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▶ To cite this version:

Michiel Postema, Georg Schmitz. Bubble dynamics involved in ultrasonic imaging. Expert Review of Molecular Diagnostics, 2006, 6 (3), pp.493-502. 10.1586/14737159.6.3.493. hal-03193345

HAL Id: hal-03193345

https://hal.science/hal-03193345

Submitted on 11 Apr 2021

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Bubble dynamics involved in ultrasonic imaging

Running title: Bubble dynamics

Michiel Postema, Georg Schmitz

Institute for Medical Engineering, Ruhr-Universität Bochum, Bochum, Germany

Affiliations

Dr. Michiel Postema Prof. Dr.-Ing. Georg Schmitz

Institute for Medical Engineering | Institute for Medical Engineering

Department of Electrical Engineering | Department of Electrical Engineering

and Information Technology and Information Technology

Ruhr-Universität Bochum Ruhr-Universität Bochum

Building IC, 6/146 Building IC, 6/135

D-44780 Bochum D-44780 Bochum

Germany Germany

Tel: +49 234 3227740

Fax: +49 234 3214872

michiel.postema@rub.de georg.schmitz@rub.de

Summary — In clinical ultrasound, blood cells cannot be differentiated from surrounding tissue, due to the low acoustic impedance difference between blood cells and their surroundings. Resonant gas bubbles introduced in the blood stream are ideal markers, if rapid dissolution can be prevented. Ultrasound contrast agents consist of microscopically small bubbles encapsulated by an elastic shell. These microbubbles oscillate upon ultrasound insonification. Microbubbles with thin lipid shells have demonstrated highly nonlinear behavior. To enhance diagnostic ultrasound imaging techniques and to explore therapeutic applications, these medical microbubbles have been modeled. Several detection techniques have been proposed to improve the detectability of the microbubbles. A new generation of contrast agents, with special targeting ligands attached to the shells, may assist in imaging nonphysical properties of target tissue. Owing to microbubble-based contrast agents, ultrasound is becoming an even more important technique in clinical diagnostics.

Keywords: microbubbles, ultrasound, ultrasound contrast agent, scattering, resonance, RPNNP equation, Herring equation, harmonic imaging, targeted imaging, detection.

Bubble dynamics involved in ultrasonic imaging

Ultrasonic imaging is a relatively cheap, reliable diagnostic technique. Typical diagnostic ultrasonic frequencies range from 1 MHz (heart, liver) to 100 MHz (eye, skin). A signal generated by an ultrasonic transducer typically consists of a pulse of a few μ s with an angular center frequency ω . Part of this signal propagates through target tissue, part is reflected by macroscopic tissue structures, part is absorbed by tissue, and part is scattered by structures in the tissue smaller than the acoustic wavelength. A small portion of the transmitted acoustic energy is received by the transducer, which is used to build an ultrasonic image. The received signal is the superposition of specular reflections at tissue boundaries and echoes from tissue backscattering. The quantity of signal specularly reflected from a boundary separating tissues 1 and 2 depends on the acoustic impedance change in the tissue:

$$R_{\rm I} = \frac{(Z_2 - Z_1)^2}{(Z_2 + Z_1)^2} \,, (1)$$

where $R_{\rm I}$ is the intensity reflection coefficient. The acoustic impedance of a medium is defined by:

$$Z = \rho \, c = \sqrt{\frac{\rho}{\kappa}} \,, \tag{2}$$

where c is the speed of sound in the medium, κ is the compressibility, and ρ is the density.

Scattering

Scattering is far more complicated [1]:^a An inhomogeneity in the path of a sound wave causes it to spread out in a variety of directions [2]. The quotient representing the power scattered per unit solid angle and per unit incident intensity, is referred to as the differential scattering cross section [2]. When the direction toward the receiver extends back to the source, such as in pulse-echo systems, we speak of backscattering. Under this condition, the backscattering cross section equals 4π times the differential scattering cross section [2]. When multiplying the differential scattering cross section and the backscattering cross section with the density of the scatterers per unit volume, one obtains the scattering coefficient and the backscattering coefficient, respectively. The backscattering coefficient of a medium containing a small concentration of fluid spheres with radii r is given by [3]:

$$\eta(\omega) = 3 k^4 \gamma_0^2 r^3 \left(\frac{J_1(2kr)}{2kr} \right)^2,$$
(3)

where k is the acoustic wave number, J_1 is the first order Bessel function of the first kind, and γ_0 the variance of the density/compressibility fluctuation:

$$\gamma_0^2 = N_0 \frac{4\pi r^3}{3} \left(\frac{\kappa_1 - \kappa_0}{\kappa_0} - \frac{\rho_1 - \rho_0}{\rho_0} \right)^2, \tag{4}$$

where N_0 is the number of scatterers per unit volume, κ_1 is the compressibility of the scatterer, κ_0 is the compressibility of the surrounding medium, ρ_1 is the density of the scatterer, and ρ_0 is the density of the surrounding medium.

