

Bioeffects Considerations for Diagnostic Ultrasound Contrast Agents

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Abbreviations

CAT, chloramphenicol acetyltransferase; CK, creatine kinase; ECG, electrocardiographic; FDA, Food and Drug Administration; MCE, myocardial contrast echocardiography; MI, mechanical index; PESDA, perfluorocarbon-exposed sonicated dextrose albumin; PRPA, peak rarefactional pressure amplitude; PVC, premature ventricular contraction

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Diagnostic ultrasound contrast agents have been developed for enhancing the echogenicity of blood and for delineating other structures of the body. Approved agents are suspensions of gas bodies (stabilized microbubbles), which have been designed for persistence in the circulation and strong echo return for imaging. The interaction of ultrasound pulses with these gas bodies is a form of acoustic cavitation, and they also may act as inertial cavitation nuclei. This interaction produces mechanical perturbation and a potential for bioeffects on nearby cells or tissues. In vitro, sonoporation and cell death occur at mechanical index (MI) values less than the inertial cavitation threshold. In vivo, bioeffects reported for MI values greater than 0.4 include microvascular leakage, petechiae, cardiomyocyte death, inflammatory cell infiltration, and premature ventricular contractions and are accompanied by gas body destruction within the capillary bed. Bioeffects for MIs of 1.9 or less have been reported in skeletal muscle, fat, myocardium, kidney, liver, and intestine. Therapeutic applications that rely on these bioeffects include targeted drug delivery to the interstitium and DNA transfer into cells for gene therapy. Bioeffects of contrast-aided diagnostic ultrasound happen on a microscopic scale, and their importance in the clinical setting remains uncertain. *Key words:* acoustic cavitation; contrast agent adverse effects; echocardiography; mechanical index.

1. Introduction

The ability to “see” inside the body represents one of the most potent diagnostic tools of modern medicine. Ultrasound imaging is particularly attractive owing to the portability of imaging machines and the inherent safety of low-power acoustic interrogation of tissue. The desire for means to enhance image information has led to the development of contrast agents for pulse-echo diagnostic ultrasound, as for

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other imaging modalities (ie, radiography and magnetic resonance imaging). Although similar in concept to radiologic contrast agents, ultrasound contrast agents have unique properties designed for echogenic interaction with the ultrasonic pulses, which make them special cases for safety assurance in diagnosis.

The primary purpose of ultrasound contrast agents is to enhance the echogenicity of blood. Although modern Doppler imaging equipment is very good at showing the location and motion of rapidly flowing blood, it fails in other important diagnostic problems such as capillary perfusion. What could reasonably be added to blood to enhance its echogenicity? The answer is microbubbles. Over the past 2 decades, contrast agents consisting of suspensions of gas bodies (specially stabilized microbubbles) have been developed that can pass through the lungs after intravenous injection and persist in circulation for useful periods.¹⁻³ Ultrasound contrast agents can enhance B-mode and Doppler images, and special imaging methods can show blood distribution at the capillary level to reveal tissue perfusion.⁴ Owing to the promise of safe and more cost-effective vascular diagnosis than might be possible with contrast-enhanced radiography or magnetic resonance imaging, a substantial research and development effort has been pursued to achieve approval and bring ultrasound contrast agents to the clinic.

The interaction of ultrasound with gas bodies for enhanced echogenicity also introduces a potential for bioeffects. Ultrasonic cavitation is defined as the interaction of ultrasound with a body of gas and is a potent mechanism for biological effects of ultrasound.^{5,6} Cavitation involving microbubbles (microcavitation) has long been recognized as the most likely potential mechanism for nonthermal bioeffects of diagnostic ultrasound.^{7,8} At the megahertz frequencies used for diagnostic ultrasound, the most efficiently echogenic (ie, resonant) gas bodies are a few ($\approx 1-5$) micrometers in diameter. At low ultrasonic pressure amplitudes, less than atmospheric pressure (0.1 MPa), cavitation microbubbles pulsate linearly in response to the time-varying acoustic pressures of the ultrasound field. At higher pressure amplitudes, the pulsation becomes nonlinear, and expansion

during the rarefactional pressure phase may become much greater than the initial microbubble radius. The large expansions are followed by violent collapse of the microbubble, in which the collapse is dominated by the inertia of the inrushing fluid surrounding the microbubble. For this reason, this special case of ultrasonic cavitation is called *inertial cavitation*. The minimum threshold for inertial cavitation was calculated for the diagnostically relevant frequency range,⁹ and this theory served as the basis for the mechanical index (MI) used for display on ultrasound imaging machines.¹⁰

Gas bodies suitable for nucleation of ultrasonic cavitation are normally practically absent from the body, minimizing the possibility of cavitation-related bioeffects for diagnostic ultrasound without contrast agents.⁶ The fortuitous coincidence that the microbubble sizes most strongly activated by diagnostic ultrasound pulses to yield high echogenicity are small enough to pass through the circulatory system has allowed the creation of ultrasound contrast agents. However, the use of gas body contrast agents introduces a potential for microcavitation bioeffects into the clinical diagnostic setting. At low MIs, the stabilized microbubbles scatter ultrasound effectively, but the amplitudes of oscillation are too small to have significant effects on nearby cells. At higher MIs, the contrast agent microbubbles may be destabilized, allowing loss of the gas or nucleation of microcavitation activity.¹¹ This destabilization process can be useful for certain diagnostic procedures but can also damage nearby cells. The bioeffects possible with contrast agents and diagnostic ultrasound are sufficiently robust that several therapeutic applications are under study.¹²

The subject considered in this report is the potential for contrast-enhanced diagnostic ultrasound to induce cavitation-related bioeffects by ultrasound interaction with the gas bodies. This potential has been the subject of authoritative reviews previously.^{5,6} In addition, a symposium on the safe use of ultrasound contrast agents was conducted recently by the World Federation for Ultrasound in Medicine and Biology.¹³ This report arose from the 2005 American Institute of Ultrasound in Medicine Bioeffects Consensus Conference. Aspects of this subject considered below include (1) the nature of present contrast

agents, including diagnostic applications and ultrasound imaging modes; (2) the basic *in vitro* bioeffects of contrast gas body activation; and (3) reported *in vivo* bioeffects, including the therapeutic potential of contrast-enhanced diagnostic ultrasound. The text will only briefly revisit information included in the earlier conference report⁵ and will concentrate on bioeffects data relevant to diagnostic imaging, which have become available between 1998 and the 2005 conference. Summary conclusions are then presented with recommendations intended to optimize the safety profile of contrast-aided diagnostic ultrasound.

2. Contrast Agents for Diagnostic Ultrasound

2.1. Ultrasound Contrast Agents

First-generation contrast agents used air as the core gas. These air bubbles were surrounded by a fatty acid, lipid, or protein shell. The shell surrounding the air bubble increased the stability of the microbubbles both in the vial and in the body. Commercially available first-generation agents included Echovist (SH U 454; Schering AG, Berlin, Germany), Levovist (SH U 508A; Schering AG), both of which were only available in Europe, and Albutex (Molecular Biosystems, Inc, San Diego, CA), which was the first agent approved by the US Food and Drug Administration (FDA) for use in the United States.

One of the problems with the first-generation microbubbles was the short duration of efficacy after intravenous injection. These first-generation agents used air as the active component. However, because of an inherent unsaturation of dissolved gases (primarily oxygen, nitrogen, and carbon dioxide) in blood, the air contained within these contrast agents readily diffused out of the microspheres or microbubbles.¹⁴ With this loss of air, the first-generation contrast agents quickly lost the ability to produce ultrasound contrast. The next advancement made in ultrasound contrast agents (the second generation) was the inclusion of gases having decreased solubility and diffusivity. These microspheres or microbubbles retain their gas for a longer period of time; thus, the durations of contrast and Doppler enhancement increase from several seconds to several minutes.¹⁵

Optison (FS-069, perflutren protein type A microspheres for injection; GE Healthcare, Princeton, NJ) was the first second-generation ultrasound contrast agent approved by the FDA. This agent consists of a protenaceous shell surrounding a gas bubble of octafluoropropane gas.¹⁶ This relatively insoluble gas is inert and eliminated through normal gas exchange in the lung. Optison opacifies the cardiac ventricular chambers at doses much smaller than Albutex (0.2 mL compared with 15–20 mL, respectively) and opacifies the chamber for a much longer period of time than Albutex (>5 minutes versus 30–45 seconds).¹⁷ Currently, Optison is indicated to opacify the left ventricular chamber and to improve the delineation of the endocardial border.

