Acoustic cavitation and ionic liquid combined: A modelling investigation of the possible promises in terms of physico-chemical effects †

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† Presented at The 4th International Electronic Conference on Applied Sciences

Abstract: The present work is based on a mathematical model describing a single acoustic cavitation bubble oscillating under an ultrasonic field of 200 and 300 kHz and an acoustic amplitude of 1.8 atm within 1-Butyl- 3-methylimidazolium Acetate. The model integrates the dynamics of bubble oscillation, the thermodynamics applied to the interior of the bubble and at its interface, as well as the sonophysical and sonochemical events occurring in the presence of dissolved cellulose in the ionic liquid. The performed simulations shed light on the major physical effects of acoustic cavitation, namely shockwave and microjet, as well as the sonochemical effects in terms of the degradation rate of the dissolved cellulose in the secondary reactional site, i.e., the interface. The predominance of the effects and its dependency on the acoustic frequency is tackled from an energetic point of view, it has been demonstrated that 300 kHz offers the lowest heat flow across the bubble interface, lowering the chances for the sonochemical degradation of cellulose, whilst 200 kHz offers a significant degradation rate attaining 71.4 mol.dm$^{-3}$.s$^{-1}$, and harsher microjets and shockwaves with powers of 3300 and 900 mW at the collapse, respectively.

Keywords: Ultrasounds; Acoustic cavitation; [C4mim][CH3COO]; Cellulose; Temperature; Pressure; Sonochemistry; Microjet; Shockwave.

1. Introduction

Several researchers have described the dissolution of cellulose in ionic liquid and have offered diverse molecular theories. Most of them agreed that cellulose dissolved when anions made hydrogen bonds with it, but others demonstrated that cellulose dissolves in ionic liquids when an anion and a cation make hydrogen bonds with the hydrogen and oxygen atoms of the cellulose, which are precisely positioned between the C6 and C3 hydroxyl groups of the nearby cellulose chain [1]. Besides, The exposure of the ionic liquid to ultrasounds revealed according to several studies some interesting physical and chemical effects that may be beneficial in terms of cellulose degradation [2]. Nonetheless, the joint investigation of the sonophysical and sonochemical effects with this configuration has not been performed yet, particularly through modeling and simulation. The present paper proposes a numerical model describing the behavior of single acoustic cavitation bubble within [C4mim][CH3COO] in the presence of dissolved cellulose, with the aim to evaluate the chemical and physical impact of sonication, under the specific acoustic frequencies of 200 and 300 kHz, generally adopted in sonochemistry.

2. Numerical model

The 1-Butyl-3-methylimidazolium Acetate [Camin][CH3COO] is an ionic liquid characterized by a high cellulose solvation power [3,4]. In the present model, the gaseous medium is saturated with argon. Despite the fact that only a limited number of studies examined the solubility of argon in ionic liquids [5–8], the reported solubility rates are judged sufficient to create gaseous inclusions and generate acoustic cavitation by heterogeneous nucleation process [9] when enough tensile strength is induced [10].

The ionic liquid is submitted to ultrasonication at 200 and 300 kHz of frequency, and an acoustic amplitude of 1.8 atm, which is the lowest value of acoustic amplitude (under the lowest temperature) that allows acoustic cavitation phenomenon and consequently the expected chemical effects. The operating temperature is fixed at 353 K in order to

Citation: To be added by editorial staff during production.

Academic Editor: Firstname Lastname

Published: date

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attain an acceptable value of dynamic viscosity of the ionic liquid. The ambient radii at
the respective acoustic frequencies are taken as 5 and 3.5 µm. The oscillation of the bubble
is governed by the Keller-Miksis equation as indicated in Eq. 1.

\[
R = \frac{(c + R)}{\rho_L(cR - RR)} + 4\mu \left( \frac{P_g - 26}{R} - 4\mu \frac{\dot{R}}{R} + P_o + P_a \sin \left( 2\pi f \left( t + \frac{R}{c} \right) \right) \right) + \frac{R}{\rho_L(cR - RR)} + 4\mu \left( \frac{\dot{R}}{R^2} + \frac{26R}{R^2} + 4\mu \frac{\dot{R}^2}{R^2} \right) - \frac{\rho_L(3c - R)\dot{R}^2}{2(\rho_L(cR - RR) + 4\mu)}
\]

According to [12], Safarov et al. [11] provided the physical parameters of 1-Butyl-3-
methylimidazolium acetate at 353 K in relation to temperature. These properties include
surface tension \( \sigma \), dynamic viscosity \( \mu \), density \( \rho_L \), and sound speed \( c \).

