Original Article

Stabilization of Gas Bubbles Released from Water-Soluble Carbohydrates Using Amphiphilic Compounds: Preparation of Formulations and Acoustic Monitoring of Bubble Lifetime

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Abstract The ultrasound contrast agents Echovist® and Levovist® (Bayer AG, Schering AG, Germany) are based on the release of gas bubbles from milled α-D-galactose. In diagnostic ultrasound, for this class of contrast agents, there is a need for prolonged contrast duration. To investigate if new carbohydrate compositions could prolong the lifetime of the gas bubbles, α-D-galactose was mixed with other carbohydrates or amphiphiles with varying log P. Acoustic attenuation vs. time (390 s) area under the curve (A_{390}) and bubble half-time (t/2) were used as measures of prolonged lifetime of gas bubbles. The products, to which 0.1% of a lipophilic carboxylic acid (5β-cholanic acid, behenic acid, and melissic acid) has been added, showed more than 5, 7 and 11 times enhancement of A_{390}, respectively, compared with the reference compound 2 (RC2) corresponding to the commercial product Levovist®. The half-time t ½ of the same compounds was prolonged more than 6 times compared with RC2. A partial least square (PLS) statistical analysis confirmed that, for additives, high log P carboxylic acids lead to the highest A_{390}. The present results bear a promise of products with a more persistent in vivo ultrasound contrast effect than the commercially available agents.

Keywords Gas bubbles · Ultrasound · Contrast agents · Carbohydrates · Gas-release · Amphiphiles · Colloid · Fatty acids · Acoustics

Introduction

Gas bubbles in the micrometer range are ideal contrast agents for ultrasound imaging because of their high compressibility and low density, giving excellent reflectance of ultrasound waves (acoustic backscatter) [1]. Ultrasound contrast signals from free gas bubbles in vivo in the human heart were first observed in 1968 after direct intracardiac injection [2]. Short bubble lifetime urged development of more stable agents giving more persistent contrast on the arterial side after intravenous injection and passage through the lung capillaries. The ultimate goal is to generate contrast in the heart muscle (myocardium). Suspensions of solid particles, emulsified liquid droplets, free and encapsulated gases or liquids [3] have been proposed as ultrasound contrast agents (USCA). Efficient USCA may comprise micron-sized bubbles stabilized by a thin, potentially biodegradable flexible membrane consisting of polymers [4] or phospholipids [5]. However, carbohydrate-based gas-releasing systems, generating free gas bubbles after intravenous injection of a particle suspension, eliminate the challenge of developing a stabilized bubble suspension as a...
drug product, with demands of long term shelf stability. In
the commercially available agent SHU 454 (Echovist®,
Schering AG, Berlin), milled, particulate water-soluble α-D-galactose acts as a precursor for gas bubbles, and is in
clinical use in enhanced hystero-sonography [6]. The gas
releasing powder is prepared using ball-milling of com-
mercial α-D-galactose, giving clusters of particles with a
mean size of about 40 μm, with air-filled voids between
them [7]. Upon dissolution in water, the clusters release the
gas trapped in the clusters and gas bubbles are generated.
The α-D-galactose crystals dissolve in the water phase while
acting as short-living nucleation sites for the gas bubbles
[7]. Lack of myocardial contrast enhancement (MCE) with
SHU 454 in humans led to the improved agent SHU 508
(Leovist®, Schering AG, Berlin), wherein α-D-galactose is
milled with 0.1% palmitic acid [8]. However, in spite of
more persisting contrast on the arterial side of the heart with
SHU 508 [9], it is not in clinical use to study early reduced
blood flow in the myocardium using MCE. Since pre-
formed, lyophilized phospholipid-stabilized gas bubbles on
the market have been a success [10], the simplicity of the
gas-releasing carbohydrate-based products in the form of a
dry, particulate, water-soluble precursor of gas bubbles
encouraging further research. In the present research, our
first aim was to investigate if lifetime of the populations of
gas bubbles released could be prolonged, by investigating a
broad range of amphiphilic compounds. Our second aim
was to investigate if acoustic attenuation vs. time (390 s)
area under the curve could be used as a suitable screening
tool in this research.