Definity (MRX-115, perflutren lipid microspheres; Bristol-Myers Squibb Medical Imaging, North Billerica, MA) is another agent that encapsulates octafluoropropane, but this agent uses phospholipids to coat the microbubbles. Before injection, Definity must be activated by agitation in a “dental shaker” for approximately 45 seconds. Small volumes of these microbubbles opacify the left ventricular chamber and enhance the Doppler signal from the peripheral vasculature for prolonged periods of time.^{18,19} The microvascular rheology of Definity microspheres has been shown to be similar to that of erythrocytes in the circulation.²⁰ Currently in the United States, Definity is indicated for opacification of the left ventricular chamber and to improve the delineation the endocardial border. In Canada, Definity is indicated for both cardiology and radiology applications.

Imagent (AFO150, perflorane lipid microspheres; IMCOR Pharmaceuticals, San Diego, CA) is another ultrasound contrast agent that uses a perfluorocarbon to increase *in vivo* stability. This agent is composed of phospholipids, phosphate buffers, sodium chloride, and a blend of perfluorohexane and nitrogen.²¹ As with the other two approved agents, Imagent is indicated for opacification of the left ventricular chamber and to improve delineation of the endocardial borders.

Although not approved for sale in the United States, SonoVue (BR-1, sulfur hexafluoride; Bracco International BV, Amsterdam, the Netherlands) is also a second-generation ultrasound contrast agent approved for use in Europe. This agent is composed of phospholipids, ethylene glycol,

and sulfur hexafluoride gas.²² Originally, SonoVue was approved in March 2001 to increase the echogenicity of the blood, opacify the cardiac chambers, improve delineation of the endocardial borders, improve the Doppler signal-to-noise ratio in the cerebral, extracranial carotid, and peripheral arteries, and improve the visualization of the vascularity of liver and breast lesions (SonoVue package insert). In May 2004, the European Medicines Agency restricted the use of SonoVue to noncardiac imaging procedures.²³ This restriction was later removed, and a precautionary statement was issued stating that extra caution should be exercised in patients with conditions such as severe hypotension, bradycardia, cardiac arrest, and myocardial infarction.

The second-generation ultrasound contrast agents relied on core gases that possessed low solubility in blood to prolong their duration in the circulation. A third generation of contrast uses engineered changes in the microsphere shell to impart unique features to the contrast materials. Although these agents are still in development, two are in the later stages of their initial clinical development.

CARDIOSphere (PB127; Point Biomedical Corp, San Carlos, CA) is composed of a bilayer shell made up of polylactide and albumin. The polylactide inner layer is a biodegradable polymer that provides specific physical characteristics, which control acoustic properties of the contrast agent. The outer layer, composed of human albumin, functions as the biological interface and provides biocompatibility. These microspheres use nitrogen as the gas core. They were designed to collapse under very specific ultrasound conditions.²⁴ Once the microspheres are destroyed, the encapsulated nitrogen is released, and the gas quickly dissolves into the surrounding blood. This rapid loss of gas produces an intense signal using harmonic power Doppler imaging techniques.²⁵ Another third-generation agent, AI-700 (Acusphere, Inc, Watertown, MA), uses a synthetic porous microparticle to trap an insoluble gas. These microparticles are composed of D,L-lactide co-glycolide, a biodegradable polymer. These microparticles appear to be more resistant to the destructive effects of ultrasound.^{26,27}

Several companies continue to develop new ultrasound contrast agents with unique properties and niche applications. Their developmental

status, physical characteristics, and imaging profiles are less well known. A description of these various materials must wait until they move further along the developmental pathway. In addition, several other agents (eg, Ecogen, [Sonos Pharmaceuticals, Bothell, WA], Sonazoid [Nycomed, Zurich, Switzerland], Sonovist [Schering AG], and Quantison [Andaris Ltd, Nottingham, England]) failed to make it through the entire developmental process and gain regulatory approval for one reason or another. Although these agents did not make it to the clinical market, the information derived from their use both preclinically and clinically has contributed to a better overall understanding of ultrasound contrast agents, their effects and their applications.

2.2. Diagnostic Applications of Ultrasound Contrast Agents

Echocardiographic applications for Optison, Definity, and Imagent approved for use in the United States include left ventricular opacification and border delineation. Contrast echocardiography can also be used (but is not yet approved in the United States) for measurement of myocardial perfusion.²⁸ Levovist and SonoVue are used in other countries for cardiac, microvascular, and transcranial indications. Possibly the most prominent clinical application of contrast agents in Canada, Europe, and Asia is the detection and characterization of liver lesions. There are many other potential applications for imaging, including tissue perfusion, inflammation, and tumors, in any tissue accessible to ultrasound interrogation, including liver, kidney, breast, spleen, and others.²⁹⁻³¹ Ultrasound contrast agents are useful also for imaging body structure and function, such as for the diagnosis of vesicoureteral reflux^{32,33} and fallopian tubal patency.^{34,35}

2.3. Diagnostic Ultrasound Modes Used With Contrast Agents

Ultrasound imaging enhanced by the use of ultrasound contrast agents has been the subject of several recent reviews.³⁶⁻³⁸ Ultrasound contrast agents can be used most simply by bolus injection or infusion to enhance the echogenicity of blood in B-mode or Doppler images. High MI values, such as those used for harmonic imaging without contrast agents, result in the destruction

of the contrast agent and loss of contrast. This property of contrast agents has been used by intermittent imaging with agent infusion, which allows the tissue vasculature to refill with the contrast agent before the next image frame is processed (eg, see Porter et al³⁹ and Kuersten et al⁴⁰). This procedure can be used to image perfusion by varying the interframe trigger timing, as for myocardial contrast echocardiography (MCE).⁴¹ Doppler techniques are well suited to image microbubble destruction. Harmonic power Doppler imaging combined with some form of triggering has been used extensively to image microbubbles in the macrocirculation and microcirculation.

The increased sensitivity provided by newer imaging techniques makes it possible to image contrast microbubbles relatively nondestructively in real time at very low acoustic pressures. Low-MI imaging is important for two reasons. First, at a low MI, bubble destruction is avoided. Although microbubbles differ in their shell composition, work completed to date indicates that at an MI of about 0.15, the microbubbles examined are not significantly destroyed yet give a good harmonic (nonlinear) contrast signal.³⁷ The second major reason for low-MI scanning is the reduction of the harmonic component in the tissue echoes relative to bubble echoes. While tissue harmonics have benefited routine diagnostic scanning, it is the background “noise” signal that the contrast signal must rise above. Because tissue is less nonlinear than bubbles, it requires a higher MI than the contrast microbubbles for a certain harmonic response. Therefore, at a low MI, the contrast-to-tissue ratio is higher than at a high MI, helping remove the tissue signal and leave only the contrast. For quantification purposes, both high and low MIs may be combined, in which a high-MI pulse is sent to destroy the contrast microbubbles in a scan plane, and then a low MI is used afterward to monitor the contrast replenishment.

3. Ultrasound Interaction With Contrast Agent Gas Bodies: Physical Theory

3.1. Pulsation

The physical interaction between ultrasound pulses and contrast agent gas bodies produces microbubble pulsations, which are responsible for their high echogenicity but which also increase

the potential for local bioeffects. Theory has been developed for the interaction of ultrasound with encapsulated gas bubbles in contrast agents such as Optison.^{42,43} The theory is similar to that for a free cavitation microbubble but with added elasticity and viscosity-related damping parameters to account for the shell. This theory has been used to describe the scattering properties of suspensions of the agents, leading to empirical values for the shell parameters.⁴⁴ The theories appear to be reasonable models of the gas body behavior for low levels of excitation. In addition, the presence of blood cells around the contrast agent gas bodies has a relatively small effect on the theoretical dynamics of the pulsation.⁴⁵

3.2. Influence of Contrast Agents on Tissue Heating

Although thermal effects of ultrasound are treated in other articles,^{46,47} the presence of contrast agents may have an effect on tissue heating. The theory for this effect has been advanced⁴⁸ and tested in nonbiological materials such as tissue-mimicking phantoms.⁴⁹ Because of the ability of bubbles to oscillate nonlinearly and produce higher-frequency components than are present in the insonifying beam, acoustic energy can be dissipated much more effectively as heat. Moreover, the effects of fluid viscosity near oscillating bubbles are enhanced because of the large component of acoustic radiation (especially for bubbles above the resonance size⁵⁰). Numerical modeling of the theory agrees reasonably well with *in vitro* experiments.^{51–53} In addition, Stride and Saffari⁵⁴ have noted that the viscous properties of the stabilizing shells could enhance local heating near the gas bodies.

Measurements of excess heating of tissue due to the presence of echo contrast agents fall mostly in the insonification regimen of therapeutic ultrasound. Fujishiro et al⁵⁵ insonified beef samples using 1.5-MHz continuous ultrasound at 0.9 W/cm² for 3 minutes and found an equivalent temperature rise to doubling the intensity without contrast agents present. Wu⁵⁶ found excess temperature elevation of several degrees Celsius due to the presence of contrast agents in suspensions insonified at 3.5 W/cm² and 1 MHz. Sokka et al⁵⁷ monitored lesion formation in rabbit thighs after bubbles were detected following a 7-W con-

tinuous insonation at 1.7 MHz for 20 seconds. The lesions were larger in volume by up to a factor of 3 for the bubble-enhanced exposures.