^aThe References in this paper are far but complete. They should, however, provide the reader with sufficient background reading material for further study.

From equations (1) and (4) it follows that blood cells are poor scatterers in the clinical diagnostic frequency range. Since imaging blood flow and measuring organ perfusion are desirable for diagnostic purposes, a marker has to be added to the blood that helps to differentiate between blood and other tissue types, by providing additional and desirably characteristic backscatter [4]. Gas bubbles are suitable markers, not only because they have a high compressibility (gaseous microbubbles are 10,000 times more compressible than red blood cells [5]) and low density compared to the surrounding medium, but also because they resonate. For illustration, De Jong computed the scattering cross section in water of a rigid iron sphere and a gas sphere, both with a $1 \mu m$ radius [6]. At a driving frequency of 3 MHz, the scattering cross section of the gas sphere is

Ultrasound contrast agents

Microbubbles that are used for ultrasonic imaging purposes are referred to as *ultrasound contrast agents*. Their development has gone through several generations [7]. Table 1 gives an overview of the ultrasound contrast agents used most in imaging research. Free microbubbles represent generation 0. These bubbles rapidly dissolve owing to diffusion.

The change of gas bubble radius as a function of time, due to diffusion, is given by [8]:

$$\frac{\mathrm{d}r}{\mathrm{d}t} = DL \left(\frac{\frac{C_{\rm i}}{C_0} - 1 - \frac{2\,\sigma}{r\,p_0} - \frac{p_{\rm s}^+}{p_0}}{1 + \frac{4\,\sigma}{3\,r\,p_0}} \right) \left(\frac{1}{r} + \frac{1}{\sqrt{\pi\,D\,t}} \right) ,\tag{5}$$

where $\frac{C_1}{C_0}$ is the ratio of the dissolved gas concentration to the saturation concentration (saturation ratio), D is the diffusion constant, L is the Ostwald coefficient, p_0 is the ambient pressure, p_s^+ is the applied static overpressure, r is the instantaneous bubble radius, t is the time starting (t = 0) when the bubble surface is exposed to the liquid surface, and σ is the surface tension. Equation (5) shows that the disappearance of gas bubbles in a liquid medium is highly influenced by gas diffusion parameters and applied overpressure, and that the disappearance time of gas bubbles is shorter when the liquid medium is under pressure. The difference, however, is of a much smaller order than the half-size time of the bubbles (the time it takes for a bubble to dissolve until it has reached half its initial size [9]), as Figure 1 demonstrates.

To prevent them from rapid dissolution, generation 0 microbubbles could have diameters of 80 μ m, which withheld them from passing the lung capillaries. First generation ultrasound contrast agents consist of air bubbles encapsulated by a stabilizing shell. With mean diameters below 6 μ m, these bubbles are small enough to pass through capillaries.

For encapsulated microbubbles, shell stiffness parameters χ , $S_{\rm sh}$ have been introduced [6, 10]:

$$\chi = \frac{S_{\rm sh}}{8\pi} = \frac{E\,\epsilon}{1-\nu}\,,\tag{6}$$

where E is Young's modulus, ϵ is the shell thickness, and ν is the Poisson ratio. For albumin and lipid nanoshells, we presume $0.499 < \nu < 0.500$ [11]. The shell properties of an ultrasound contrast agent have to be derived from experimental data. Second generation agents consist of encapsulated gas microbubbles with an elastic shell. The linear angular resonance of microbubbles with a viscous shell can be approximated by a Newtonian model [12, 13]:

$$\omega_0^2 \approx \frac{3\Gamma}{r_0^2 \rho} \left(p_0 + \frac{2\sigma}{r_0} + \frac{2\chi}{r_0} \right) - \frac{2\sigma + 6\chi}{r_0^3 \rho} ,$$
 (7)

or by a viscoelastic model [14, 15]:

$$\omega_0^2 \approx \frac{1}{r_0^2 \rho} \left(3 \Gamma p_0 - \frac{4\sigma}{r_0} + \frac{4E}{r_0} \right) .$$
 (8)

Here, r_0 is the equilibrium bubble radius, Γ is the polytropic exponent of the gas, ρ is the density of the host medium, and ω_0 is the angular resonance frequency. Clearly, if the size distribution of the microbubbles is wide, the agent will respond to a wide ultrasonic frequency band. If the size distribution of the microbubbles is narrow, the agent will be selective to a narrow frequency band.

