OVER-PRESSURE SUPPRESSES ULTRASONIC-INDUCED DRUG UPTAKE

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Abstract—Ultrasound (US) is used to enhance and target delivery of drugs and genes to cancer tissues. The present study further examines the role of acoustic cavitation in US-induced permeabilization of cell membranes and subsequent drug or gene uptake by the cell. Rat colon cancer cells were exposed to ultrasound at various static pressures to examine the hypothesis that oscillating bubbles, also known as cavitating bubbles, permeabilize cells. Increasing pressure suppresses bubble cavitation activity; thus, if applied pressure were to reduce drug uptake, cell permeabilization would be strongly linked to bubble cavitation activity. Cells were exposed to 476 kHｚ pulsed ultrasound at average intensities of 2.75 W/cm² and 5.5 W/cm² at various pressures and times in an isothermal chamber. Cell fractions with reversible membrane damage (calcein uptake) and irreversible damage (propidium iodide uptake) were analyzed by flow cytometry. Pressurization to 3 atm nearly eliminated the biological effect of US in promoting calcein uptake. Data also showed a linear increase in membrane permeability with respect to insonation time and intensity. This research shows that US-mediated cell membrane permeability is likely linked to cavitation bubble activity. (E-mail: pitt@byu.edu) © 2009 World Federation for Ultrasound in Medicine & Biology.

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INTRODUCTION

Ultrasound currently finds significant application in diagnostic and therapeutic medicine. Diagnostic ultrasound, typically employing frequencies greater than 1 MHz, takes advantage of the ability of ultrasound to penetrate soft tissue and reflect from interfaces, thus, allowing imaging of internal body structures. In physical therapy applications, ultrasound is used to warm targeted tissues and typically employs frequencies around 1 MHz.

A recent development in therapeutic ultrasound is its application to deliver drugs and genes to a variety of tissues (Husseini et al. 2008; Mitragotri 2005; Muller et al. 2007; Pitt et al. 2004). Many drugs and genes have been delivered to specific tissues and locations such as tumors (Frenkel 2008; Nelson et al. 2002; Reddy 2005), cardiac tissue (Chen et al. 2003; Mayer et al. 2008; Muller et al. 2007), the brain (Hynynen 2008; Kinoshita et al. 2006; Meairs et al. 2007) and occluded vessels (Francis 2001). Ultrasound is associated with permeabilization of cell membranes, but the exact mechanism by which this occurs remains under investigation (Prentice et al. 2005; Schlicher et al. 2006; Tachibana et al. 1999). Because ultrasound is composed of oscillating pressure waves, it causes direct mechanical effects on tissue. It also acts physically on bubbles, causing them to oscillate in size—a phenomenon called cavitation. It has been hypothesized that intense cavitation, resulting from sufficiently intense ultrasonication, creates an increase in cell membrane permeability (Guzman et al. 2001; Prentice et al. 2005; Schlicher et al. 2006).

There are several mechanisms whereby ultrasonication and cavitation may be involved in drug delivery. Even in the absence of gas bubble cavitation, the oscillating pressure waves produce a slight oscillatory motion of fluid. This action will increase molecular transport by increasing local molecular motion. Another phenomenon called acoustic streaming occurs when the momentum of the sound wave is transferred to an absorbing fluid (like water), thus, generating convective fluid flow in the direction of the propagating ultrasonic waves. Although these phenomena will enhance the transport of drug to a cell surface, it is doubtful that these low energy motions can change cell membrane permeability.

A more pronounced effect of ultrasonication is mediated by cavitation when gas bubbles are present. There are two general categories of cavitation: stable and collapse cavitation, also known as noninertial and inertial cavitation. Stable cavitation is the cyclic volumetric oscillation of microbubbles without a violent collapse event. This stable oscillation phenomenon induces convective flow around the bubble called microstreaming, which causes two important effects. First, genes or drugs in solution are convected at high velocities. Second, the rapid movement of fluid and changes in pressure...
associated with bubble oscillation generate high shear stresses adjacent to the bubble that may create stress on nearby cell membranes (Van Wamel et al. 2004). Under carefully controlled conditions, stable cavitation was found to be sufficiently energetic to permeabilize cell membranes (Nyborg 2001; Rooney 1970).

