio signal used corresponds to the resonance frequency of the transducer, which is equal to 2.5 MHz. The second device is the acoustical part of the set up and consists of a table supporting the piezoelectric transducer. On top of the transducer is a quartz tube which contains water for drug release studies. The said quartz tube is hermetically fixed on the plate of the table while the piezoelectric transducer represents a spherical acoustic lens made out of piezoceramics. Both surfaces of the transducer are covered by silver electrodes. Our transducer is obliged with a high acoustical quality factor Q and generates a focused ultrasonic beam with a focal distance of about 12 mm in water, used as acoustical media and the sound conductor. The diameter (D) of the sound beam in the focal plane was as small as $500 \times 500 \mu\text{m}^2$.

Defined finite element simulations were performed with Comsol Multiphysics 4.7 (Comsol, Stockholm, Sweden) utilizing the acoustothermal plugin. The parameters used were obtained from Comsol's internal databases, as well as from ref. 39–41. The modelled chambers had a diameter of $5 \mu\text{m}$ and a wall thickness of 300 nm, which is in the range between 1 and 2% PLA chambers produced in this study.

3. Results

3.1. PLA microchamber fabrication

In order to produce polylactic acid (PLA) microchamber arrays, a silicone rubber (polydimethylsiloxane (PDMS)) stamp with microwells needs to be produced. The said PDMS stamp can be fabricated by curing PDMS on top of a silicon master, resulting in a negative replica of the silicon master, as displayed in ESI Fig. S1.†⁴² A PLA thin film was formed on top of the PDMS stamp (microwells on the surface) by dipping it into the PLA solution for 5 seconds (s), lifting the PDMS stamp@PLA out of the PLA solution, followed by air drying. Furthermore, the PDMS stamp@PLA microchamber was sonicated inside the drug solution for 30 s to remove trapped air bubbles and to load the drug solution at the same time. The PLA microchambers were sealed by microcontact printing^{35,43} them onto another PLA thin film by small applied pressure, after the cargo was *in situ* precipitated by evaporation of the



solvent within the microchambers (Fig. 1A). Afterwards, the sealed microchambers were delaminated from the PDMS sheet allowing freestanding drug delivery microchamber arrays, as the top view and the backside view in Fig. 1B and C demonstrate. Sealing by microcontact printing enables the individual PLA microchambers to entrap an air bubble within its containment, which is proven by focused ion beam and scanning electron microscopy (FIB-SEM) slicing (Fig. 1D).

Please note that the uneven wall thicknesses in Fig. 1D are caused by PLA melting upon FIB slicing. It is worth noting that a successful PLA thin film and therefore chamber production requires a proper adjustment of the deposition conditions whereby three main parameters define the PLA thin film thickness. The first parameter is the PDMS stamp exposure time within the PLA solution: despite PDMS being crosslinked and insoluble in chloroform it can swell and deform the chamber structure⁴⁴ at exposure times exceeding 10 s. Therefore, exposure times around 5 s were chosen. The second parameter is the PDMS stamp's withdrawal speed. As demonstrated in Fig. 2, PLA withdrawal speeds above 3 mm s^{-1} result in slightly thinner films than below, which is caused by shear force. It should be noted that the PLA thin film ($m\% = 2\%$) thickness is quite stable around $1 \mu\text{m}$. The third and most important parameter for reproducible PLA thin film production is the PLA solution concentration, which results in PLA nanofilms (50–150 nm) for 1% up to $7 \mu\text{m}$ thick films causing filled PLA pillars instead of empty chambers at 5% (Fig. 2).