When using perfluorocarbon gases instead of air, the microbubbles will first swell by a factor, because of the diffusion of dissolved gases into the bubbles, and then dissolve. The low diffusion rate of high molecular weight perfluorocarbons prolongs microbubble presence from seconds to minutes [4]. Often, the surface of the bubble shell has a negative charge, to prolong its presence in target tissue [16].

Agents can be designed to specifically target a receptor system [17], thus facilitating ultrasonic molecular imaging [18]. Third generation ultrasound contrast agents consist of bubbles with such special targeting shell properties. Owing to primary radiation forces, microbubbles can be forced to translate away from the transducer, to vessel walls [18, 19], increasing the success rate of targeting.

Submicron contrast agents have also been of interest, because they can travel through the lymphatic system and are small enough to be extravasated from tumor neovasculature [5, 20].

Ultrasound contrast agents have been involved in therapeutic applications, as well. Because any drug might in some form be attached to a bubble, the number of potential therapeutic applications of bubbles is virtually unlimited. To limit our scope to diagnostic applications, here we describe the physical mechanisms involved in bubble-imaging techniques.

Table 1: Overview of the ultrasound contrast agents used most in diagnostic imaging research [21, 22, 23, 24, 25, 26, 27, 28, 29, 7, 30]. First generation: encapsulated air bubbles, second generation: encapsulated low solubility gas bubbles, third generation: particulated gas bubbles with controlled acoustic properties [31]. This classification is approximate, because not all agents fit neatly into the generation categories.

Agent	Manufacturor	Shell	Gas/vapor	Mean diameter (μm)
First generation				
Albunex	MolecularBiosystems	Albumin	Air	4.3
Levovist	Schering	Lipid/galactose	Air	2–3
Sonovist	Schering	Cyanoacrylate	Air	1–2
Second generation				
BR14	Bracco	Lipid	C_4F_{10}	2.5 – 3.0
Definity	Bristol-Myers Squibb	Lipid/surfactant	C_3F_8	1.1–3.3
EchoGen	Sonus	Surfactant	C_5F_{12}	2–5
Imagent	Alliance Pharmaceutic	Lipid/surfactant	N_2/C_6F_{14}	6.0
Optison	GE Healthcare	Albumin	C_3F_8	2.0 – 4.5
Quantison	Upperton	Albumin	Air	3.2
SonoVue	Bracco	Lipid	SF_6	2.5
Third generation				
AI-700	Acusphere	PLGA^a	C_4F_{10}	2
CARDIOsphere	POINT Biomedical	Polyactide	N_2	4.0
Sonazoid	GE Healthcare	Lipid/surfactant	C_4F_{10}	2.4 – 3.6

^aPolyactide co-glycolide

Originally, ultrasound contrast studies were performed for left ventricular function and myocardial perfusion [31]. Nowadays, ultrasound contrast agents have, among others, been implicated in the following diagnostic techniques: imaging the heart [27], vasculature including vasa vasorum, liver, spleen [7], kidneys [32], brain [33, 34], measuring tissue perfusion [4], ejection fractions [35], detecting focal lesions in the liver [36], angiogenesis assessment [37], characterizing tumors, and detecting sites of inflammation [26].

The ultrasound contrast agents currently approved for certain clinical applications are Levovist (Canada, China, Europe, Japan), Definity (Canada, United Kingdom, United States), Optison (Canada, Europe, United States) and SonoVue (China, Europe) [38, 27].

Contrast bubble models

To enhance detection techniques, predicting the dynamic behavior of ultrasound-insonified encapsulated microbubbles has been of much interest. Their behavior has been studied with acoustic methods such as the measurement of backscattering under different conditions [39], (high-speed) photography [40, 41, 42, 9], and other optical methods, such as the dynamic measurement of light scattering [13].