Experimental Section

Materials

Chemicals

Starch was purchased from Reppal PSM 70, Reppe Glykos
(Sweden). α-D-Xylose was purchased from BDH (Basel,
Switzerland). 1,2-dipalmitoyl-sn-glycero-3-phosphati-
dylcholine (DPPC) and 1,2-dioleoyl-sn-glycero-3-phos-
phatidylethanolamine (DOPE) were purchased from Avanti
Polar Lipids, USA. Human serum albumin (HSA), sodium
dodecyl sulfate (Sodium lauryl sulfate, Na-LS), 2-sul-
fobutanedioic acid 1,4-bis(2-ethylhexyl) ester sodium
salt (Na-docusate), Pluronic® F68 (Mw 8.4 kDa) a very
hydrophilic polyoxyethylene polyoxypropylene triblock
copolymer (abbreviated HTBCP), sorbitan mono-oleate
isomer mixture (Span 80) abbreviated SMO, sorbitan tri-
oleate isomer mixture Span 85 abbreviated STO, α-lino-
lenic acid ((Z,Z,Z)-9,12,15-octadecatrienoic acid) and other
fatty acids were purchased from Sigma Chemical Company
(St. Louis, MO). Tricosa-10,12-diynoic acid was purchased
from ABCR GmbH & Co. (Karlsruhe, Germany). Hexa-
decanedioic acid, α-cyclohextrin, dextran (Mw 20 kD),
maltodextrin, glycogen, α-D-galactose and all other chem-
icals were purchased from Fluka Chemie AG (Buchs,
Switzerland) or E. Merck AG (Darmstadt, Germany).
Scheme 1 gives the structures of the used compounds
except for the oligo and polysaccharides. Deionized water
was used throughout the experiments. Chemical structures
and names of the compounds used in the present work are
given in Scheme 1.

Visual Inspection of the Products After Suspension
in the Carrier Liquid

Screening of the rough ability of the products to release gas
microbubbles was performed using conventional light
microscopy.

Area Under the Curve of Acoustic Attenuation
and Bubble Half-Time

A carrier liquid for determining the acoustic attenuation of
the products consisted of 10 mL of propylene glycol mixed
with 90 mL of 5% dextrose in water. Each product (1.0 g)
was dispersed in the carrier liquid (3.0 mL) and gently
handshaken for 15 s. The resulting mixture was added to
5% human serum albumin in phosphate buffer (52 mL).
The mixture was placed in a measurement cell with a
5 MHz broadband transducer mounted on one side wall,
and an ultrasound-reflecting plate on the opposite side of
the cell. A pulse-reflection technique measuring the
acoustic transmission through the product dispersions was
used to calculate the acoustic attenuation taken as the
inverse value of the measured acoustic transmission. The
acoustic attenuation (dB/cm) was plotted as a function of
time for 390 s. Temperature in the measurement cell was
37 ± 1 °C with constant circulation of the liquid. The
results were normalized with regards to measurements of a
reference consisting of 55 mL of 5% human serum albumin
buffer solution. The calculated integral of the area Α
under the curve of acoustic attenuation vs. time (390 s),
was denoted A390. This parameter expressed the total
amount of gas phase in the dispersions up to and including
390 s. Average bubble lifetime was characterized by the
half-time (t½) of the acoustic attenuation up to 390 s. Both
A390 and t½ were necessary to decide if persistence of
ultrasound contrast effect in vitro was improved for the
various formulations compared with the reference stan-
da
d

Scheme 1 gives the structures of the used compounds
except for the oligo and polysaccharides. Deionized water
was used throughout the experiments. Chemical structures
and names of the compounds used in the present work are
given in Scheme 1.
Particle Size Distribution

The particle size distribution of selected products was analyzed using a Coulter Counter LS 100 or a Malvern Mastersizer light-scattering apparatus by suspending 2 g of the product using a vibrating screen before measuring.