Utilizing 1% PLA solution and a PDMS stamp exhibiting wells with 500 nm diameter and $1 \mu\text{m}$ length, nanochambers can be produced, as demonstrated in ESI, Fig. S4.† Such thin nanostructures were however at the stability limit as can be seen by chamber bending in ESI Fig. S4.† In addition, the

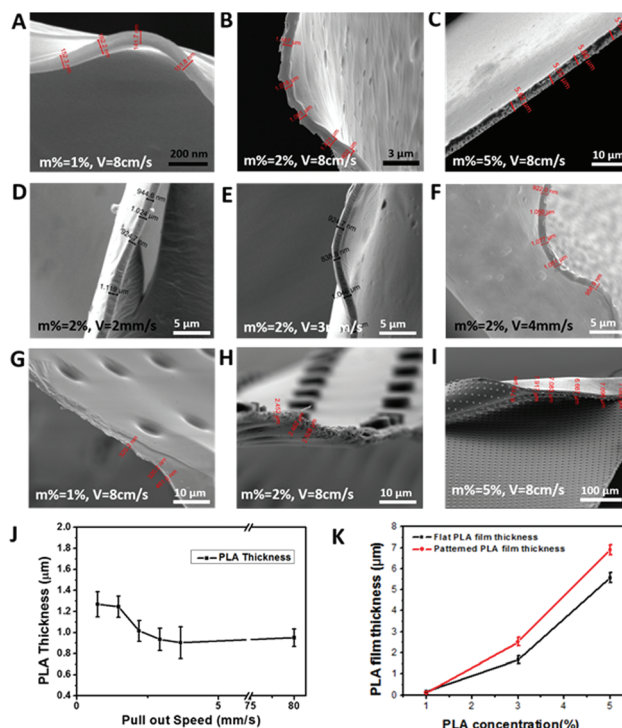


Fig. 2 Characterization of the PLA thin film thickness. SEM images of PLA films: (A–C) Same dip-coating pull-out speed ($v = 8 \text{ cm s}^{-1}$), PLA thin film thickness versus different PLA concentrations ($m\% = 1\%$, 2% and 5%); (D–F) the same PLA concentration ($m\% = 2\%$), PLA thin film thickness versus different pull-out speeds ($v = 2 \text{ mm s}^{-1}$, 3 mm s^{-1} and 4 mm s^{-1}); (G–I) the same dip-coating pull-out speed ($v = 8 \text{ cm s}^{-1}$), patterned PLA microchamber thickness versus different PLA concentrations (1%, 2%, 5%). (J) Influence of pullout speed on PLA thickness with the same PLA concentration (2%), (K) PLA film thickness in relation to PLA solution concentration with the same pull-out speed ($v = 8 \text{ cm s}^{-1}$).

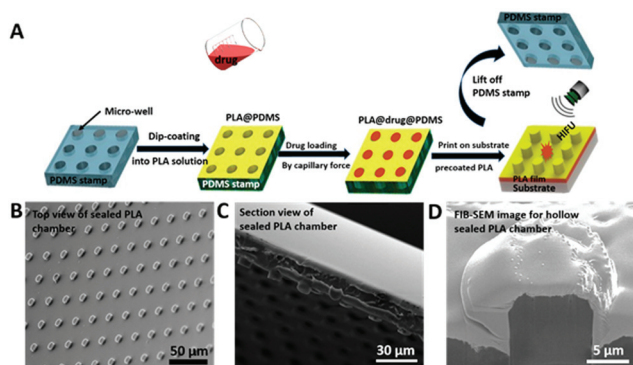


Fig. 1 Preparation and resulting PLA ($m\% = 2\%$) thin film chambers. (A) Schematic illustration of the fabrication of drug loaded poly(lactic acid) (PLA) microchambers and drug release upon HIFU sonication. Scanning electron microscopy (SEM) image with the top view (B) and section view (C) of fabricated free-standing hollow PLA microchambers sealed by imprinting onto a flat PDMS sheet pre-coated with a PLA thin film. (D) SEM image of a PLA microchamber sealed on PLA and cut by a focused ion beam (FIB) proving that the microchamber retains its hollow structure after sealing by imprinting. Uneven wall thicknesses in (D) are caused by PLA melting upon FIB slicing.

films suffered from sensitivity to the electron beam, melting under zoomed in observations. For this reason we focus for the rest of this study on microchambers.