Inertial cavitation is much more violent than stable cavitation, and it occurs when the oscillations in bubble radius become so large that the inertia of the imploding wall compresses the internal gas to extremely high temperatures and pressures, which in turn forms free radicals and releases energy in the form of heat and powerful shockwaves (Brennen 1995). In addition, it has been shown that an asymmetric collapse may occur near cell surfaces, thus causing a high-speed jet of extracellular solution to be injected into the cell or tissue (Prentice et al. 2005). It is hypothesized that through shear stresses and shockwaves associated with inertial cavitation events, cell membranes are permeabilized, therefore causing an increase in passive drug uptake (Schlicher et al. 2006).

Several methods have been used to study the role of cavitation in biological systems. Using sonoluminescence as an indicator of cavitation, Cochran measured flashes of light that were believed to be emitted upon the collapse of microbubbles (Cochran et al. 2001). Other evidence of the presence of cavitation has been to “listen” to acoustic signatures present in the fluid during sonication by using an acoustic detector (Ciaravino et al. 1981; Hill 1971; Hussein et al. 2005, 2007; Tang et al. 2001; Tezel et al. 2002).

Cavitation phenomena can be reduced by pressurizing the bubbles (Brennen 1995; Ciaravino et al. 1981; Delius 1997; Richardson et al. 2007). Higher static pressure reduces the initial bubble radius and increases the internal gas density. Therefore, if a phenomenon associated with ultrasound decreases as static pressure increases, then the phenomenon is most probably caused by cavitation, not by fluid oscillation (independent of bubbles) or acoustic streaming. The unique aspect of this study is the use of a pressurized chamber to investigate the effect of ultrasound and cavitation on cellular uptake of a model drug. This study investigated the hypothesis that cavitation is involved in permeabilizing cell membranes and not simply fluid oscillation or acoustic streaming. Specifically, we show that the application of static pressure correlates with a reduction in membrane permeability.

MATERIALS AND METHODS

Reagents

Calcein (622.5 g/mol) (MP Biomedicals, Inc., Aurora, OH, USA) was dissolved into 1X Ca²⁺/Mg²⁺-free Dulbecco’s phosphate-buffered saline (DPBS) to create a stock solution of 0.1 mM, stored at 4°C and sheltered from light. Fifty L of stock solution were then added to 0.5 mL of cell suspension. Propidium iodide (PI; Sigma, St. Louis, MO, USA) was added to the cell suspension approximately 10 min before flow cytometry so that the final concentration was 10 M.

Cell culture

DHD/K12 TRb rat colon cancer cells (#90062901, European Cell Culture Collection) were cultured at 37°C, 5% CO₂ in RPMI 1640 medium (Gibco BRL, Gaithersburg, MD, USA) and supplemented with 20% cosmic calf serum (HyClone, Logan, UT, USA). The cells were harvested during their exponential phase of growth and suspended at a concentration of 1 × 10⁶ cells/mL in medium containing 10 M calcein. These cells were selected because they are responsive to ultrasonic-activated chemotherapy in vivo in a rat model (Nelson et al. 2002).

Apparatus

The chamber used for sonicating the cells was an aluminum box of 2 L volume lined with about 1 L of acoustically absorbing rubber on the bottom and sides. This rubber had a corrugated surface (to scatter any nonabsorbed sound). Placed on one wall was a 476-kHz ultrasonic transducer (model H-104; Sonic Concepts, Woodinville, WA, USA), which has an active diameter of 64 mm, a radius of curvature of 62.64 mm, a focal length of 51.74 mm and a center frequency of 476 kHz. Mapping of the interior of the box with a needle hydrophone (ONDA Corporation, Sunnyvale, CA, USA) showed that there were no standing waves inside the box when insonified at 476 kHz.