3.3. Cavitation Nucleation

At low pressure amplitudes, the contrast agent gas bodies may remain stable; that is, their shells may remain intact as the gas body oscillates. However, destabilization of gas bodies appears to occur at modest peak rarefactional pressure amplitudes (PRPAs), particularly for shell-encapsulated designs. At relatively high pressure amplitudes, the stabilized gas bodies are destroyed but can serve as cavitation nuclei. As the incident ultrasound pressure amplitude increases from 0, the stabilizing shell experiences oscillating stresses. Above rather modest excitation PRPAs, the theoretical stresses may be sufficient to expect shell failure.⁵⁸ From *in vitro* bioeffects research on contrast agents, it is evident that bioeffects are often associated with gas body destabilization, and that observation of the loss of gas bodies appears to agree with the theory for expected shell failure as a function of frequency.⁵⁹ Uncertainty remains as to how the theories can be used to describe the destabilization of the gas bodies and the complex behavior at higher amplitudes. The fate of destabilized gas bodies can include gradual shrinkage or rapid fragmentation, depending on physical conditions.^{60,61}

Nucleation of cavitation involves a transition from restricted pulsation of gas bodies to the free pulsation of cavitation microbubbles. Cavitation nucleation by ultrasound contrast agents is important with regard to the bioeffects potential *in vivo* because there normally are few, if any, cavitation nuclei in the body suitable for direct activation by diagnostic ultrasound pulses.⁶² Ultrasound contrast agents can supply such nuclei. For example, direct evidence of cavitation activity has been obtained by detection of broadband noise emissions from the myocardium during contrast echocardiography.⁶³

At relatively modest PRPAs, the pulsation of free microbubbles results in collapse driven by the inertia of the surrounding liquid during the compression phase of the oscillation. This phenomenon defines inertial cavitation, which is strongly associated with many bioeffects. These

bioeffects are caused by fluid jets, extreme heating, and free radicals generated on collapse.⁵ The threshold for inertial cavitation derived for optimum-sized nuclei was the basis for the MI. By assuming the presence of these optimally sized nuclei, the inertial cavitation threshold p for blood at frequency f was found to fit the formula⁹

$$\frac{p^{1.67}}{f} = 0.13.$$

The square root of this formula approximates the form of the MI (ie, the PRPA in MPa divided by the square root of the frequency in megahertz). The minimum inertial cavitation threshold in blood can therefore be expressed approximately in terms of the MI as $MI_1 = 0.4$. As noted below in section 5, several *in vivo* bioeffects associated with contrast-aided diagnostic ultrasound have been reported at and above this MI value, although the frequency dependence of ultrasound bioeffects thresholds in the presence of an echo contrast agent may differ from that suggested by the MI (see Figure 1).

3.4. Shear Stress Theory for Bioeffects

Bioeffects induced by ultrasound interaction with contrast agents are amenable to theoretical consideration. However, detailed theory for estimating the amount of biological perturbation expected from a given exposure situation is not presently available for use in medical applications of contrast agents. Even for moderate pressure amplitudes, a number of different mechanisms may plausibly have a role in bioeffects.⁶⁴ Large oscillatory and steady fluid shear stresses occur for gas body pulsation near solid surfaces. The contrast agent gas bodies can destabilize and nucleate inertial cavitation. Cavitation, in the absence of contrast agents, is rare in most tissues and is discussed in other conference reports.^{65,66} However, the use of ultrasound contrast agents introduces the potential for cavitation bioeffects into the risk/benefit equation for diagnostic ultrasound.

For low PRPAs and simple *in vitro* conditions, cellular bioeffects have been modeled theoretically with some success. Fluid shear stress generated in the vicinity of a pulsating gas body can produce mechanical membrane damage.^{67,68} The microstreaming shear stress model has proven useful in describing several bioeffects situations

involving gas body activation in terms of the damage of cell membranes by shear stress in acoustic microstreaming fluid flow near the oscillating gas body.⁶⁹ Approximate theory is available for the nonoscillatory steady shear stress generated in near-boundary acoustic microstreaming and can be used to estimate the pressure amplitudes required to exceed the critical shear stress for biological membranes. The average steady stress can also persist with a time-average value given by the peak stress times the fractional duty cycle. Miller⁷⁰ considered this theoretical model with respect to ultrasound contrast agents. Significant shear stress was found to be possible for relatively low pressure amplitudes, particularly for the case of free microbubbles. The pressure amplitude for which the shear stress was expected to exceed critical levels increased approximately in proportion to frequency for encapsulated gas bodies. Wu⁷¹ used the microstreaming shear stress theory to model the effects of contrast agent gas bodies attached to cells when exposed to ultrasound. Shear stresses generated by 1- to 2-MHz ultrasound were found to be sufficient for sonoporation (transient permeabilization with resealing) of cells at a pressure amplitude of only 0.12 MPa. Destabilization with liberation of free bubbles was shown to produce a much higher shear stress with potential cell killing. The shear stress model was applied to a specific experimental system for which contrast agent gas bodies were exposed to pulsed ultrasound while attached to monolayer cells.^{72,73} The observed PRPA thresholds p for destabilization at different ultrasound frequencies f correspond to approximately constant relative pulsation amplitudes (amplitude divided by the initial radius). The constant relative amplitudes theoretically yield approximately constant shell stresses, which were sufficient to induce destabilization. The observed cell death thresholds corresponded theoretically to approximately constant radial velocity amplitudes. The constant velocity amplitudes theoretically yield slowly increasing microstreaming shear stress for 1.8- to 0.2-microsecond pulse durations in the 1- to 10-MHz range. For this model system, contrast agent gas body destabilization and pulsation-induced cell membrane damage was expected to occur for similar critical values of the parameter p/f . The comparison to in vitro bioeffects is noted below in section 4.2.

4. Basic In Vitro Studies of Bioeffects Potential

In vitro experimental models can provide insights into the fundamental processes and minimum ultrasound exposures needed for bioeffects induced by the interaction of ultrasound with contrast agents. There are two qualitatively different situations for cultured cells: suspensions and monolayers. Selected PRPA values for thresholds of in vitro bioeffects are plotted as a function of frequency in Figure 1.

4.1. Effects Produced in Cell Suspensions

Sonolysis of red blood cell suspensions (or hemolysis) containing contrast agent gas bodies has been studied extensively.⁵ The frequency dependence of thresholds for hemolysis in whole blood with added Alunex in a stationary chamber was determined by Miller et al⁷⁴ for pulsed (10-microsecond pulses) ultrasound (Figure 1). The threshold at 2.4 MHz with Optison was found to be somewhat lower, presumably due to the greater presence of the microbubbles.⁷⁵

A correlation between contrast-aided-ultrasound induced hemolysis in a rotating chamber and the amount of passively detected inertial cavitation activity (an inertial cavitation “dose”) has been established.^{78–81} This correlation is so robust that hemolysis can be used as a cavitation dosimeter.^{81,82} In blood containing ultrasound contrast agent gas bodies, exposure to 1-MHz ultrasound with a 2.0-MPa PRPA produces detectable hemolysis and inertial cavitation dose with pulses as brief as 2 cycles.⁸⁰ The inertial cavitation dose and hemolysis evolve and reach limiting values very rapidly in vitro as microbubbles are destroyed. Both end points are influenced strongly by the PRPA, with thresholds at 1 MHz of approximately 0.5 to 1 MPa.⁸¹ Hemolysis induced by inclusion of contrast agents in ultrasound-exposed samples has a very strong frequency dependence at suprathreshold PRPAs.^{79,82,83} That is, the PRPA to produce a given level of hemolysis increases with frequency much faster than the $f^{0.5}$ frequency dependence of the MI. The MI, therefore, has poor predictive value for the magnitude of observed hemolysis above the threshold.

Suspended phagocytic cells, which have bound or phagocytosed contrast agent gas bodies, are lysed when exposed to megahertz-frequency acoustic pressures within the capabilities of diagnostic imaging machines.^{84,85} When the expansion phase of the phagocytosed bubble oscillation exceeded the cell's ability to expand, cell membrane damage was observed. For 3- μm -diameter gas bodies and a 2.25-MHz frequency, a 0.9-MPa pulse did not appear to damage the membrane, while a 1.6-MPa pulse resulted in membrane rupture.