The oscillating behavior of ultrasound contrast agent microbubbles in Newtonian fluids has been frequently described by a Rayleigh-Plesset-Noltingk-Neppiras-Poritsky (RPNNP) equation modified for the stiffness of a shell [6, 43, 44] and by a Herring equation modified for a viscous shell [41, 13, 18]. Other models include

an RPNNP equation modified for the mechanical properties of the shell [45], a zero-thickness interface model [14], and a model accounting for variations in surface tension [46]. Generally, the presence of blood has a relatively small effect on bubble dynamics [47]. The RPNNP equation has been modified to include the shell properties stiffness and friction [48]:

$$\rho \, r \, \ddot{r} + \frac{3}{2} \rho \dot{r}^2 = \left(p_0 + \frac{2\sigma}{r_0} \right) \left(\frac{r_0}{r} \right)^{3\Gamma} - \frac{4\mu \, \dot{r}}{r} - \frac{2\sigma}{r}$$

$$-2\chi \left(\frac{1}{r_0} - \frac{1}{r} \right) - \omega^2 \, \rho \, r^2 \frac{\dot{r}}{c} - \frac{S_{\rm f} \, \dot{r}}{4\pi \, r^2} - \delta_{\rm t} \, \omega \, \rho \, r \, \dot{r} - (p_0 + p_{\rm a}(t)) \,,$$
(9)

where c is the speed of sound in the medium, $p_{\rm a}(t)$ is the driving acoustic pressure in time, $S_{\rm f}$ is the shell friction, μ is the shear viscosity of the medium, ω is the angular driving frequency, and $\delta_{\rm t}$ is the damping coefficient due to heat conduction [49]:

$$\delta_{t} = \frac{\frac{\sinh z + \sin z}{\cosh z - \cos z} - \frac{2}{z}}{\frac{z}{3(\Gamma - 1)} + \frac{\sinh z - \sin z}{\cosh z - \cos z}},$$
(10)

where $z = r_0/l_D$, in which l_D is the thermal boundary layer thickness:

$$l_{\rm D} = \sqrt{\frac{K_{\rm g}}{2\,\omega\,\rho_{\rm g}\,C_{\rm p}}}\,. (11)$$

Here, C_p is specific heat of the gas, K_g is the thermal conductivity of the gas, and ρ_g is the density of the gas. Note that the vapor pressure of the liquid has been neglected, which is valid in our situation [50]. The modified RPNNP equation is used for simulating bubble response to insonification at moderate acoustic pressures, i.e. $\text{MI}^b \lesssim 0.6$ [9]. An example of simulated oscillating behavior of a $\frac{1}{p}$ On clinical ultrasound devices, the intensity of the ultrasonic field is generally adjusted with a switch for the mechanical index instead of the acoustic amplitude. The mechanical index is defined as $\text{MI} = p^-/\sqrt{f}$, where p^- is the peak rarefactional acoustic pressure normalized by 1 MPa and f is the center frequency of the ultrasound normalized by 1 MHz.

microbubble is shown in Figure 2. The microbubble with half resonance size oscillates exactly $\phi = \pi$ rad out of phase with the driving pressure $p_a = p^+ \sin \omega t$ in which p^+ is the peak-positive acoustic pressure. At resonance, $\phi = \frac{3}{2}\pi$ rad, and at double resonance $\phi = 2\pi$ rad. As the damping becomes less, the transition in offset becomes more abrupt [49]. For microbubbles with thin elastic shells, the modified RPNNP equation gives a slightly conservative estimate of the radial excursion, as opposed to the modified Herring equation [9]. The modified Herring equation is given by [41, 13]:

$$\rho r \ddot{r} + \frac{3}{2} \rho \dot{r}^{2} = \left(p_{0} + \frac{2\sigma}{r_{0}} + \frac{2\chi}{r_{0}} \right) \left(\frac{r_{0}}{r} \right)^{3\Gamma} \left(1 - 3\frac{\dot{r}}{c} \right) - \frac{4\mu \dot{r}}{r} - \frac{2\sigma}{r} \left(1 - \frac{\dot{r}}{c} \right) - \frac{2\chi}{r} \left(\frac{r_{0}}{r} \right)^{2} \left(1 - 3\frac{\dot{r}}{c} \right) - 12\epsilon \,\mu_{s} \frac{\dot{r}}{r(r - \epsilon)} - (p_{0} + p_{a}(t)),$$
(12)

where $\mu_{\rm s}$ is the shell shear viscosity.

Equations (9) and (12) can only be solved numerically. Figure 3 shows solutions of the modified Herring equation and Fourier-spectra thereof, at different acoustic amplitudes, for a bubble with a negligible shell. At 10 kPa the bubble oscillates linearly, but at higher acoustic amplitudes the bubble has a longer expansion than a contraction phase, and a higher outward than inward excursion. At 391 kPa, a slow expansion is followed by a rapid collapse, which is followed by a number of rebounds. Furthermore, the maximal bubble excursion is more than twelve times the equilibrium radius. The nonlinear behavior described here generates harmonics. Examples of the acoustic emission from ultrasound contrast agent bubbles can be appreciated in [20]. Figure 4 demonstrates the spectra of the ultrasound contrast agent Levovist, insonified at different driving

amplitudes [34]. Similar results were presented in [51]. The quasi-continuous part in the spectrum of the acoustic response to the highest driving amplitude has been attributed to microbubble disruption, which is discussed in a following section.