Monitoring of Amphiphiles

The surfactants, phospholipids and fatty acids milled with the carbohydrates were monitored by analyzing the product (1.0 g) in carrier liquid (3 mL) using thin layer chromatography (TLC) and detecting spots using Nile blue fluorescence and copper complexation as described in the literature [11].

Scheme 1 Chemical structures and names of the compounds used. The structures of starch, α-cyclodextrin, dextran Mw 20 kD, maltodextrin and glycogen are omitted for clarity.
log $P$ Values

The concept of hydrophilic-lipophilic balance (HLB) is based on calculations of the ratio of hydrophilic and lipophilic groups in the molecules [12]. However, HLB values approach zero for many of the compounds used in the present study with low water solubility. Therefore calculated log $P$ was found to be a more useful parameter characterizing the amphiphiles used herein. The log $P$ values of the amphiphilic compounds were calculated using the chemist drawing and calculation software ChemBioDraw, version 11.0.1, CambridgeSoft Europe Office, 1 Signet Court, Swanns Road, Cambridge CB5 8LA UK.

Data Analysis

The parameters defined below were used (abbreviations in parentheses) in a partial least squares analysis (PLS) model with $A_{390}$ as a response parameter: % (w/w) amphiphilic material added (% amph); calculated log $P$ of the amphiphiles (log $P$); ionic character in sodium salt form (ionicity: 0 or 1); mean particle size in freshly prepared suspensions (psize); % (w/w) $\alpha$-D-galactose (%Gal); molecular weight of the amphiphilic additive (LipidMw); degree of unsaturation in the additives as number of double bond equivalents (Unsat: 1–4); number of rings in the chemical structure of the amphiphilic (rings); presence of carboxylic acid groups (Carboxyl: 0 or 1). The PLS analysis was performed using the computer program Unscrambler, version 7.01 from CAMO Software AS, Oslo, Norway.

Preparation of Products

All concentrations are given as % w/w in the procedures. Two literature procedures for preparation of carbohydrate-based gas-releasing ultrasound contrast agents [7, 13] were modified to establish the three general preparation procedures below. Ball-milling was performed with a Retsch centrifugal ball-mill in a stainless steel ball-mill having a 50 mL grinding cup and 3 × 20 mm balls for 10 min. All products appeared as white powders after milling.

General Procedure I: Mixtures of Carbohydrates Without Amphiphiles Added, Products 1–4

Products (1) and (2) consisted of commercial qualities of $\alpha$-d-galactose (1) and $\alpha$-d-xylose, respectively, that were ball-milled. The $\alpha$-d-galactose/Starch Mixtures (3a-c) were prepared by mixing $\alpha$-d-galactose (3.2, 2.0 and 0.8 g) with starch (0.8, 2.0, and 3.2 g, respectively) followed by ball-milling. The $\alpha$-d-galactose/Dextran (Mw 20 kD) mixtures (4a-b) were prepared by combining 25.8% solutions of $\alpha$-d-galactose in water (29.2 and 19.4 g) with 25.7% solutions of Dextran in water (9.7 and 19.4 g, respectively). The combined solutions were evaporated to dryness while stirring under reduced pressure (10 Torr, 40 °C) and dried in a desiccator overnight. The products were then ball-milled. Experimental data and results for products 1–4 are given in Table 1.

General Procedure II: $\alpha$-d-galactose-Based Products with Surfactants (5–11) and Lipophilic Acids (12–19)

For each of the products 5–19, the amphiphilic additives were each dissolved in 96% ethanol or water at 50–78 °C. The resulting solutions were filtered and then added to 1 (24.2 g) under stirring. The resulting mixtures were evaporated to dryness under reduced pressure (10 Torr, 40 °C) and the resulting solid products were dried in a desiccator overnight. The residues were ball-milled to give the final products. Experimental data and results for products 5–19 are given in Tables 2 and 3.