Exposure of whole blood to 3.5-MHz ultrasound produced by a diagnostic ultrasound scanner (MI = 1.9) induced platelet activation at a high gas body concentration of Levovist.⁸⁶

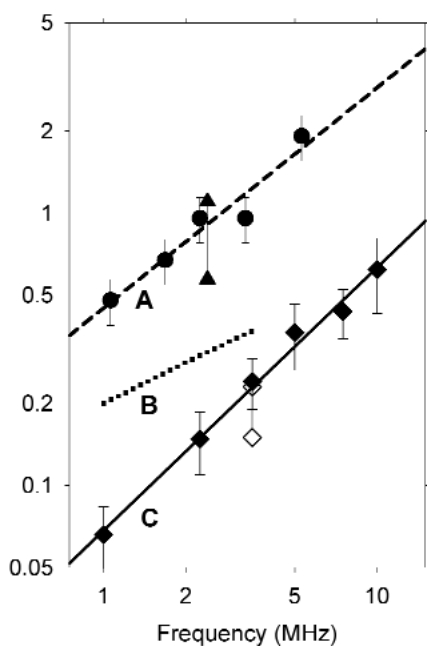
Killing of lymphocytes cosuspended with contrast agent gas bodies in vitro results from exposure to low-amplitude (0.2-MPa PRPA, 20/180 cycles on/off) 2-MHz ultrasound in the rotating tube system.⁸⁷

4.2. In Vitro Studies of Bioeffects on Cell Monolayers

Miller and Bao⁸⁸ studied cell killing in Chinese hamster ovary monolayers insonified in the presence of Albunex. Monolayers were exposed to 10-microsecond pulses of 3.3-MHz ultrasound while "inverted" (ie, the monolayer was located at the top of the vessel during exposure so that the gas bodies would rise to become adjacent to the monolayer). This model system was designed to maximize the potential for cellular effects and may simulate the concentration of gas bodies at a distal vessel wall by acoustic radiation forces.⁸⁹ A second transducer was used to detect subharmonic emissions associated with oscillating bubbles. Cell killing and bubble acoustic emissions were correlated with the PRPA, with a threshold of 0.56 MPa. Similar experiments using Optison and 3.5-MHz diagnostic ultrasound showed epidermoid cell sonoporation to be strongly dependent on gas body concentration.⁹⁰ Sonoporation was detected at PRPAs as low as 0.23 MPa in the pulsed Doppler mode and 0.39 MPa in the B-mode. Experiments to characterize gene transfer and cell killing were conducted with similar inverted epidermoid cell monolayers and used a diagnostic scanner operating in the harmonic mode (1.5-MHz transmit frequency) to produce the ultrasound exposures. With 2% Optison, the PRPA threshold for cell killing was less than 0.7 MPa, and gene transfer was detected above 1.7 MPa.⁹¹

Brayman et al⁷⁶ modeled the endothelial layer of blood vessels with V79 fibroblast monolayers whose orientation was varied to simulate either the proximal or distal walls of blood vessels scanned by ultrasound. Contrast agents increased damage to both distal and proximal monolayers at 1.0-, 2.1-, and 3.5-MHz frequencies with a sharp frequency dependence (Figure 1). Distal monolayers were damaged more than proximal monolayers. Kudo et al⁹² used a high-speed camera to directly observe ultrasound-induced (1 MHz, 0.6-MPa PRPA) oscillations of

Figure 1. Critical (apparent threshold) measurements of the PRPA for cell damage by pulsed ultrasound interaction with contrast agent gas bodies in in vitro test systems. Circles⁷⁴ are fitted by line A for hemolysis in a whole-blood suspension with Albunex. Triangles are for hemolysis with an Albunex (top) or Optison (bottom) suspension in whole blood.⁷⁵ Line B is for erosion of endothelial-like cell monolayers by pulsed ultrasound interaction with Albunex suspended in the medium.⁷⁶ Filled diamonds⁷² are fitted by line C for killing of phagocytic monolayer cells with Optison gas bodies attached to the cells. Open diamonds represent apparent thresholds for cell killing (top) and sonoporation (bottom) of epidermoid cells with Optison gas bodies allowed to rise and contact the monolayer cells.⁷⁷ Reproduced with permission from Miller D. Overview of experimental studies of biological effects of medical ultrasound caused by gas body activation and inertial cavitation. *Prog Biophys Mol Biol* 2007; 93:314–330.



contrast agent gas bodies adjacent to bovine endothelial cell monolayers. Obvious cell shape distortions and killing were associated with non-spherical bubble collapse.

Phagocytic cell monolayers (RAW-264.7) preincubated with Optison and then washed to remove unbound gas bodies were killed by exposure to ultrasound produced by a diagnostic ultrasound machine operated in the spectral Doppler mode, with a PRPA threshold of approximately 0.2 MPa.⁷⁷ Similar experiments using Optison showed the PRPA threshold to be 0.8 MPa for exposures consisting of a single 2-cycle pulse.⁷² Using 2-cycle ultrasound pulses, the PRPA thresholds for killing RAW-264.7 cells preloaded with Optison showed a linear correlation ($r^2 = 0.982$) with frequencies over the 1- to 10-MHz range (Figure 1), increasing with a slope of approximately 0.06 MPa/MHz.⁷³ The authors note that these pressure thresholds are lower than those for nucleation of inertial cavitation and have a markedly different frequency dependence. As noted above in section 3, the frequency dependence of gas body destabilization and cellular bioeffects observed for this *in vitro* system can be modeled by the theory for shell stresses and acoustic microstreaming shear stress on cells.⁵⁹ The theory substantiated the observed linear dependence of thresholds on frequency. Owing to the design of this model monolayer system for maximum sensitivity for cellular bioeffects, the thresholds observed may approximate the lowest PRPAs for which biologically significant bioeffects (ie, cell killing) can be expected for contrast-aided pulsed ultrasound.

5. In Vivo Studies of Bioeffects

In this section, available reports on *in vivo* bioeffects associated with contrast ultrasound will be reviewed. Some reports of bioeffects at high pulse amplitudes, which did not involve actual diagnostic ultrasound or pulsed ultrasound intended to simulate diagnostic ultrasound, are briefly noted to indicate the types of bioeffects that might occur above the guideline upper limits for diagnostic ultrasound. Results of clinical research conducted during the approval process are generally not available for review. In addition, research on possible pharmacologic side effects

(ie, without ultrasound interaction) of the contrast agents are not reviewed here. No epidemiologic studies are available on possible adverse (or favorable) health effects of the use of ultrasound contrast agents. Most *in vivo* research has been conducted using mice or rats. Reports have centered mostly on skeletal or cardiac muscle, as listed in Table 1. Several studies on other tissues are listed in Table 2. Research on the possible therapeutic use of diagnostic ultrasound aided by gas bodies is also reviewed briefly.

5.1. Skeletal Muscle and Myocardium

The behavior and resulting bioeffects of contrast agent gas body destruction by diagnostic ultrasound were observed by intravital microscopy of the spinotrapezius muscle in rats.⁹³ The muscle was positioned in a custom-built chamber filled with Ringer's solution containing adenosine for vascular dilation and propidium iodide to stain nuclei of dead cells. A phased array diagnostic ultrasound system was used in the harmonic mode at 2.3 MHz to image the muscle. Optison with fluorescently labeled gas bodies was infused into the femoral vein for 1 minute before obtaining a single image frame at a specific MI, with MIs of 0.4, 0.5, 0.7, and 1.0 used for each animal. After exposure, the muscle was examined for microvessel rupture and dead (stained) cells. The number of capillary rupture sites and stained cells was near 0 at an MI of about 0.4 and increased rapidly for the higher MI values.

The induction of petechiae by contrast-aided diagnostic ultrasound was confirmed for skeletal muscle by Miller and Quddus.⁹⁴ A 2.5-MHz probe was directed at the abdomens of anesthetized mice mounted in a water bath to provide for free-field exposure conditions. A tissue-mimicking phantom was used to simulate intervening tissue. Evans blue dye, used to indicate microvascular leakage, and Optison were introduced by retro-orbital injection. After imaging, the abdominal muscle and intestines were examined for microvascular injury. For 10 periods with 10 seconds on and 10 seconds off with 5-mL/kg Optison, petechiae counts in the tissue were significantly elevated relative to shams at PRPAs above 0.64 MPa (measured equivalent MI = 0.4) and were proportional to the square of the PRPA. A single image frame was sufficient to pro-

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duce petechiae. The petechiae number was approximately proportional to the contrast agent dose. Evans blue leakage was evident, and excess dye could be extracted from tissue within the

scan plane relative to sham samples. Miller and Quddus⁹⁴ also reported capillary rupture in fat, small intestine, and Peyer patches (intestinal lymph nodes).