Figure 5 shows r-t curves computed with both models, for a free gas microbubble, and for an encapsulated microbubble, insonified at a relatively high acoustic pressure of 210 kPa. Both for the free gas bubble and for the encapsulated bubble, the curves are very similar.

Harmonic imaging

At lower driving pressures, microbubbles produce linear backscatter enhancement, only resulting in an augmentation of the echo from blood [31]. By making use of nonlinear behavior of microbubbles resulting in harmonic backscatter, blood can be discriminated from the surrounding tissue. The increase of nonlinear behavior with acoustic driving pressure has been demonstrated in Figures 3–5. To suppress signal from tissue, a band pass filter around the second harmonic frequency component of the echo can be used to produce the ultrasound image [52]. This imaging technique is ineffective, due to the wide overlap of the baseband and the second harmonic in broadband systems. Furthermore, this technique is not effective at high driving pressures, since tissue harmonics may interfere with the signals from the perfused areas. Therefore, nondestructive imaging is preferably done at moderate acoustic pressures.

Given the lack of subharmonic generation in tissue, subharmonic imaging is

an alternative [53]. However, subharmonic generation occurs only, when the acoustic amplitude exceeds a certain threshold level. Goertz *et al.* recently demonstrated that microvessels can be successfully detected *in vivo* using subharmonic imaging with a transmit frequency of 20 MHz [54].

Multipulse techniques have been developed to allow for the separation of contrast agent signal from tissue signal [52]. In these techniques, multiple pulses with modifications in amplitude (power modulation) [55, 56], phase (pulse inversion) [57, 58, 56], or pulse length (pulse subtraction) [59] are transmitted rapidly after one another, after which the linear contribution in the signal echoes is canceled out by summation or subtraction, leaving a residual signal containing harmonics (cf. Figure 6). Overviews of multipulse imaging methods have been given in [58, 60, 59]. With the introduction of affordable broadband transducers, far more complicated pulse schemes have become possible, such as multifrequency excitation [59, 61]. Since the bubble oscillation amplitude depends on the ambient pressure, a relatively low frequency can be used as if to modulate the ambient pressure, whereas a frequency close to bubble resonance is used to excite the bubble. Two-frequency excitation may be sufficient to induce nonlinear behavior of the microbubbles at moderate incident pressures [59]. Coded excitation, another technique, operates by transmitting a relatively long pulse (high energy) with a low amplitude (low destruction rate), that is decoded (compressed) to obtain sufficient axial resolution. The use of chirps, long bursts with increasing or decreasing instantaneous frequency, has been proposed for coded excitation [59].

Microbubble disruption

During the collapse phase, when the kinetic energy of the bubble surpasses its surface energy, a bubble may fragment into a number of smaller bubbles [42, 62, 63]. Fragmentation has been exclusively observed with contrast agents with thin elastic shells. For these bubbles, fragmentation is the dominant disruption mechanism [62].

Thick-shelled bubbles have demonstrated gas release during a high-amplitude ultrasonic cycle [64, 9, 65]. The increased pressure difference between inside and outside of the bubble during the ultrasonic wave causes the shell to be stretched across a critical deformation, resulting into its mechanical cracking. The released bubble has an oscillation amplitude much higher than an encapsulated bubble of the same size. Therefore, the acoustic signal from a contrast agent after gas release differs from that of the same contrast agent before gas release, until the released gas has dissolved.

Although the coalescence (merge) of ultrasound contrast agent microbubbles has been observed *in vitro*, this phenomenon is negligible *in vivo*, because long low-amplitude bursts are needed for forced approach of microbubbles [9].

Ultrasound-induced disruption of the microbubbles will result in strong, transient harmonic echoes [66]. These strong echoes reveal instantly which areas on the resulting image represent perfused regions. After a disruptive ultrasonic burst, the disappearance of microbubble fragments or released gas may be traced with low-amplitude ultrasound, as well as the wash-in rate of fresh contrast agent [67]. With the pressure-dependant changes in the echo signal, hydrostatic

overpressures might be determined. Making use of the subharmonic response leads to more precise sizing [9, 68] and to more precise pressure measurements [44, 69].