General Procedure III: Palmitic Acid Added to a Mixture of $\alpha$-d-galactose (1) and Starch (20) and Carbohydrates Other than $\alpha$-d-galactose (21–24)

For 20, $\alpha$-d-galactose (1, 12.1 g) was heated to 60 °C and mixed with a 14.3% w/w starch solution in water (35.0 g) before addition of the fatty acid solution. For products 21 and 22, $\alpha$-d-galactose (1) was replaced by 41.3% w/w solutions (24.2 g) of $\alpha$-d-xylose and maltodextrin, respectively. For 23 and 24, $\alpha$-d-galactose (1) was replaced by 22.5% w/w solutions (22.2 g) of glycogen and $\alpha$-cyclo-dextrin, respectively. The resulting mixtures were evaporated to dryness under reduced pressure (10 Torr, 40 °C). The resulting solid products were dried in a desiccator overnight and the residues were ball-milled to give the products. Experimental data and results for products 20–24 are given in Table 4.

| $\alpha$-d-galactose + carbohydrate (C2) | Prod. | % C2 | $A_{390}$ | $t$ 1/2 (s) |
|----------------------------------------|-------|------|----------|-------------|
| $\alpha$-d-galactose (reference 1, C1) | 1     | 0    | 5        | 16          |
| $\alpha$-d-xylose                      | 2     | 0    | 3        | 0           |
| $\alpha$-d-galactose + Starch          | 3a    | 20   | 164      | 30          |
|                                       | 3b    | 50   | 161      | 35          |
|                                       | 3c    | 80   | 70       | 38          |
| $\alpha$-d-galactose + Dextran         | 4a    | 25   | 23       | 54          |
|                                       | 4b    | 50   | 13       | 48          |

Milled $\alpha$-d-galactose (1) is reference product 1 (RC1) and corresponds to the commercial product Echovist®. Carbohydrate 1 is milled $\alpha$-d-galactose or $\alpha$-d-xylose. The added carbohydrate is denoted C2.
Results and Discussion

Results

The particle size distributions of the solid products 1–24 were in the range of 1–20 μm (Coulter Counter, data not given). In example, a typical accumulated particle volume distribution of 18b showed that 90% was <4.4 μm and 10% was <0.5 μm, the mean diameter was 2.2 μm and the median diameter was 2.1 μm. Based on visual microscopy observation, the solid particles tended to release air bubble dispersions in the size range of 1–15 μm, with some larger bubbles.

For product comparison, two reference compounds were needed: according to the literature and product descriptions [7, 13, 14], milled α-D-galactose (1) corresponds to the commercial product Echovist®/C210 and was chosen as reference product 1 (denoted RC1). According to the literature [8] and public information, the product consisting of α-D-galactose milled with 0.1% palmitic acid (13a) corresponds to the commercial product Levovist®/C210, and was chosen as reference product 2 (denoted RC2). The results of the acoustic characterization expressed as $A_{390}$ are given in Tables 1, 2, 3 and 4. The results of the measurement of $A_{390}$ after mixing with the carrier liquid are given in Figs. 1, 2, 3, 4 and 5.

Table 2 $A_{390}$ for products 5–11 comprising α-D-galactose, milled with amphiphilic compounds lacking a carboxylic acid functionality