Due to films above 3% resulting in filled chambers, all the following experiments were performed with 2% PLA solution concentration (except stated otherwise), due to an optimum stability to chamber volume ratio.

3.2. Cargo loading and sealing

Cargo loading into PLA ($m\% = 2\%$) microchambers was performed *via* sonication within NaCl solution (3 M), and drying in air (shown in Fig. 3). The said treatment allows chamber *in situ* NaCl crystal growth (Fig. 3B). Utilizing 1% PLA solution, several dozens of nanometer thin PLA membranes which are vacuum stable (Fig. 3D) and mainly electron transparent (despite a 10 nm gold conduction layer) are formed.⁴⁵ The electrons penetrate through the PLA sealing layer, arrive at the underlying NaCl crystal and stimulate X-ray emission, which escapes the chamber, reaching the detector.⁴⁵ Fig. 3E is the resulting EDX-mapping micrograph, proving successfully the presence of sealed NaCl crystals within microchambers. Subsequent FIB-



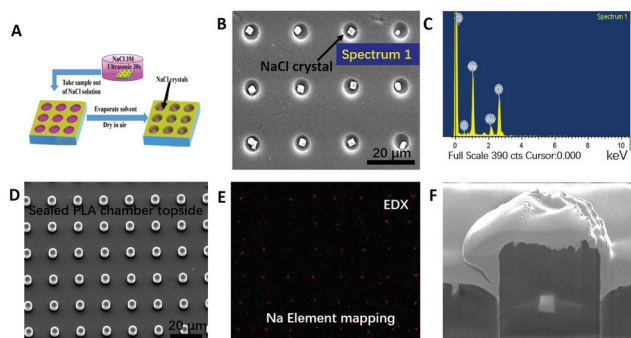


Fig. 3 Encapsulation of small hydrophilic molecules within air bubbles in PLA thin films and proof of successful encapsulation. (A) Scheme of NaCl crystal encapsulation within hydrophobic PLA microchambers. (B) SEM image of PLA@NaCl crystals in microchambers before sealing (PLA $m\%$ = 2%), the size of NaCl crystals is $1 \pm 0.2 \mu\text{m}$. (C) EDX analysis of NaCl crystals loaded into PLA chambers. (D) SEM image of the sealed PLA@NaCl microchambers (PLA $m\%$ = 1%). (E) EDX mapping analysis of PLA@NaCl microchambers, proving that NaCl is within the chambers of (D). (F) Focused ion beam (FIB) image of a half-cut and sealed PLA@NaCl microchamber (PLA $m\%$ = 2%), uneven wall thickness from PLA melting upon FIB cutting.

cutting of loaded and sealed 2% microchambers proves the successful encapsulation of NaCl crystals (Fig. 3F).

The long term encapsulation of small hydrophilic molecules was also studied in this work. Compared to single NaCl crystal encapsulation, the small molecular dye RhB ($\text{MW} = 479 \text{ g mol}^{-1}$) was also successfully loaded into PLA microchambers in the form of several nano-sized precipitates, as demonstrated in Fig. 4A–D. The fact that these dyes remain in the precipitated state under submerged conditions for weeks proves the entrap-

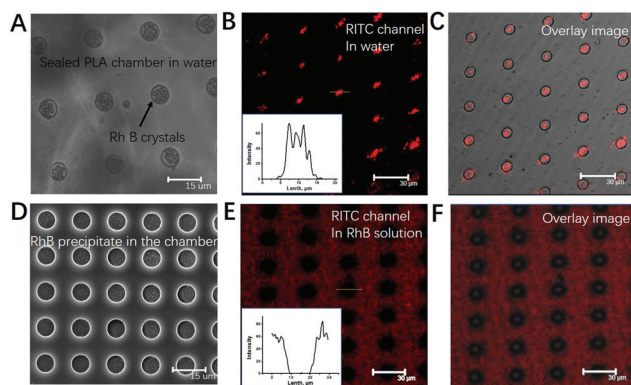


Fig. 4 Determination of the sealing properties of submerged PLA chambers: (A) optical micrograph of the sealed PLA@RhB precipitate within microchambers under submerged conditions after 2 hours. Confocal laser scanning microscopy (CLSM) image of the sealed PLA@RhB precipitate within microchambers in an RITC channel (B) and an overlay micrograph (C). (D) SEM image of the unsealed PLA@RhB precipitate within microchambers. (E), (F) CLSM micrographs of sealed, empty PLA microchambers in Rh-B solution in an RITC channel and an overlay image proving efficient chamber sealing. The line scan insets show the relative fluorescence intensity of the corresponding microchambers.