Table 1. Bioeffects Induced by Contrast-Aided Diagnostic Ultrasound in Skeletal and Cardiac Muscle

Reference	Frequency, MHz	Mode	Agent	Animal Tissue	Effect	Critical MI
Skyba et al ⁹³	2.3	B-mode	Optison	Rat spinotrapezius	Microvessel rupture, cell killing	≈0.4
Miller and Quddus ⁹⁴	2.5	B-mode	Optison	Mouse abdominal	Petechiae, capillary leakage	0.4
van der Wouw et al ⁹⁵	1.66	Triggered B-mode	AIP101	Human heart	PVCs	1.1–1.5
Ay et al ⁹⁶	1.8	Triggered B-mode	PESDA, Sonazoid, Optison, Levovist	Rabbit isolated heart	Function, lactate, capillary rupture	1.0
Chen et al ⁹⁷	1.3	Triggered B-mode	Optison, Definity	Rat heart	Troponin T elevation, negative histologic findings	≈1.2
Borges et al ⁹⁸	1.7–1.9	Triggered B-mode	Optison	Human heart	Negative for PVCs, troponin I, CK, CK-MB	1.4–1.7 only
Raisinghani et al ⁹⁹	NS	Triggered Doppler	PB127	Human heart	Negative for PVCs	<1.0
Li et al ¹⁰⁰	1.7	Triggered B-mode	Optison	Rat heart	PVCs	0.77
					Petechiae, leakage	0.41
Kobayashi et al ¹⁰¹	1.8	Harmonic B-mode	Definity	Rat heart	Negative for petechiae	≈0.61 only
Miller et al ¹⁰²	1.7	Triggered B-mode	Optison	Rat heart	Microvascular leakage <20 min	1.5 only
Li et al ¹⁰³	1.7	Triggered B-mode	Optison, Imagent, Definity	Rat heart	PVCs	0.77
					Petechiae	0.31
Miller et al ¹⁰⁴	1.5	Triggered B-mode	Optison	Rat heart	Histologic microlesions	1.7 only
Chapman et al ¹⁰⁵	1.7	Triggered B-mode	PESDA	Human heart	PVCs, arrhythmia	<0.8

NS, not specified.

Table 2. Bioeffects of Diagnostic or Pulsed Ultrasound With Contrast Agents in Various Tissues Other Than Muscle

Reference	Frequency, MHz	Mode	Agent	Animal Tissue	Effect	Critical MI
Miller and Gies ^{106,107}	1.09	10-μs pulses	Albunex, Levovist	Mouse intestine	Petechiae	0.81
	2.3		Optison, PESDA			1.2
Kobayashi et al ¹⁰⁸	1.8	Harmonic B-mode	Levovist	Intravital rat mesentery	Endothelial cell killing	≈0.61 only
Schlachetzki et al ¹⁰⁹	2.0–3.5	Transcranial color duplex	Levovist, Optison	Human brain	Negative for magnetic resonance contrast leakage	<1.9
Wible et al ¹¹⁰	1.8	Continuous or triggered B-mode	Optison, MP1950	Rat kidney	Glomerulus, hemorrhage	0.94
	4.0		MP2211			1.0
Kobayashi et al ¹⁰¹	1.8	Harmonic B-mode	Definity, Levovist	Intravital rat mesentery	Endothelial cell killing, hemorrhage	≈0.1
Shigeta et al ⁸⁶	8	B-mode	Levovist	Rat liver	Platelet aggregation, endothelial injury	≈1.8 only
	12					≈0.7 only
O'Brien et al ¹¹¹	3.1	1.2-μs pulses	Optison	Rat lung	Negative for enhanced hemorrhage	1.6, 3.3
Miller and Dou ¹¹²	1.5	Harmonic B-mode	Definity	Mouse tumor	Negative for enhanced metastasis	1.9 only

*The different MI values correspond to the different frequencies.

Increased numbers of premature ventricular contractions (PVCs) were reported in humans undergoing contrast echocardiography by van der Wouw et al.⁹⁵ The contrast agent AIP101 (not commercially available) was infused intravenously into healthy volunteers. Cardiac scans were conducted at 1.66 MHz with intermittent image frames, which allowed refill of the tissue with the contrast agent between frames, and MI values of 1.1 and 1.5. A significant increase in PVCs to about 1 per minute was seen for end-systolic triggering at an MI of 1.5 but not at an MI of 1.1.

The microvascular effects of contrast echocardiography were examined in isolated rabbit hearts by Ay et al.⁹⁶ A cardiac ultrasound system was operated at 1.8 MHz with 1-Hz triggering of image frames using the machine display MI as a measure of the exposure. The perfusate contained the laboratory-made perfluorocarbon-exposed sonicated dextrose albumin (PESDA) contrast agent (effects were also confirmed with commercial agents); however, it is difficult to relate the dosage to an *in vivo* intravenous dose. Scanning at an MI of 1.6 led to a transient decrease in left ventricular pressure. Lactate release was significant at MIs of 1.0 and higher. Light microscopy revealed capillary damage and erythrocyte extravasation.

Potential injurious effects associated with the use of Optison or Definity with diagnostic ultrasound exposure were examined for *in vivo* rat hearts.⁹⁷ Imaging was performed at 1.3 MHz with electrocardiographic (ECG) triggering at every 4 cardiac cycles. The contrast dosage was 0.1 mL in saline delivered by infusion into the jugular vein. Left ventricular function was not perturbed (maximum MI = 1.6). Elevations in troponin T in blood plasma, indicating myocardial damage, were detected after 30 minutes for MIs of 1.2 and 1.6. The elevation was significant at an MI of 1.6 for both agents, declining to normal by day 4. Histologic examination of scanned tissue obtained on day 7 did not show evidence of necrosis, vascular damage, or inflammation.

A clinical study was reported for humans undergoing MCE by Borges et al.⁹⁸ A dose of 3 mL of Optison in saline was injected as a bolus into the cubital vein over 3 minutes followed by a saline flush. A diagnostic scanner was used in the

harmonic imaging plus power Doppler imaging duplex mode at 1.7 to 1.9 MHz for MIs of 1.4 to 1.7 and with end-systolic triggering at every 1 to 3 beats. Blood samples were taken before and up to 24 hours after scanning and were analyzed for myoglobin, troponin I, creatine kinase (CK), and CK isoenzyme MB. One patient had a transient increase in troponin I after the examination, but there were no consistent changes in measured parameters that could be related to the ultrasound examinations.

A large group (135) of humans was studied with regard to induction of PVCs during MCE using a new contrast agent, PB127.⁹⁹ Several cardiac ultrasound machines were used in the dual-frame triggering mode with MI settings of 0.9 to 1.0. One group of patients also had dipyridamole infusion for a stress test. Premature ventricular contractions were observed in the patient group but were not associated with the frame triggers. There was no significant increase in PVC frequency during or after imaging. The negative result was reassuring, but higher MI values would have been needed for direct comparison with the results of van der Wouw et al.⁹⁵

An *in vivo* rat model of MCE was used to examine microvascular permeabilization and PVCs with respect to the method of imaging, ultrasound exposure, and agent dose.¹⁰⁰ A 1.7-MHz diagnostic ultrasound system was used to scan the rats in a water bath for 3 minutes. Evans blue dye, a marker for microvascular leakage, and a bolus of Optison were injected intravenously. Neither PVC nor microvascular leakage was seen in controls, rats imaged without the injected contrast agent, or rats with the injected contrast agent but not imaged. Triggering 1:4 at end systole produced the most PVCs, petechiae, and microvascular leakage, followed by end systole 1:1 triggering, continuous scanning, and end diastole 1:1 triggering. All effects increased with increasing Optison doses over the range of 25 to 500 $\mu\text{L}/\text{kg}$. Threshold PRPAs above which effects were significant were 1.0 MPa for PVCs and 0.54 MPa for microvascular leakage.

The possible occurrence of microvascular injury to rat hearts was also examined by Kobayashi et al.¹⁰¹ A phased array ultrasound system was used at 1.8 MHz and a 4-cm focus with frame rates and on-screen MI combina-

tions of 1.6 and 1 Hz, 0.2 and 30 Hz, and 1.6 and 30 Hz. After scanning of the rat mesentery, described below, the probe was moved to the chest wall for 3 minutes of exposure while Definity at 0.1 or 1.0 mL/kg was administered. This arrangement may have placed the rat heart in the near field of the array (ie, at lower PRPAs than implied by the on-screen MI), but the heart images were monitored on the ultrasound screen. After scanning, the hearts were fixed for histologic examination. No hemorrhages were found in the rat heart sections.

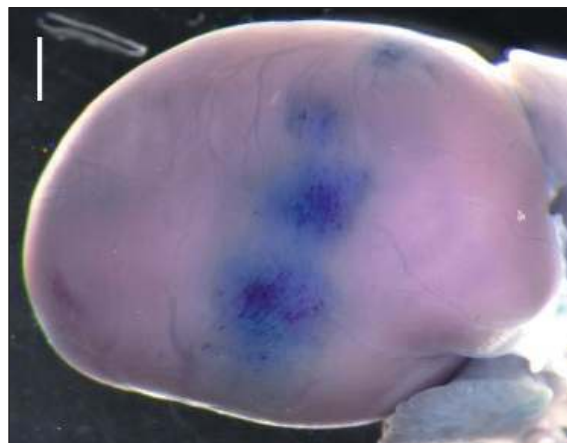
The timing and influence of vasoactive drugs on microvascular leakage induced by MCE in rats were investigated by Miller et al.¹⁰² Hairless rats were anesthetized and transthoracically scanned with a diagnostic ultrasound system (GE VingMed System V) at 1.7 MHz with 1:4 triggered frames at end systole in a water bath with the heart at the focal position. Optison, vasoactive medications, and Evans blue dye were injected via the tail vein. Effects were similar to those in the previous study.¹⁰⁰ Propranolol and isoproterenol had little effect on the microvascular leakage, which suggests that the microvascular leakage was primarily a mechanical effect rather than a physiologic response. Capillary leakage occurred during and after exposure but diminished for Evans blue injections administered 20 minutes after scanning.