| Amphiphile A with trivial name (formula, Mw, log $P$) milled with α-D-galactose | Prod. % | $A_{390}$ | $t/2$ (s) |
|---|---|---|---|
| DPPC (C$_{40}$H$_{81}$NO$_{8}$P, 736.1, 9.8) | 5a | 0.1 | 366 | 53 |
| | 5b | 1.0 | 3,712 | >500 |
| Na-LS (C$_{12}$H$_{25}$NaO$_{5}$S, 266.4 (anion), 1.8) | 6a | 0.1 | 22 | 23 |
| | 6b | 1.0 | 67 | 66 |
| SMO (C$_{24}$H$_{46}$O$_{7}$, 446.6, 5.0) | 7a | 0.1 | ≈0 | 0 |
| | 7b | 1.0 | 34 | 26 |
| STO (C$_{60}$H$_{110}$O$_{9}$, 975.5, 21.2) | 8a | 0.1 | 6 | 34 |
| | 8b | 1.0 | 28 | 72 |
| HTBCP (Average Mw 8,400, log $P$ n.a.) | 9a | 0.1 | 55 | 23 |
| | 9b | 1.0 | 74 | 23 |
| Na-Docusate (C$_{20}$H$_{37}$NaO$_{7}$S, 422.6 (anion), 6.1) | 10a | 0.1 | 22 | 23 |
| | 10b | 1.0 | 67 | 66 |
| DOPE (C$_{44}$H$_{86}$NO$_{8}$P, 744.1, 13.6) | 11a | 0.1 | 343 | 500 |
| | 11b | 1.0 | 552 | 72 |

Table 3 $A_{390}$ values of products 12–19 comprising α-D-galactose, milled with amphiphilic compounds possessing a carboxylic acid functionality

For each product, abbreviation or trivial name of the carboxylic acid C is followed by the formula, molecular weight and calculated log $P$ in parentheses. In the last three columns, amounts of A, $A_{390}$ values and half-times are given. The chemical names of the amphiphiles are given under "Materials and Methods"

| Carboxylic acid C, trivial name (formula, Mw, log $P$) milled with α-D-galactose | Prod. % | $A_{390}$ | $t/2$ (s) |
|---|---|---|---|
| Capric acid (C$_{10}$H$_{22}$O$_{2}$, 172.3, 4.0) | 12a | 0.1 | ≈0 | 0 |
| | 12b | 1.0 | ≈0 | 0 |
| Palmitic acid (C$_{16}$H$_{34}$O$_{2}$) reference 2 (RC2) = 13a (0.1% palmitic acid) | 13a | 0.1 | 357 | 31 |
| | 13b | 0.2 | 871 | 35 |
| | 13c | 1.0 | 1,061 | 107 |
| Hexadecane-dioic acid (C$_{16}$H$_{34}$O$_{4}$, 286.4, 5.1) | 14 | 1.0 | 474 | 40 |
| Linolenic Acid (C$_{18}$H$_{30}$O$_{2}$, 278.4, 7.3) | 15 | 1.0 | 401 | 79 |
| Behenic acid (C$_{22}$H$_{44}$O$_{2}$, 340.6, 9.9) | 16a | 0.1 | 2,745 | >500 |
| | 16b | 1.0 | 4,653 | >500 |
| 10,12-tricosa-diynoic acid (C$_{23}$H$_{32}$O$_{2}$, 346.6, 9.9) | 17 | 0.2 | 884 | 51 |
| Melissic acid (C$_{30}$H$_{60}$O$_{2}$, 452.8, 13.8) | 18a | 0.01 | 322 | 150 |
| | 18b | 0.1 | 4,095 | >500 |
| | 18c | 1.0 | 4,564 | >500 |
| 5β-cholanic acid (C$_{24}$H$_{40}$O$_{2}$, 360.6, 7.7) | 19a | 0.01 | 162 | 47 |
| | 19b | 0.1 | 1,962 | 203 |
| | 19c | 1.0 | 4,048 | >500 |
Figures 6 and 7 show the statistical analysis of the results, using $A_{390}$ as a response parameter. A high log $P$, the presence of $\alpha$-D-galactose and that of a carboxylate group in the amphiphilic molecule correlates positively with $A_{390}$. In addition, the PLS analysis reveals some interaction effects between parameters. There is a strong interaction between log $P$ and the presence of a carboxylate group; an interaction between log $P$ and the presence of $\alpha$-D-galactose is also evident from the analysis.