An XYZ stage on top of the box positioned a pressure tube containing the sample at the focal point of the transducer (Fig. 1). Within this tube of cellulose butyrate, a hollow rod held a small polyethylene pipette bulb filled with the cell suspension. A hole in the hollow rod connected the gas in the pressure tube with the gas above the cell suspension. Pressure was applied to the interior of the tube (and to the sample) from a compressed air source and regulator at 0, 1.2 and 3 atmosphere (gauge pressure). The local pressure at Brigham Young University is 86 kPa, so the experimental pressures were 86, 187, 288 and 389 kPa absolute pressure.

The wall thicknesses of the pressure tube and PE sample holder were 0.46 and 0.42 mm, respectively. Over pressurization to 3 atm was calculated to expand the tube radius by 0.6% and, although it may have shifted the center of the focal volume by 1%, any changes in cell permeabilization were estimated to be less than the noise level of experimental error in this system. The calculated peak rarefaction pressure within the PE sample holder was not corrected for any attenuation of the thin films of cellulose butyrate and PE.
The water in the chamber external to the pressure tube was recirculated through a heater at 37°C and then through a 0.2-m filter to remove particulate and gas bubbles that might serve as nuclei for cavitation bubbles.

**Electronics**

A sine wave at 476 kHz was produced by a signal generator (Hewlett Packard, model 33120A). A 1:10 duty cycle was produced with a 100-cycle burst at a frequency of 476 Hz and was monitored on a digital oscilloscope. This signal was amplified (ENI Model 240L; ENI Inc., Rochester, NY, USA) and then sent through a matching network to the ultrasonic transducer. The 10% duty cycle was selected because it was a good compromise between a very low duty cycle (say 1% that would require long experimental times) and continuous insonation that would heat up the sample. The intensity values reported herein are spatial peak temporal averages \( I_{\text{ptt}} \) unless otherwise noted.

The intensity of US at the focal point was determined in separate experiments using a pendulum radiation meter consisting of a 1.5-mm-diameter stainless steel ball suspended by two Kevlar fibers in an inverted V-configuration (Chen et al. 2004; Dunn et al. 1977). The deflection of the steel ball was measured with a micrometer (0.005 mm precision) as follows. The pendulum assembly was mounted on an XYZ micrometer stage and the steel ball located at the acoustic focal point and illuminated by a laser. When the transducer was activated the ball was deflected backward and then the horizontal micrometer of the stage was adjusted to bring the ball back to the focal point illuminated by the laser. The difference in micrometer readings gave the deflection from which the acoustic intensity was calculated.

**Insonation of cells**

The polyethylene bulb was filled with the suspension of cells in calcein and then attached to the hollow rod. The pressurizable tube was filled with degassed water and sealed to the lid of the box with a sealing ring. Once the tube and sample were pressurized to the desired static pressure, the lid was placed on the Al box and the sample positioned at the focal point with the XYZ micrometer stage. Ultrasound was applied for various times and intensities, after which the static pressure was released and the sample recovered.

**Flow cytometry**

After insonation, cells were washed three times in 1X DPBS Ca\(^2+\)/Mg\(^2+\) free. PI was added to 0.5 mL of cell solution to a final concentration of 10 M. Cells were analyzed in a flow cytometer (Epics XL; Beckman Coulter, Fullerton, CA, USA) for forward and side scatter, cell fractions with reversible (calcein uptake) and irreversible membrane damage (PI uptake).

**RESULTS**

**Acoustic intensity versus exposure time**

A minimal, and often negligible, amount of calcein uptake occurred between temporal average intensities of 0.125 W/cm\(^2\) and 1.17 W/cm\(^2\) (data not shown). However, at temporal average intensities of 2.75 W/cm\(^2\) and 5.5 W/cm\(^2\), there were much higher levels of uptake that were more easily detected; therefore, these intensities were selected for subsequent experiments.