ment of air bubbles within the microchambers. Fig. 4A–C display micrographs of the sealed PLA microchamber@RhB precipitates submerged under aqueous conditions after 2 hours, indicating that the RhB crystals are not in contact with water, otherwise the RhB crystals would be immediately dissolved. The amount of encapsulated RhB that precipitates in each microchamber is $8 \times 10^{-12} \text{ g}$ – $1.7 \times 10^{-11} \text{ g}$ (or 8–17 pg), which amounts to $1.28 \mu\text{g}$ – $2.72 \mu\text{g cm}^{-2}$ as also shown in ESI, Fig. S5.† This rationale is supported by contrast experiments, proving exclusion of aqueous RhB solution from sealed PLA chambers as CLSM micrographs in Fig. 4E and F demonstrate.

3.3. Drug release

High intensity focused ultrasound (HIFU) is a highly efficient and non-invasive technique, which is already employed as a diagnostic and therapeutic method for many diseases.⁴⁶ Here, we demonstrate that apart from long term encapsulation, one-step fabricated PLA microchambers support rapid drug release triggered by HIFU. The SEM images of sealed biodegradable PLA microchambers after exposure to different HIFU sonication times are displayed in Fig. 5, demonstrating that sealed

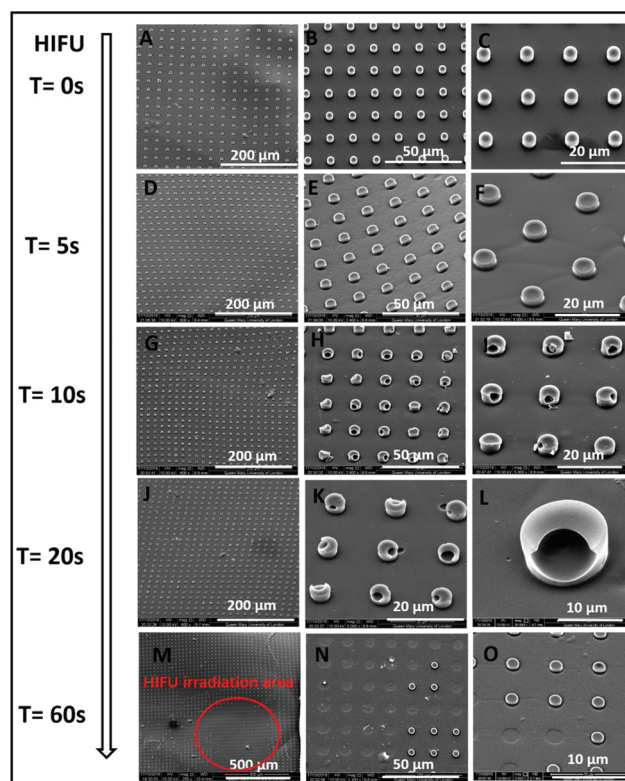


Fig. 5 High intensity focused ultrasound (HIFU) triggered chamber opening versus exposure time. SEM images of sealed PLA@NaCl microchambers before (0 s first row) and after HIFU exposure for 5 s (second row), the microchambers look comparable to non-ultrasound exposed chambers; after 10 s (third row) and 20 s (fourth row) HIFU treatment, wells appear in chambers, whereby the absence of crystals proves cargo release; within 60 s (last row), the PLA microchamber "caps" are blown away by HIFU. The red circle in M shows the HIFU irradiation area (D ~500 μm). The images in the same row shows at different magnifications.