Discussion

As shown in Figs. 1 and 2, the use of other carbohydrates than $\alpha$-D-galactose does not improve performance ($A_{390}$ and $t/2$) as compared with the $\alpha$-D-galactose-based reference products RC1 and RC2. The observed significant increase in both $A_{390}$ and $t/2$ when $\alpha$-D-galactose is mixed with 0.1% of $\beta$-cholanic acid (19b), behenic acid (16a), and melissic acid (18b) compared with RC2 (Fig. 5), correlates well with the log $P$ values for the acids. The products 16a and 18b show half-time $t/2 > 500$ s compared with $t/2 = 31$ s for RC2 (Table 3). For instance, log $P$ for 5$\beta$-cholanic, behenic and melissic acids are calculated to be 7.7, 9.9 and 13.8, respectively. The $A_{390}$ values for the corresponding products 19b, 16a and 18b are 1,962, 2,745 and 4,095, respectively. When the amount of these fatty acids is increased to 1% (products 16b, 18c and 19c, Table 3, Fig. 4) the correlation of $A_{390}$ and $t/2$ with log $P$ is even more pronounced.

An interesting comparison is the results for the products 13c, 14 and 15, with 1% (w/w) palmitic acid, hexadecanedioic acid and linolenic acid, respectively (Table 3). Both $A_{390}$ and $t/2$ are more than 100% higher for 13c

| Table 4 | $A_{390}$ values for products 20–24 different carbohydrates milled with 0.2% palmitic acid |
|---------------------------------|-----------------|-----------------|-----------------|
| Carbohydrate milled with        | Product          | $A_{390}$       | $t/2$ (s)       |
| palmitic acid                   |                  |                 |                 |
| $\alpha$-D-galactose + Starch   | 20               | 142             | 152             |
| $\alpha$-Xylose                 | 21               | 706             | 245             |
| Maltodextrin                    | 22               | 254             | 225             |
| Glycogen                        | 23               | 358             | >500            |
| $\alpha$-Cyclodextrin           | 24               | 94              | 47              |

Fig. 1 Acoustic attenuation (dB/cm) vs. time (390 s) of carbohydrate mixtures without amphiphiles added, compared with the reference standard 1 (RC1), milled $\alpha$-D-galactose (1). When no amphiphiles are present, mixing of $\alpha$-D-galactose with polymeric carbohydrates like starch or Dextran with $Mw$ 20 kD only slightly increases $A_{390}$ compared with RC1.

Fig. 2 Acoustic attenuation (dB/cm) vs. time (390 s) of different carbohydrates with 0.2% palmitic acid compared with the reference standard 2 (RC2), milled $\alpha$-D-galactose with 0.2% palmitic acid (13b). Adding palmitic acid to different carbohydrates shows that $A_{390}$ of RC2 is superior for the first 100 s compared with the products containing maltodextrin, glycogen and $\alpha$-cyclodextrin with the same amount of palmitic acid.

Fig. 3 Acoustic attenuation (dB/cm) vs. time (390 s) of $\alpha$-D-galactose with 1% of different amphiphiles, surfactants and linolenic acid compared with $\alpha$-D-galactose containing 1% palmitic acid. The intensity and duration of the acoustic attenuation of all products containing 1% amphiphiles with various log $P$ values, but lacking carboxylic acid functionality, is significantly lower than for RC2.

Fig. 4 Correlation of $A_{390}$ and $t/2$ with log $P$ for the products 16b, 18c and 19c, Table 3, Fig. 4. The correlation of $A_{390}$ and $t/2$ with log $P$ is even more pronounced.
compared with 14. The dicarboxylic acid 14 will need to bend its lipophilic chain at the gas–water interface, reducing the lipophilic interaction. For linolenic acid (15), a highly unsaturated carboxylic acid with a log P comparable to palmitic acid, A390 is at the level of 14 and with a half life of only 79 compared with 107 for 13c (Table 3 and Fig 3). This may be due to the high number of cis double bonds in linolenic acid, giving a non-linear conformation of the lipophilic chain, reducing the hydrophobic interactions between the hydrocarbon chains [15].