![Fig. 2. Average (n = 3) calcein uptake as a function of time and average intensity at atmospheric pressure (86 kPa). At 5.5 W/cm\(^2\) \( I_{\text{ptt}} \), there were consistently higher amounts of drug uptake than at 2.75 W/cm\(^2\) \( I_{\text{ptt}} \). The lines are a least square fit of the data.](image-url)
As measured by calcein uptake into viable cells, the amount of molecular uptake was directly proportional to the time and intensity of insonation (Fig. 2). For example, at 10 min of ultrasonic exposure, the average percentage of cells with calcein uptake was 17.5% at 2.75 W/cm², while the average uptake was 34.3% at 5.5 W/cm². Similarly, after 5 min of exposure, the average uptake at 2.75 W/cm² and 5.5 W/cm² were 5.6% and 16.7%, respectively. The data showed that increasing either time or intensity increased the calcein permeability of cells.

Additionally, experiments showed a similar linear relationship between calcein uptake and exposure time at different pressures (Figs. 3 and 4). Most remarkably, pressurization of the samples during the insonation procedure drastically decreased the percent of cells with calcein uptake.

The total energy density delivered to the sample is the product of the average intensity (W/cm²) and the exposure time. The relationship between calcein uptake and total energy density is depicted in Fig. 5. Acoustic energy density appears to be a fundamental parameter because data from both intensities correlate in a linear relationship at a given overpressure.

**Cell death**

Flow cytometry data of propidium iodide uptake (cell death but not cell rupture) showed that there was not a significant increase in dead cells above the usual amount found in the control, with the exception of samples exposed to 5.5 W/cm² for 15 min. In other words, the cells that were permeabilized enough to uptake calcein, were able to re-establish membrane integrity after insonation at the specified energies except at the most severe treatment. At this energy density (4950 J/cm²), more than half of the cells were lysed by the ultrasonication process. Therefore, this data point was not included in the subsequent analysis and higher energy densities were not investigated. Cells that received 3300 J/cm² or lower were permeabilized enough to uptake calcein and apparently were able to re-establish membrane integrity after insonation.

**Increased pressure decreases uptake**

Figures 3 to 5 show that the application of pressure above ambient decreases the cell membrane damage as assessed by calcein uptake. Again, the uptake increased linearly with time (Figs. 3 and 4) and acoustic energy density (Fig. 5) but the rate of uptake decreased as overpressure increased. For example, application of 3 atm overpressure decreased calcein uptake by about 90% for samples at all the times and intensities studied (Fig. 4).
DISCUSSION

Our observations that cell membrane damage and calcein uptake correlates with the intensity, time and total energy of ultrasonic exposure is not novel but is confirmatory of the observations of Prausnitz et al. (Guzman et al. 2001a, 2001b; Keyhani et al. 2001; Schlicher et al. 2006). However, our observation that model drug uptake is a strong function of ambient pressure is novel and aids in explaining the nature of the mechanism that causes membrane damage. We posit that the ultrasound itself does not have a direct effect on the cell membrane, because the ultrasonic acoustic vibrations themselves are not a function of the ambient pressure, except for a negligible change in acoustic wave velocity. However, the behavior of cavitating bubbles is strongly influenced by the ambient pressure; thus, the ultrasound is apparently acting indirectly through the action of the cavitating bubbles, which are strongly responsive to the ambient pressure. Richardson et al. have also shown that increasing the ambient pressure during 80-kHz insonation reduced the background emission attributed to inertial cavitation (Richardson et al. 2007).