Expert opinion

According to the World Health Organization, in 20–30% of cases worldwide, clinical considerations alone are not sufficient to come to a correct diagnosis [70]. Of these cases, 80–90% can be diagnosed using common X-ray or ultrasound examinations [70]. The most performed diagnostic imaging technology is X-ray, followed at a distance by ultrasound. For example, in Ontario, Canada, 63% of the diagnostic imaging examinations in hospitals are done with X-ray, 16% with ultrasound, 9% with computed tomography, 7% with nuclear medicine, 4% with magnetic resonance imaging, and 1% with catheterization [71]. When taking into account the absolute hospital operating expenses [71], X-ray and ultrasound have approximately the same price per examination. Other imaging techniques are roughly three times as expensive, except for catheterization, which is twenty times as expensive. However, X-ray is a less desirable imaging technique than ultrasound, due to the negative radiation effects. Therefore, novel ultrasound-based imaging techniques are being developed that may compete with other imaging techniques.

The main disadvantage of ultrasonography has been that only physical properties of the tissue are imaged. A new generation of ultrasound contrast agents may help to overcome this limitation, with ligands having specific targeting properties attached to the shell of the microbubbles. Such agents would certainly be more expensive than the current second generation agents. But, by providing accurate and reliable early diagnosis, contrast sonography could reduce downstream resource use and with that overall health care costs, justifying incremental examination costs [4].

Five-year view

Ultrasonic contrast-enhanced imaging applications depend on the detectability of microbubbles. The detectability in turn depends on the ultrasonic pulsing scheme. Initiatives have been undertaken to find the optimal pulse sequence for a maximal contrast-to-tissue-ratio in combination with an imaging resolution as high as possible. Enhancement in image quality owing to coded excitation has been anticipated in the near future.

Note that ultrasound contrast agent bubble responses will become better predictable, if the bubble populations have a narrower size distribution and uniform shell thicknesses.

Ultrasound images acquired using lipid-shelled microbubbles targeted to leukocytes were presented in [72]. The high resolution of these *in vivo* images in comparison to the *ex vivo* gamma camera images of the same tissue samples is striking. The potential of targeted ultrasonic imaging is clearly demonstrated.

Ultrasound-induced release of therapeutic substances from microbubbles has been proposed in numerous papers [73, 74, 75]. Combining diagnostic, targeted microbubbles with therapeutic substances may lead to a simple method to instantly cure upon diagnosis.

Contrast agents for magnetic resonance (MR) imaging have been designed to accumulate in specific cells [1, 76]. As opposed to ultrasound contrast agents, whose microbubbles are limited to flowing through vessels, MR contrast agents consist of nanoparticles small enough to access cells. Undesirable accumulation of MR contrast agent takes place in organs other than the imaged, too. Nanometer-sized magnetic resonance markers may be coupled to microbubbles. In the organ to be imaged, these markers are to be released with a high-MI ultrasonic burst. This will result in an increase of the uptake MR contrast agent by the targeted organ. Radionuclides used for positron emission tomography might also be encapsulated by microbubbles, to reduce unspecific binding.

Increasing concern has been shown for ultrasound safety issues when using ultrasound contrast agents [77]. Although no side effects have been officially reported [4], current acoustic amplitudes allowed are based on tissue without cavitation nuclei, such as microbubbles, present. Since the use of ultrasound contrast agents has become increasingly popular, and their applications have been multiplied, a new safety standard in ultrasonic imaging will have to be defined in the near future.

In conclusion, the authors think that ultrasound is becoming an even more important technique in clinical diagnostics.

Key issues

• Predicting the transient behavior of microbubbles.

• Improving detection presence of bubbles by:

improving pulsing schemes,

improving processing methods,

improving bubble properties.

- Combining ultrasonic imaging with other techniques, such as magnetic resonance imaging and positron emission tomography.
- Defining a new safety standard in ultrasonic imaging.

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 $[^]c$ •: of interest, ••: of considerable interest.

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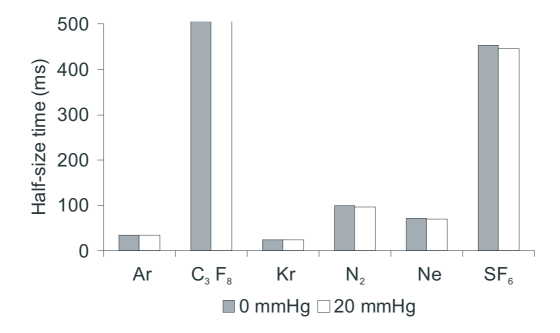


Figure 1: Half-size times of $r_0 = 3 \,\mu\text{m}$ bubbles containing different gases, at ambient pressure and at an overpressure of 20 mmHg. The saturation ratio was taken $\frac{C_i}{C_0} = 0$ [9]. The half-size time of C₃F₈ exceeds 1000 ms. The difference in dissolving time between 0 and 20 mmHg is on the order of 1 ms.