The PLS analysis illustrated by the regression coefficient plot (Fig. 6) provides further support for the trends observed in the A390 plots. An absolute high and stable regression coefficient for many principal components indicates a robust effect of the actual descriptor (Fig. 6). The strongest positive correlation is shown between amphiphiles possessing a carboxylic acid functionality and high log P added to α-D-galactose. This points at fatty acids with a number of carbon atoms >20, in good compliance with the experimental data. These data are further strengthened by the experimental results for amphiphiles with high log P but lacking the carboxyl functionality; all have low A390 (e.g. the phospholipid-based product 11). Mw alone has negligible effect on A390 as shown by its regression coefficient (Fig. 6). The good correlation between predicted and observed A390 values (Fig. 7) shows that a suitable model explaining the main effects on A390 has been established. The reliability of the PLS estimated trends was also tested by regressing various logarithmic transformations of the response having a more uniform variation than the original A390, and unimportant regression coefficients were successively removed. This process gave similar regression coefficients patterns as shown in Fig. 6 regarding the most important parameters affecting A390 as well as t/2.

Ultrasonic waves are heavily attenuated at gas–water interfaces [1]. It has earlier been reported [16] that a linear correlation between ultrasonic attenuation and interfacial area in a gas–liquid bubble column could be used to estimate the total interfacial area in the system. The same acoustic phenomenon was used in the present study to establish a method to monitor the amount and lifetime of bubble populations generated by milled carbohydrates.

The amphiphiles in the present study with the highest log P are practically insoluble in water [17]. Thus, when the ethanolic solutions of the amphiphiles are added to the aqueous carbohydrate solutions (products 5–19), colloidal suspensions may form. It was shown in a study of fatty acid particle formation in water containing a surfactant [18] that the particle size of such suspensions depended on the chain length of the fatty acid. Gas bubbles have an affinity for lipophilic particulate surfaces providing stabilization of the gas phase [19–22], providing one possible mechanism for the observed bubble stabilizing effect during the dissolution of the carbohydrate-amphiphile admixtures. However, a more plausible mechanism is the formation of fatty acid Langmuir films at the gas–water interface, systems extensively described in the literature [17, 23–26]. If the amphiphilic compounds are spread on the gas–water interface, reduction of surface tension would result, as shown by isotherm studies in the cited literature. Reduced surface tension would increase bubble stability, explaining the observations in the present study. It is also well known from reports of investigation of aerosol systems that fatty acids with high log P tend to be organized at the gas–liquid as Langmuir films [17]. In that study, it was shown that chain length and the carboxylic acid functionality were important for the interfacial lifetime of the Langmuir films. One question arising is how the fatty acids with the highest log P and very limited solubility in water could achieve a
dissolved state to form layers at the gas–water interface in the carbohydrate formulations. The ultrasonic influence in the present experiments may aid the dissolution process, as described in literature for systems comprising substances with low water-solubility [27].

Conclusion

In conclusion, the response parameters $A_{390}$, expressing the amount of gas-phase present during 390 s, and the bubble half-time $t^{1/2}$ was shown to be useful main parameters in a screening method for increased persistence of the ultrasound contrast effect in vitro. Using these parameters, it was shown that the persistence and amount of released gas bubbles in formulations used in the commercial carbohydrate-based, gas-releasing available and improved ultrasound contrast agents (Echovist® and Levovist® respectively) could be improved more than 10 times using fatty acids with a higher log $P$ relative to palmitic acid. Saturated fatty acids with chain length higher than 20 carbon atoms are commercially available, are found in many nutritional products and should find convenient use in pharmaceutical products. Specifically, the increases in $A_{390}$ and $t^{1/2}$ shown for product 19b-c, and especially products 16a-b and 18b-c containing 5β-cholanic, behenic and melissic acid, respectively, are results that encourage further research and in vivo studies. Studies are ongoing to investigate the in vivo properties of the most promising test substances.

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