The effects of PVCs, petechiae, and microvascular leakage reported by Li et al¹⁰⁰ were compared for Optison, Definity, and Imagent.¹⁰³ On the basis of the volume dose, MCE using Definity produced more microvascular leakage. An example of the petechial hemorrhages and microvascular leakage seen in rat hearts after Definity MCE is shown in Figure 2. However, when expressed in terms of the number of gas bodies, there was no apparent difference between the three agents' microvascular damage potential, which increased linearly with the gas body dose at low doses, as shown in Figure 3. Myocardial contrast echocardiography using Definity resulted in fewer PVCs than the other agents. The effects increased strongly with the PRPA, with calculated thresholds for petechiae at about 0.4 MPa (MI = 0.31) and for PVCs at about 1.0 MPa (MI = 0.77). There was no apparent threshold for the visual detection of an Evans blue leakage area on the heart surface, which was significant for the lowest exposure of 0.54 MPa (MI = 0.41).

Histologically defined microlesions with inflammatory cell infiltration induced by MCE were reported by Miller et al.¹⁰⁴ Myocardial contrast echocardiography with 1:4 end-systolic triggering was performed in rats at 1.5 MHz and an MI of 1.7 in a short-axis view of the left ventricle in rats. Two high doses (500 μ L/kg) of Optison were given 5 minutes apart during 10 minutes of echocardiography. In rats killed 10 minutes after MCE, microvascular leakage and petechiae were evident. After 24 hours, microlesions with inflammatory infiltrates were scattered primarily over the anterior half of the sections. Lesion areas in the anterior wall were scored from photomicrographs, and there was inflammatory cell infiltration in areas of $0.5\% \pm 0.8\%$ (SD) for shams and $7.4\% \pm 5.0\%$ for MCE ($P < .02$). For rats killed 1 and 6 weeks after MCE, the microlesions healed to form small fibrous regions interspersed with normal myocytes.

Dalecki et al¹¹³ examined the induction of premature contractions in mice by 1.2-MHz pulsed ultrasound with Alunex or Optison in the circulation. Pulses were triggered during diastole from the ECG. Using 5-millisecond pulses, the threshold for premature contractions was low-

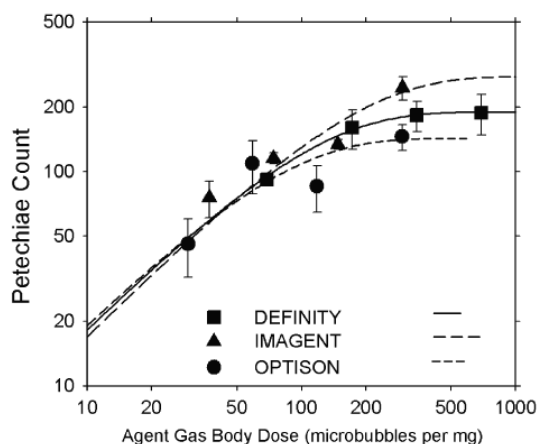
Figure 2. Microvascular leakage of Evans blue dye and petechial hemorrhages in a rat heart model of MCE.¹⁰³ Diagnostic B-mode ultrasound at 1.7 MHz with an in situ PRPA of 1.9 MPa (MI \approx 1.5) was used to image the heart using frame triggering at end-systole at each fourth heartbeat. A bolus of Definity at the recommended dose of 10 μ L/kg was injected into the tail vein with scanning continuing for 5 minutes after contrast appeared in the heart. Scale bar indicates 2 mm. Reproduced with permission from Miller D. Overview of experimental studies of biological effects of medical ultrasound caused by gas body activation and inertial cavitation. *Prog Biophys Mol Biol* 2007; 93:314–330.



ered by a factor of 10 below a previously determined threshold without the contrast agent to about 0.2 MPa, which suggests that cavitation was responsible for the bioeffect. For 10-microsecond pulses, the threshold was about 1 MPa. The thresholds were similar for the two different agents.

The occurrence of arrhythmias, such as PVCs, was examined and compared for physical therapy ultrasound and for triggered diagnostic ultrasound in humans.¹⁰⁵ The therapeutic mode involved an unfocused 1-MHz ultrasound beam with continuous or burst mode operation at measured PRPAs up to only 0.39 MPa. The diagnostic ultrasound machine was operated at 1.7 MHz with frames triggered at every 4 cardiac cycles and had a measured PRPA of 1.0 MPa for a 1.3-MI setting on the machine. The therapy device produced significantly more arrhythmia than the diagnostic imager for the same transthoracic exposure windows for the heart, which indicates that factors other than PRPA, such as a continuous versus pulsed mode, are important for this bioeffect. The low numbers of PVCs induced by the diagnostic ultrasound with a moderate PRPA,

Figure 3. Comparison of the counts of petechiae seen on the rat heart surface after MCE at 1.7 MHz, 1:4 end-systolic triggering, and a PRPA of 1.9 MPa. The doses of the three agents are compared on the basis of the numbers of gas bodies in the bolus injections. The curves are simple exponential functions with linear dependence at low doses and saturation at higher doses. Reproduced with permission from Li P, Armstrong WR, Miller DL. Impact of myocardial contrast echocardiography on vascular permeability: comparison of three different contrast agents. *Ultrasound Med Biol* 2004; 30:83–91.



$$\frac{P}{\sqrt{f}} < 0.8,$$

was consistent with other observations in humans⁹⁵ and also in rats.¹⁰³

5.2. Bioeffects on Other Tissues

The intestine has been of interest with regard to potential nonthermal bioeffects, owing to its natural content of free gas bubbles. Miller and Gies^{106,107} used pulsed ultrasound to simulate diagnostic ultrasound (10-microsecond pulses and a 1-kHz pulse repetition frequency for 100 seconds) to expose mouse intestines and search for petechiae in the intestinal wall. Contrast agents were introduced by retro-orbital injection. Peak rarefactional pressure amplitude thresholds for induction of petechiae in the presence of 10-mL/kg Albunex increased from 0.85 MPa at 1.09 MHz to 2.3 MPa at 2.4 MHz. Levovist, Optison, and PESDA all yielded more petechiae than Albunex at 2.3 MHz, and thresholds were as low as 1.8 MPa (MI \approx 1.2) for Levovist. Owing to the continuous exposure in the near field of unfocused transducers, the apparent thresholds are not comparable to intermittent diagnostic scanning.

The occurrence of microvascular injury to the rat mesentery was examined by Kobayashi et al¹⁰⁸ using diagnostic ultrasound in an intravital preparation. A phased array ultrasound system was used at 1.8 MHz and a 4-cm focus with frame rates and MI combinations of 1.6 and 1 Hz, 0.2 and 30 Hz, and 1.6 and 30 Hz. Field measurements indicated a PRPA of 0.82 MPa (MI \approx 0.61). Levovist was injected via the femoral vein. Propidium iodide was used to stain venule and capillary endothelial cells killed by contrast-aided scanning. Capillary ruptures were rare. This same system was used to compare Levovist with Definity for induction of endothelial cell injury and microvessel bleeding.¹⁰¹ The system was adjusted for a measured PRPA value of either 0.14 MPa (MI \approx 0.1) or 0.82 MPa (MI \approx 0.61). No microvessel injury was seen for either ultrasound alone or the contrast agent alone. Microvessel bleeding was rare and only seen for the 30-Hz frame rate at the higher exposure. Endothelial cell damage was observed for all conditions at the higher exposure. Significant cell killing was produced using the low exposure and 30-Hz

frame rate with 1.0-mL/kg Definity in capillaries and venules (but not arterioles).

The possible alteration of the blood-brain barrier by contrast-aided ultrasound was investigated by Schlachetzki et al.¹⁰⁹ Transcranial color-coded sonography was performed on human volunteers with a 2- to 3.5-MHz phased array probe with maximal output settings. Frames were triggered from the ECG at each heartbeat, with speckling in the color Doppler images indicating microbubble destruction. Levovist and Optison were used for the contrast ultrasound, and the magnetic resonance contrast agent Magnevist (Bayer HealthCare Pharmaceuticals, Leverkusen, Germany) was also injected intravenously. Evidence of microvascular leakage was sought using magnetic resonance imaging of the brain. There were no indications of focal signal enhancement attributable to extravasation of the Magnevist.