PLA microchambers with entrapped air bubbles are responsive to ultrasound. HIFU exposure times below 5 s cause no adverse effects to the PLA chambers (Fig. 5A–F), but 10 s already open small wells in the microchamber “caps”, as indicated in Fig. 5G–I. 20 s of HIFU irradiation causes the wells in the PLA microchambers to grow slightly bigger (Fig. 5J–L). After 60 s the PLA microchamber “caps” are completely eradicated, leaving craters on the irradiated areas ($D \sim 500 \mu\text{m}$) behind, as shown in Fig. 5M–O. The corresponding fluorescence images of the PLA microchamber under HIFU radiation for RhB released into water are in good agreement with the SEM data (ESI, Fig. S6†).

Comparison experiments employing PLA microchambers sealed by one more dip-coating-step without air-bubble entrapment were also performed (ESI, Fig. S7†). Sealed PLA microchambers without air-bubble entrapment cannot be opened by HIFU treatment, even after increasing the exposure time up to 30 minutes. The explanation for this >300 times enhanced ultrasound sensitivity possessed by sealed PLA microchambers with air-bubble entrapment is the high refractive index differences between air and water.

The quantitative measurement of released RhB triggered by HIFU was studied (Fig. 6). To ensure the release was not induced by diffusion, the release behaviour of the PLA@RhB microchambers (PLA $m\%$ = 2%) kept in the dark (to prevent photobleaching of RhB) without any treatment was also monitored.

Less than ~5% RhB is released by diffusion within 14 days due to low PLA permeability for gases and water vapour.³⁰ For the same reason PLA is also widely used for food packaging.⁴⁷

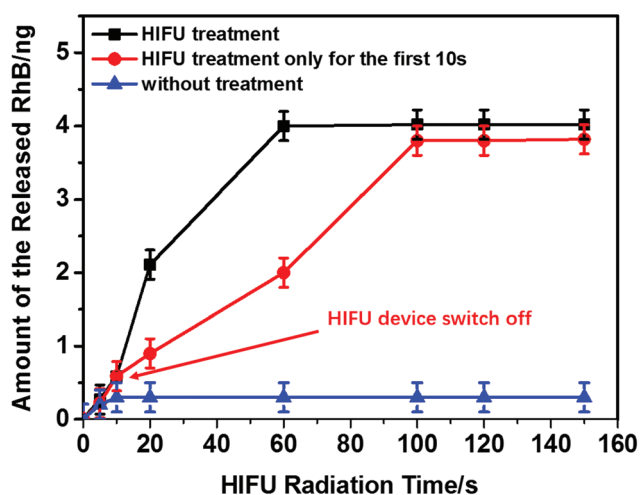


Fig. 6 Amount of released RhB from sealed PLA@RhB microchambers triggered by HIFU. The black line shows microchambers exposed to continuous HIFU radiation and the red line shows the PLA microchambers with HIFU treatment for 10 s; then the HIFU device was switched off; the blue line visualizes a control experiment without any HIFU exposure for the PLA microchambers corresponding to the HIFU treatment duration (0 s, 5 s, 10 s, 20 s, 60 s, 100 s, 120 s, 150 s). Values from calibration and the release spectra of ESI Fig. S2.†