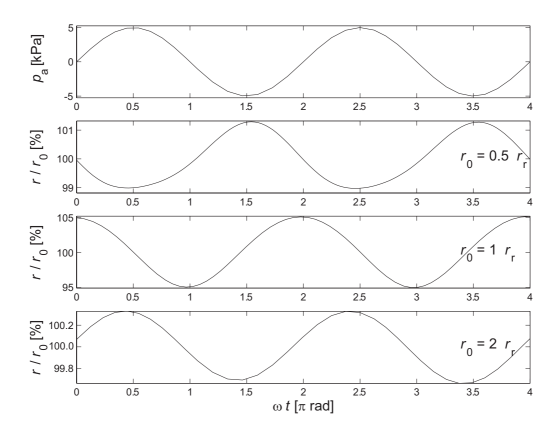


Figure 2: Simulations of the relative bubble surface excursion as a function of the driving phase for underdamped microbubbles with equilibrium radii $r_0 = \frac{1}{2}r_{\rm r}, 1r_{\rm r}, 2r_{\rm r}$, where $r_{\rm r}$ is the resonant radius, during insonification at $\omega/2\pi=0.5$ MHz and $p^+=5$ kPa. © 2005 Schiele & Schön GmbH. Reprinted with permission from Postema M, de Jong N, Schmitz G. The physics of nanoshelled microbubbles. *Biomed. Tech.* 50(S1), 748–749 (2005).

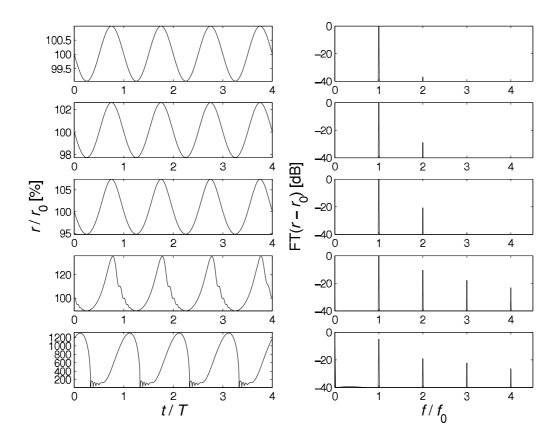


Figure 3: Computed solutions of the modified Herring equation and Fourier-spectra thereof for a bubble with a negligible shell (i.e. $\mu_s = \mu$ and $\chi = 0\,\mathrm{kg\,s^{-2}}$), at five different acoustic amplitudes, taking $c = 1480\,\mathrm{m\,s^{-1}}$, $\omega/2\pi = 0.5\,\mathrm{MHz}$, $p_0 = 1\,\mathrm{atm}$, $r_0 = 0.75\,\mu\mathrm{m}$, $\mu = 10^{-3}\,\mathrm{Pa\,s}$, $\rho = 998\,\mathrm{kg\,m^{-3}}$, and $\sigma = 0.072\,\mathrm{N\,m^{-1}}$. The acoustic amplitudes were 10.0, 25.0, 62.5, 156, and 391 kPa, respectively. Clearly, the harmonic components increase with the acoustic amplitude.

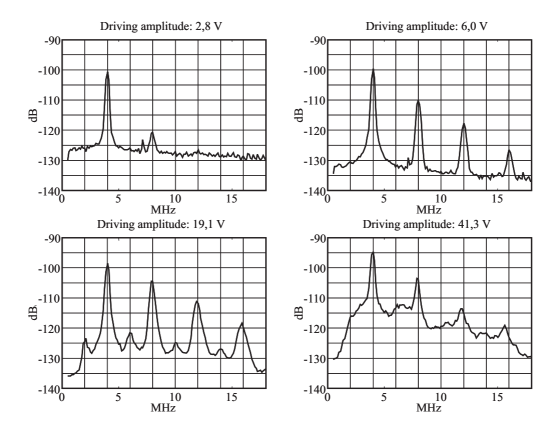


Figure 4: Measurements on a Levovist sample. The driving pulse was a narrow-band Gaussian pulse with a center frequency of 4 MHz and a -20 dB bandwidth of 512 Hz. The spectra of the acoustic responses were normalized by the respective driving amplitudes. The echo-amplitude rises out of proportion with the driving amplitude. Moreover, higher harmonics, subharmonics, and ultraharmonics appear. At the highest driving amplitude, the spectrum appears quasi-continuous, with a shift of the higher harmonics to lower frequencies. © 2003 Wilko G. Wilkening. Reprinted with permission from Wilkening WG. Konzepte zur Signalverarbeitung für die kontrastmittelspezifische Ultraschallabbildung. Ph.D. thesis, Ruhr-Universität Bochum (2003).