Since 1-Butyl-3-methylimidazolium acetate has a very low vapor pressure [13], argon
is assumed to be the only gas present in the bubbles. The initial pressure of the gas inside
the bubble, as determined by the mechanical equilibrium condition, is given by Eq. 2.

\[
P_{\text{eq}} = P_o + \frac{2\pi}{R_0}
\]

The energy balance applicable to the acoustic cavitation bubble considering its oscilla-
tion is given in Eq.3. It allows the determination of the variation and distribution of
the temperature as a function of time and space.

\[
\dot{T} = \frac{1}{nC_v(T)} (-P_g 4\pi R^2 \dot{R} - \varphi(t))
\]

\[
\varphi(t) \text{ represents the heat flow passing through the thermal boundary layer of width } \xi, \text{ its expression is based on the continuity of the thermal flow across the bubble interface at each instant } t, \text{ as seen in Eq.4.}
\]

\[
\varphi(t) = \frac{4\pi R(R - \xi)}{\xi} \int_{\tau(t)}^{T(t)} \lambda(T) dT
\]

The isochoric molar heat capacity of argon \( C_v(T) \) and its thermal conductivity \( \lambda(T) \),
appearing respectively in Eqs. 3 and 4, vary in function of the temperature of the bulk
volume of the bubble. Both are expressed as polynomial form as shown in [14].

The pressure of gases inside the bubble volume is expressed using the Van der Waals
state equation as shown in Eq.5.

\[
\left( \frac{P_g + \frac{n^2a}{V^2}}{V - nb} \right) = nR_gT
\]

Due to the ionic liquid’s high order of magnitude thermal conductivity, which is almost
ten times higher than that of argon [15,16], the resistance to thermal diffusion is
found on the gaseous side. As a result, the boundary layer of thermal diffusion is taken
into account on the gaseous side; Eq. 6 [17] gives its width \( \xi \). This equation includes the
thermal diffusivity \( \kappa \), which is temperature-dependent and depends on the density, heat
capacity, and thermal conductivity of argon.

\[
\xi = \min \left( \frac{Rk}{|R|}, \frac{R}{\pi} \right)
\]

With the exception of the thin boundary layer with a width \( \xi [18] \), it is assumed that
the temperature is spatially consistent throughout the majority of the bubble. Resolving
Eq. 7 yields the radial distribution of temperature throughout the thermal boundary layer
(the result of the integration of Eq. 4) for each value of \( r \) comprised between 0 and \( \xi ).

\[
1.79435 \times 10^5 T(R - r)^2 + 6.81277 \times 10^{-3} T(R - r) - 4.6408 = -\frac{r \varphi(t)}{4\pi R(R - r)}
\]

The thermal effect caused by the hot spots at the bubble interface (second reactivity
site), according to the two sites model [2], constitutes the seat of chemical reactions. The
bubble’s main body is thought to be chemically inert. The cellulose is anticipated to ther-
ma!y disintegrate in accordance with the process put forward Diebold [19] and shown in
when the bulk temperature is high enough and the surrounding temperature is adequate to start pyrolysis reactions.

According to Table 2, the chemical kinetics occurring in the second reactivity site is described as a chain mechanism of first order reactions, with the molar yield of each species \( k \) being controlled by Eq. 8.

\[
\frac{dn_k}{dt} = V_r \sum_{i=1}^{7} (\theta_{ki} - \vartheta_{ki}) A_{i} e^{-\left(\frac{E_i}{RT}\right)} \prod_{k=1}^{K} (c_k)_{\theta_{ki}}
\]

(8)

In this equation, \( T_r \) constitutes the temperature governing the chemical kinetics of the decomposition of cellulose. This temperature varies spatially as a function of the bubble radius \( r \), as shown in Eq. 7, the median value is considered to approximate the conditions of the thermal decomposition of cellulose. The molar concentrations \( c_k \) of each chemical species \( k \) involved in the mechanism are reported to the volume of the thermal boundary layer of width \( \xi \) shown previously, this volume is denoted \( V_r \), and is given in Eq. 9.

\[
V_r = \frac{4}{3} \pi \left( R^3 - \left( R - \min \left( \frac{R K}{R \pi} \right) ^3 \right) \right)
\]

(9)

When the bubble collapses, a spherical shockwave is generated around the bubble in the liquid medium. This shockwave can provoke changes in the transport properties, by inducing dispersion of molecules in the liquid phase and consequently promoting mass transfer. The power emitted by the single acoustic cavitation bubble in the form of a shockwave \( P_s \) is expressed by Eq.10.

\[
P_s = \frac{4\pi \rho_{el} (R^2 \ddot{R} + 2R \dot{R}^2)^2}{c}
\]

(10)

Additionally, during the bubble collapse near to a solid surface (cellulose particles), the bubble loses its sphericity. A modification of the shape of the bubble is observed from the original approximate sphere to a slender shape and then flattened shape. At the final stage, a microjet breaks the bubble wall near the solid and impacts towards it. Micro-jet velocity is given by Eq.11 [20].