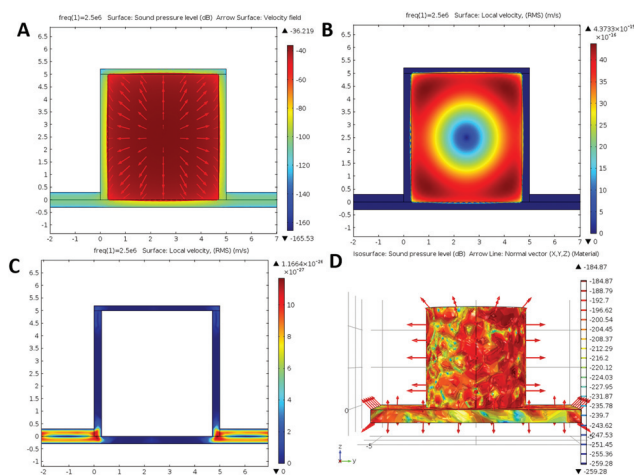


Fig. 7 Defined finite element simulation of non-gas filled and gas filled chambers: (A) 2D simulation of acoustic pressure interaction and pressure vectors within a gas filled micro-chamber, showing slight deformations due to the pressure; (B) 2D simulation of particle velocity for an air-filled chamber proving peak maxima in the corners and slight deformations; (C) 2D simulation of the water filled microchamber proving low particle speed and no deformation; (D) air filled 3D micro-chamber proving pressure peaks and force vectors pointing outwards from the chamber, with a pressure peak in the upper corner.

3.4. Ultrasound harvesting mechanism

Defined finite element (DFE) simulations were used to determine the interaction between the ultrasound pressure wave and the herein utilized PLA microchambers. A significant acoustic pressure in the range of 40–160 Db (decibel) is determined within the aforementioned microchambers. This results in a maximum energy harvesting of $\sim 10 \text{ kW m}^{-2}$ out of 1.27 MW m^{-2} generated by the HIFU (more device details are shown in the Experimental section and ESI, Fig. S3†), corresponding to an $\sim 0.8\%$ conversion efficiency of PLA chambers with a force vector pointing outward from the chamber, even slightly deforming them (Fig. 7A). This pressure is accompanied by an 11 orders of magnitude higher air molecule speed compared to water filled chambers (Fig. 7B and C). A decreased particle speed and harvested energy for chambers not possessing air bubbles explains the failure to open them in the experiment. A comparison between Fig. 5 and 7D reveals that 3D chambers possess local pressure peaks on positions corresponding to the appearance of wells in Fig. 5; these pressure peaks facilitate a diagonal as well as a sideward chamber opening observed in the experiment.

4. Conclusion

In summary, we have developed a one-step dip-coating strategy for fabricating free-standing biocompatible polylactic acid microchamber arrays (whereby the production limit is $\sim 500 \text{ nm}$ sized chambers), able to encapsulate small hydrophilic molecules as well as to entrap air bubbles. To the best of our knowledge, this is the first report about the *in situ* crystalli-



zation of NaCl and RhB inside the individual microchambers (~8–17 pg per chamber) with long term encapsulation in aqueous solution up to 14 days. Biodegradable PLA nano- and microchamber fabrication requires a relatively short production time (5 s, compared to several days for the present PEM chambers¹⁷ and about a day for capsules⁴⁸), low permeability, and high ultrasound sensitivity due to air bubble entrapment. Numerical simulations reveal an 11 order of magnitude higher molecule speed for PLA microchambers with air bubbles compared to water filled chambers. The determined ultrasound energy harvesting efficiency is ~0.8%. The existence of ultrasound induced local sideward directed pressure peaks was revealed to cause diagonally upwards directed chamber piercing. In addition, this highly ordered array of PLA microchambers enables the option of programmed release-on-demand for precisely defined active quantities from individual chambers in a site- and time specific manner. With these features, we expect PLA microchambers to offer considerable potential for medical applications and chemical and biochemical studies (e.g. single cell level, implant coatings, catalyst release).

It is worth pointing out that the applied method might aid dispersed drug delivery systems like PEM capsules and micro motors,⁴⁹ if the sealing mechanism can be transferred to disperse drug delivery systems. The approach might be used in the future also to capture desired cell types if utilized in the bloodstream, for example to capture cancer cells, or in the case of stents that capture and promote the growth of blood vessel cells. For such purposes the surface might be coated with cellular membranes and biomolecules by protocols outlined in ref. 50 and 51. Another potential application is the utilization of our system in combination with or as a replacement for electrospun PLA scaffolds which are currently used for tissue growth and as implants.^{52,53} In such a case electrospun fibres can be used to adjust the surface area or mechanical properties.

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