Wible et al.¹¹⁰ studied renal capillary hemorrhage induced by contrast ultrasound in rats. A diagnostic ultrasound probe was placed 1 cm from the kidney with one kidney exposed at a 30-Hz frame rate and the other at a 1-Hz frame rate. Frequencies of 1.8, 4, and 6 MHz were used at displayed MI values of 0.4 to 1.6, with the actual PRPA values measured at 1.5 cm. The contrast agents MP1950, MP2211, and Optison were administered at a dose of 40 million microbubbles/kg. Contrast-aided ultrasound caused areas of small hemorrhages visible on the kidney surface within the scanned plane. The small hemorrhages involved escape of red blood cells from the glomerular tuft into the Bowman capsule and proximal convoluted tubules. Intermittent exposure was more effective at producing the small hemorrhages than continuous scanning and gave a significant increase in renal hemorrhages for a PRPA of 1.26 (MI = 0.94). The severity decreased with increasing ultrasound frequencies but was significant at a PRPA of 2.02 at 4 MHz (MI = 1.0). At 6 MHz, no significant hemorrhagic effect was seen, but the maximum measured PRPA was only 1.6 MPa (MI = 0.65).

The liver is often the subject of ultrasound examinations, and these can be improved by contrast agents. Effects of Levovist-aided ultrasound on rat liver were investigated by Shigeta et al.¹¹⁴ The on-screen MI values were 1.8 at 8 MHz

and 0.7 at 12 MHz, and both were used on each rat. The transducers were moved to expose the entire liver. Electron microscopy was performed on the rat livers with control, sham, and ultrasound-only groups and two groups with ultrasound plus contrast, 1 killed immediately and the other killed 5 hours later. Qualitative observation of the specimens revealed platelet aggregation in the liver sinusoids for ultrasound-only and ultrasound-plus-contrast groups. Endothelial cell damage was seen in the ultrasound-plus-contrast group with the 5-hour delay.

Pulsed ultrasound has been shown to induce lung hemorrhage under some conditions, but the exact mechanism is not clear. O'Brien et al.¹¹¹ tested the hypothesis that cavitation was the mechanism by using saline or Optison injection with pulsed ultrasound exposure. Ten-second exposures were performed on rats at 3.1 MHz with 1.2-microsecond pulses and a 1-kHz pulse repetition frequency and in situ PRPAs of 2.74 and 5.86 MPa (equivalent to in situ MI values of 1.6 and 3.3, respectively). The contrast agent groups did not have an increase in lung lesion occurrence or size relative to the saline groups, which suggests that microbubble cavitation was not the mechanism for the lung hemorrhage. This finding confirms an earlier test using Albunex.¹¹⁵

Contrast-aided ultrasound scanning of various tissues can assist in the identification of malignant tumors but might cause microvascular perturbations. Melanoma tumors growing on the thighs of mice, which undergo metastatic spread to the lungs, were scanned with 1.5-MHz diagnostic ultrasound during or after Definity injection.¹¹² Image frames were triggered at a 1-Hz rate, and four 10- μ L/kg retro-orbital injections of the contrast agent were made over a 100-second exposure. Sham exposure involved scanning for 100 seconds followed by Definity injection with the ultrasound off. For ultrasound plus the contrast agent, observation of a brightening of the tumor image confirmed the interaction of ultrasound with the contrast agent within the tumor. One day after scanning, the primary tumor was surgically removed, and the possible lung metastasis was allowed to develop for 28 days. No significant increase in lung metastases was seen in the lungs for the contrast-aided ultrasound group.

5.3. Bioeffects Above the Diagnostic Limit

Several researchers have studied bioeffects of ultrasound contrast agents with pulsed ultrasound (for which cavitation might not normally occur) that had pressure amplitudes greater than the US FDA guideline upper limit for diagnostic ultrasound. Miller and Gies¹¹⁶ injected hairless mice with Alunex and exposed the abdominal region to lithotripter shock waves. An increase in mortality was found after exposure to several hundred shock waves, with increasing mortality for increasing Alunex doses. Hynynen et al¹¹⁷ found that 1.63-MHz focused ultrasound with 10-microsecond pulses at 6.3 MPa produced histologically observable tissue damage in rabbit brains. For this work, a window was created in the skull for exposure, and a bolus of Optison was injected 10 seconds before exposure. Hwang et al¹¹⁸ examined effects of pulsed 1-MHz ultrasound on rabbit ear veins with Optison in the circulation. Vessel wall damage with Evans blue extravasation was increased by the contrast agent at 6.5 MPa, and a small percentage of the endothelial surface was damaged at 3.35 MPa with the contrast agent. Hemolysis in suspensions has been studied for many years as an indicator of cavitation bioeffects. Dalecki et al¹¹⁹ detected hemolysis in vivo in mice with Alunex in the circulation. For a 10-microsecond pulsed exposure of the heart, thresholds were 3.0 MPa (peak positive or 1.9-MPa negative pressure amplitude) at 1.15 MHz but in excess of 10 MPa (peak positive) at 2.35 MHz.

The capillary hemorrhage effect was studied in mice injected with 0.1 mL of Alunex using positive or negative pulses from an endoscopic lithotripter (≈ 0.4 MHz).¹²⁰ One hundred pulses of 3.6 or -3.6 MPa in amplitude were delivered to the mouse abdomen. The negative pulses were significantly more effective than the positive pulses in producing hemorrhage in various tissues, including kidney, intestine, muscle, and stomach, which showed that the hemorrhage resulted from inertial cavitation.

Premature complexes (ECG signals representing ventricular electrical activity) were seen for 10 of 20 rats exposed to 3.1-MHz ultrasound with 1.3-microsecond pulses at 15.9 MPa and Optison in the circulation.¹²¹ However, myocardial degeneration was identified by histologic stain-

ing in 16 rats, which suggests that the presence of myocardial degeneration alone was not a sufficient explanation of the premature complexes.

5.4. Potential Therapeutic Applications

Contrast-aided ultrasound is capable of inducing a variety of in vivo bioeffects, and some of these effects may have useful clinical applications for therapy. There have been many reports of high-power contrast-aided ultrasound use for gene therapy, thrombolysis, and surgical applications, which will not be considered here. In this document, only reports of in vivo therapeutic applications that have involved actual diagnostic ultrasound systems for treatment are noted. These reports do not directly address the problem of bioeffects risks in diagnosis and often involved special gas body agents (they are not listed in Tables 1 and 2), but they have a bearing on the perceived significance of the possible effects.

The contrast ultrasound-induced effect of vascular permeabilization has been suggested as a means of drug delivery from the blood pool to the interstitium.¹²² The method can accommodate small particles as well as molecular drugs. Drug delivery was aided by Optison and was targeted to skeletal muscle by 2.3-MHz diagnostic ultrasound, which served both as an image guidance device and as the ultrasound exposure system.

Sonoporation, which is defined as transient ultrasound-induced enhancement of cell membrane permeability, has been used in applications of gene and drug delivery. Pislaru et al¹²³ used a phased array transducer of a commercial ultrasound imaging system (GE VingMed System V) in in vitro and in vivo experiments. For in vivo experiments, Sprague-Dawley male rats were exposed with a 1.7-MHz diagnostic ultrasound system and PESDA to transfect skeletal muscle with the luciferase marker plasmid. Tissue samples were also taken from remote, noninjected muscle or from the liver, kidney, lung, and heart. Luciferase activities were about 10-fold higher than with intramuscular injections of the plasmid alone. Cationic lipid-DNA complexes incorporated into microbubbles have also been tested for DNA transfer in skeletal muscle.¹²⁴ Diagnostic ultrasound at 1.75 MHz was used in

the B-mode with in situ PRPA values of 1.04 to 1.14 MPa. The luciferase marker gene was used, and gene expression was assessed after 4 days. Intramuscular injection of the plasmid alone produced strong gene expression, which was matched by the intra-arterial treatment with plasmid-loaded microbubbles and ultrasound. No luciferase expression was seen outside the ultrasound-treated region in liver or lung tissue or in muscle treated with ultrasound and the plasmid intra-arterially but without enhancement of cavitation by added microbubbles.