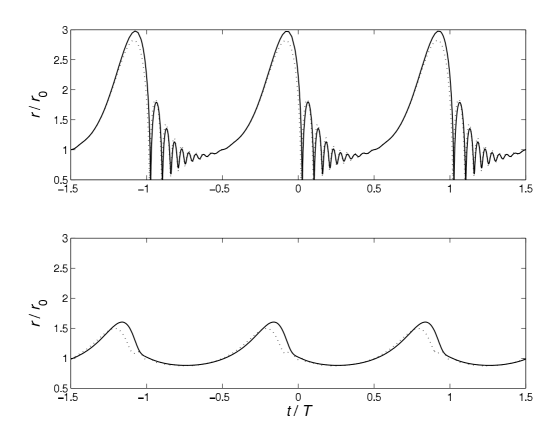
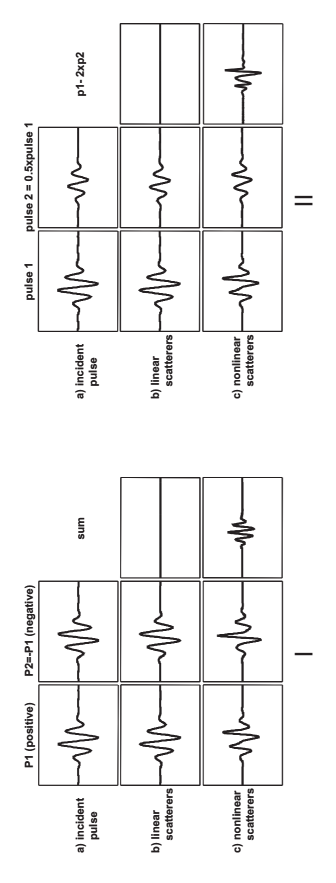


Figure 5: Computed solutions of the modified RPNNP equation (dotted line) and the modified Herring equation (bold line), taking $c=1480\,\mathrm{m\,s^{-1}},\ \omega/2\pi=0.5\,\mathrm{MHz},\ p_0=1\,\mathrm{atm},\ p^+=210\,\mathrm{kPa},\ r_0=0.75\,\mu\mathrm{m},\ \epsilon=20\,\mathrm{nm},\ \mu=10^{-3}\,\mathrm{Pa\,s},$ $\rho=998\,\mathrm{kg\,m^{-3}},\ \mathrm{and}\ \sigma=0.072\,\mathrm{N\,m^{-1}}.$ The upper frame shows r-t curves of a free gas bubble, with $S_\mathrm{f}=0\,\mathrm{kg\,s^{-1}},\ \delta_\mathrm{t}=0,\ \mu_\mathrm{s}=\mu\,\mathrm{Pa\,s},\ \mathrm{and}\ \chi=0\,\mathrm{kg\,s^{-2}}.$ The lower frame shows r-t curves of an encapsulated bubble with a thin, elastic shell, with $S_\mathrm{f}=0\,\mathrm{kg\,s^{-1}},\ \delta_\mathrm{t}=0,\ \mu_\mathrm{s}=10^{-1}\,\mathrm{Pa\,s},\ \mathrm{and}\ \chi=\frac{1.1}{8\pi}\,\mathrm{kg\,s^{-2}}.$ The shell stiffness χ has been determined in [10].



ultrasonography. In: Albrecht T, D'Onofrio M, Frauscher F, Frinking PJA, Lee SH, Nicolau C, Sidhu PS (Eds.), Contrast-Enhanced Figure 6: Multipulse contrast specific imaging by phase modulation (I) and by amplitude modulation (II). In Two incident pulses are used with the same amplitude but opposite phases. Ib After summation, signal echoes from linear scatterers cancel out. Ic Signal echoes from nonlinear scatterers do not fully cancel out after summation, leaving a residual signal containing (even) harmonics. Ila Two incident pulses are used with the same phase but amplitudes differing by a factor of two. IIb After subtraction, signal echoes from linear scatterers cancel out. Ilc Signal echoes from nonlinear scatterers do not fully cancel out after subtraction, leaving a residual signal containing harmonics. © 2005 Springer. Reprinted from Frinking PJA. Technological review of contrast-enhanced General Purpose Utrasound, Springer-Verlag, Milan, 1–8 (2005). With kind permission of Springer Science and Business Media.