Because increasing the pressure was effective in decreasing calcein uptake, the hypothesis that bubble cavitation is the cause of increased membrane permeability is supported. However, the question remains as to whether stable or inertial cavitation (or both) is responsible for the cell membrane permeability. In the presence of a wide range of bubble sizes, the likelihood of inertial cavitation occurring can be generally estimated by a parameter called the “mechanical index” (MI), which is defined as the peak negative pressure (in MPa) divided by the square root of the ultrasonic frequency (in MHz). Inertial cavitation in water containing bubbles (at atmospheric pressure and all sizes) begins to occur above a threshold in mechanical index of about 0.2 to 0.5 (Apfel 1982; Barnett 1998; Bouakaz et al. 2005; Church 2005; Husseini et al. 2005, 2007). Biological effects (again at atmospheric pressure) have been reported above an MI of 0.6 (Barnett 1998; Korosoglou et al. 2006; Miller et al. 1995).

Church points out that the MI threshold defined above is a predictor of inertial cavitation activity for a single acoustic cycle event and that with more cycles, there is a decrease in the value of the MI threshold required to produce inertial cavitation (Church 2005). His equations predict that the threshold MI required to produce inertial cavitation in water with 476 kHz US is 0.17 MPa. In our experiments at 5.5 W/cm², the MI during the 100-cycle pulse was 0.6 and the peak rarefaction pressure was 0.41 MPa. Thus, it is possible that inertial cavitation is occurring in these experiments at ambient atmospheric pressure and it is also possible that inertial cavitation is at least somewhat responsible for increasing membrane permeability. Little theoretical work has been reported on how increasing static pressure will increase the threshold of inertial cavitation, but there are several experimental observations showing that increasing pressure suppresses the amount of inertial cavitation (Fry et al. 1951; Hill 1971; Morton et al. 1983; Richardson et al. 2007) and cell damage (Ciavarrino et al. 1981; Delius 1997; Hill 1971; Morton et al. 1983).

In contrast with other studies that used contrast agents to provide bubbles, our study used no external sources of bubbles. However, micron sized gas bubbles are stable in aqueous solutions in the presence of surfactant molecules that surround and stabilize the bubble. The proteins and other biomolecules found in cell nutrient solutions are adequate surfactants to stabilize gas bubbles (Williams 1983). The solutions used herein were not degassed before resuspending the cells; thus it is postulated that normal amounts of air dissolved in these solutions were nucleated into gas bubbles when the first few low-pressure cycles of ultrasound propagated through the cell suspension.

The slopes of the lines in Figs. 3 and 4 represent the rate of cell permeabilization at various pressures. This linear increase with time is consistent with other observations, as is the observation that the cell permeabilization toward drugs is related to treatment time multiplied by acoustic intensity (Guzman et al. 2001). But perhaps even more significantly, the slopes of the lines in Fig. 5 represent the percentage of cells permeabilized per increment of energy density. This graph plots the slope of the linear regression lines of Fig. 5 at specific absolute pressures.

![Graph](https://via.placeholder.com/150)
be represented by a decaying exponential model of the following form:

\[ \text{cell damage per energy density} = Ae^{-kP} \]

where \( A \) is 0.0185 % damage/(J/cm^2), \( k \) is 0.0085/kPa and \( P \) is absolute pressure. This expression proposes that at 1 atm pressure (101 kPa), 100% of the cells would take up calcne if exposed to 27,760 J/cm^2. This might possibly be achieved by long exposures at low intensities where cell death would be a minimum.

In conclusion, our novel experimental design allowed us to increase the pressure of a cell suspension while applying ultrasound. It was demonstrated that increased pressure suppresses the effect of ultrasound on model drug uptake. The data show that the rate of calcne uptake is proportional to exposure time, exposure intensity and exposure energy, and inversely proportional to overpressure. While these experiments have no direct clinical application, the observations presented herein reveal the mechanism of ultrasonic enhanced drug uptake. Since overpressure suppresses both stable and inertial cavitation, these findings suggest that bubble cavitation has an important role in ultrasonic drug delivery. Continued research on the mechanisms and conditions of ultrasound mediated cell permeabilization will advance the effectivenes of in vivo studies and the eventual clinical application.

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