Gene delivery to the myocardium of rats was enhanced by treatment with harmonic mode diagnostic ultrasound, a microbubble contrast agent, and a viral β -galactosidase vector.¹²⁵ The contrast agent was prepared in the laboratory and was processed with the vector to attach the virus particles to the microbubbles. Three frames from a 1.3-MHz transducer destroyed the microbubbles evident in the second-harmonic image, and 3 frame bursts were triggered intermittently to allow refill of the tissue between scans. The hearts of all animals that received the combined ultrasound plus microbubble treatment showed expression of the transgene. Vannan et al¹²⁶ used cationic microbubble-linked plasmids and diagnostic ultrasound exposure to enhance transfer of the chloramphenicol acetyltransferase (CAT) marker gene in dog hearts. Diagnostic ultrasound was delivered into anesthetized closed-chest dogs at 1.3 MHz and the highest power settings (MI = 1.5–1.7). Multiple frames were triggered at every 4 to 6 cardiac cycles in an apical 4-chamber view. The specially prepared microbubbles were injected into a cephalic vein. For ultrasound with the plasmid-loaded microbubbles, CAT expression was found in several regions of the heart, with 303 ± 188 ng/g in the myocardium for 4 dogs. However, for the ultrasound treatment of the heart only, CAT expression was found in distant tissues of the lungs, liver, kidney, and skeletal muscle.

Ultrasound-enhanced gene transfer to cardiac tissue was also shown using albumin and lipid microbubbles containing a luciferase plasmid.¹²⁷ The agents were infused for 20 minutes through the jugular vein of anesthetized rats, and the hearts were scanned with a 1.3-MHz

cardiac ultrasound machine at an MI of 1.5 with 4 frames triggered from the ECG at every 4 cardiac cycles. Luciferase expression after 4 days was primarily detected in the heart, with some gene expression evident in the liver for the albumin microbubbles and in the pancreas for the lipid microbubbles. The echocardiographic treatment parameters were varied to find the optimum treatment conditions for the adenovirus- or plasmid-modified contrast agent microbubbles.¹²⁸ Cardiac scanning was performed in anesthetized rats to transfer the luciferase plasmid. Triggered imaging at 1.3 MHz was more effective than continuous imaging for gene transfer to the heart. An increase of the MI from the normal maximum of 1.6 for the diagnostic machine used in the study to 2.0 produced significantly greater gene transfer (the FDA guideline upper limit for diagnostic ultrasound is MI = 1.9).

Myocardial infarction might be treated by angiogenic gene therapies. Zhigang et al¹²⁹ used ultrasound to enhance DNA transfer of a gene vector coding for vascular endothelial growth factor in a rat model of myocardial infarction. An albumin-based contrast agent was mixed with a plasmid and incubated to attach the plasmid to the microbubbles. Three days after infarction, the plasmid vehicle was injected via the tail vein and targeted to the heart by 1.8-MHz echocardiography at the maximum MI with ECG triggering at every 6 to 8 beats. A statistically significant increase in the microvascular density in the ischemic myocardium was found in the ultrasound-plus-plasmid group. The rat model of myocardial infarction was also used by Kondo et al¹³⁰ to test the efficacy of gene therapy by hepatocyte growth factor. The naked plasmid coding for hepatocyte growth factor was injected through a catheter inserted into the left ventricle, while the femoral vein was used to infuse Optison microbubbles. The treatment involved 1.3-MHz ultrasound triggered in 3 frame bursts 1:8 at end systole at a PRPA of 2.16 MPa. The capillary density in the area around the infarct was 50% greater in the contrast-plus-plasmid group than in control groups, and staining for scar formation showed a significantly smaller scar area.

6. Conclusions

6.1. General Conclusions

Diagnostic ultrasound exposure can destabilize contrast agent gas bodies (microspheres, or stabilized microbubbles). In practice, the use of high MI values (>0.8) involves rapid gas body destruction, while use of low MI values (<0.2) involves minimal gas body destruction. Physical models of gas body behavior indicate that inertial cavitation potentially can occur with exposure conditions corresponding to MI values greater than approximately 0.4, which therefore represents a theoretical boundary between non-inertial and inertial activity regimens. The complex relationship between contrast agent destabilization and inertial cavitation remains the subject of active research.

In suspensions of nonphagocytic cells, the dominant mechanism by which extensive cell lysis is produced by ultrasound exposure with contrast agents appears to be the occurrence of inertial cavitation, and cavitation acoustic emissions can be used as cavitation “dosimeters.” In vitro studies have shown that diagnostic ultrasound exposures of very modest PRPAs can kill attached monolayer cells when in contact with contrast agent gas bodies. The cell injury can occur below the inertial cavitation threshold, apparently by a microstreaming shear stress mechanism. Because contrast agent gas bodies attach to phagocytic cells, these cells, which constitute the mononuclear phagocytic system in the body, may be particularly vulnerable. However, conditions in the body are different from specialized in vitro conditions, and translation of this basic research finding to in vivo conditions is not possible at this time.

Studies of bioeffects induced by contrast-aided diagnostic ultrasound in vivo, primarily in small animals, have shown biologically significant microscale effects, such as petechial hemorrhage with ultrasound exposures corresponding to MI values above 0.4. This value agrees with the theoretical threshold for inertial cavitation in blood, which contains potential cavitation nuclei. Additional information confirms that an MI of 0.4 is an important boundary, as stated in the American Institute of Ultrasound in Medicine statement *Bioeffects of Diagnostic*

*Ultrasound with Gas Body Contrast Agents.*¹³¹

Above an MI of 0.4, bioeffects appear to increase rapidly in magnitude, possibly as the square or high-order exponent of the PRPA. The magnitude of effects appears to be proportional to the contrast agent dose in terms of the number of gas bodies for low doses. Reported bioeffects of contrast-aided diagnostic ultrasound include sonoporation, microvascular leakage, capillary rupture (petechial hemorrhage), microlesions with inflammatory cell infiltration, and PVCs during ultrasound scanning. The bioeffects are primarily in the form of scattered microscopic injuries, which would not be expected to be clinically detectable in the short term. An exception is the induction of PVCs during contrast echocardiography, which is clinically observable but ceases on cessation of the ultrasound. The longer-term medical significance of the reported bioeffects is uncertain. Intentional bioeffects for therapeutic purposes can be produced or enhanced with diagnostic ultrasound exposure of contrast agents, such as in gene therapy. No epidemiologic studies are available on possible adverse (or favorable) human health effects of the clinical use of ultrasound contrast agents.

6.2. Specific Conclusions

1. Studies of bioeffects induced by contrast-aided diagnostic ultrasound in vivo, primarily in small animals, have shown bioeffects, such as PVCs and petechial hemorrhage with ultrasound exposures corresponding to MI values above 0.4. This value agrees with the theoretical threshold for inertial cavitation in blood, which contains cavitation nuclei.
2. Above an MI of 0.4 bioeffects appear to increase rapidly in magnitude as the square or higher exponent of the PRPA. The magnitude of effects appears to be proportional to the concentration of gas bodies for concentrations at or below the manufacturers' recommended dose.
3. In vitro studies have shown that diagnostic ultrasound exposures can induce death of attached monolayer cells when in contact with contrast agent gas bodies. Cell injury can occur below the inertial cavitation threshold; the minimum reported thresh-

- olds for cell death were approximately $p/f = 0.06$ MPa/MHz.
4. Contrast agent gas bodies can be bound and internalized by phagocytic cells, making them particularly vulnerable to injury from ultrasound exposure. The clinical implications, if any, of these results are unknown.
 5. Inertial cavitation is the dominant mechanism of cell lysis in whole blood exposed *in vitro* or *in vivo* to diagnostic ultrasound in the presence of contrast agents at suprathreshold pressure amplitudes. The clinical implications, if any, of these results are unknown.
 6. Interaction of diagnostic ultrasound with contrast agents is under investigation for therapeutic applications.

7. Recommendations

1. For imaging with contrast agents at MIs greater than 0.4, practitioners should use the minimal agent dose, MI, and exposure time consistent with acquisition of diagnostic information.
2. Practitioners of contrast-aided echocardiography should note that use of high MI values (>0.8) involves rapid gas body destruction with a potential for bioeffects (eg, PVCs), whereas bioeffects have not been observed at low values of $\frac{p}{\sqrt{f}}$ (<0.2), which involve minimal gas body destruction.
3. The ECG should be monitored during high-MI contrast cardiac-gated perfusion echocardiography, particularly in patients with a history of myocardial infarction or unstable cardiovascular disease.*

*The US FDA issued a Healthcare Provider Advisory in October 2007 stating that rare but serious cardiopulmonary complications, including fatalities, have occurred during or within 30 minutes after use of some approved ultrasound contrast agents (Definity and Optison). Changes to the package insert added at that time state that the ECG should be monitored in all patients, and monitoring should be continued for at least 30 minutes after administration of these agents. At the time of preparation of this document, the data associated with the FDA recommendations were not available to the Bioeffects Committee for review.

4. Output display indices represent important information and should be documented as part of the permanent record of the examination to enable future research.
5. The initial setting of the MI at mode selection for contrast-aided ultrasound imaging should be 0.4 or less.
6. Contrast-specific exposure indices should be developed that reflect the destruction thresholds and more accurately account for the *in situ* exposure values (derating factor).
7. Studies in laboratory animals are needed to investigate the potential for ultrasound-induced adverse fetal effects in the presence of ultrasound contrast agents or other exogenously introduced bubbles.

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