\[
v = 8.97 \left( \frac{R}{R_0} \right)^2 \sqrt{ \frac{P_{w} + P_{A} \sin \left( 2\pi ft \right) - P_{v}}{\rho_{el}}} - \frac{1}{8} \pi \rho_{el} h^2 v^3
\]

(11)

The power of the exit \( P_j \) is then shown in Eq.12 as a function of the jet velocity \( v \) and the jet nozzle exit diameter \( h \) (taken as the third of the value of the bubble diameter) [21,22].

\[
P_j = \frac{1}{8} \pi \rho_{el} h^2 v^3
\]

(12)

The system of non-linear and dependent differential equations (Eqs.1, 3 and 8) is simultaneously resolved with the non-linear equations 4 to 9 using Ode23s solver of Matlab. The shockwave and microjet powers are then assessed according to Eqs. 10 and 12.

3. Results and discussion

3.1. Dynamics of oscillation of the acoustic cavitation bubble

Figure 1(a) reports the evolution of the argon bubble radius within the sonicated ionic liquid [C₂min][CH₃COO] under both 200 and 300 kHz frequencies.

Figure 1. Bubble radius (left (a)) and wall velocity (right (b)) as functions of reduced times under 200 and 300 kHz.
The results demonstrate that 200 kHz frequency induces a harsher oscillation of the bubble wall, achieving a maximum radius of almost 13 µm, with an expansion ratio of 2.6, versus 2.18 under 300 kHz. During the collapse phase, the bubble contracts 3.81 folds under 200 kHz, and 3.45 folds under 300 kHz. This trend is confirmed by the evolution of the bubble wall velocity reported in Figure 1(b), attaining two extremums of 36.5 and 20 m/s at expansion and collapse, respectively, under 200 kHz.

### 3.2. Thermodynamics of the acoustic cavitation bubble

The evolution of the bulk and interfacial temperatures is reported in Figure 2. As predicted from the dynamics of the bubble oscillation, the highest temperature is attained under 200 kHz, with 1600 K in the bulk volume and 1000 K at the interface, at the moment of collapse. 300 kHz leads to barely 474 K at the interfacial zone, which limits the sonochemical activity there under this acoustic condition.

![Figure 2](image1.png)  
**Figure 2.** Bulk (left (a)) and interfacial (right (b)) temperatures as functions of reduced time under 200 and 300 kHz.

As a result of the radius and temperature evolutions reported earlier, we notice that the highest bulk pressure is attained under 200 kHz with a value of 48 bar. Through the bulk pressure does not directly affect the sonochemical process of cellulose transformation, it shows that 200 kHz frequency is more favorable for the sonochemical decomposition of cellulose within the sonicated ionic liquid. This point will be further discussed in the following section.

![Figure 3](image2.png)  
**Figure 3.** Bulk volume pressure under 200 and 300 kHz

### 3.3. Sonochemistry

Figure 4 reports the cellulose decomposition rate the molar yield of cellulose within one acoustic cycle under 200 kHz frequency. The decomposition rate attains its maximum at the collapse, with a value of 71.4 mol/dm$^3$.s. Though this rate lasts almost 0.05 µs, it allows the molar yield of cellulose to decrease by 9.77%.

![Figure 4](image3.png)  
**Figure 4.** Cellulose decomposition rate and molar yield under 200 kHz
3.4. Energetic reading: Heat transfer, Microjets and shockwaves

The results related to the thermal flow across the interface exhibit a significantly higher value under 200 kHz, which explains the higher interfacial temperature. The interface being the reactional site in the studied case, the prominent sonochemical activity observed under 200 kHz is explained thermally by the more important thermal flow, with a peak exceeding 3 W at collapse (almost 6 folds higher than at 300 kHz). In terms of the physical effects, 200 kHz results also in the harsher induced shockwaves and microjets, with respective peak values of 0.9 and 3.3 W at collapse. Hence, 200 KHz is both sonochemically and sonophysically more promising for the decomposition of cellulose in ionic liquid.

Figure 5. Thermal flow across the interface (upper left (a)), power of shockwave (upper right (b)) and power of microjet (down (c)) under 200 and 300 kHz.

4. Conclusion

In the present study, the sonochemical and sonophysical effects induced by sonication of the ionic liquid [C4mim][CH3COO] for the decomposition of cellulose are investigated numerically. The modelling and simulation results revealed that the harsher dynamics of bubble oscillation are achieved under 200 kHz, with expansion and collapse ratios of 2.6 and 3.81, respectively. The trend was confirmed by the bubble wall velocity, and resulted in higher temperature values both in the bulk volume of the bubble as well as the interfacial region, which is the sonochemically active zone in the studied case. The simulation of one acoustic cycle revealed that 9.77% of cellulose is decomposed under 200 kHz. The collapse is also accompanied of shockwave and microjet effects, characterized by peak powers of 0.9 and 3.3 W, respectively per bubble. Finally, 200 kHz sonication was demonstrated to be sonochemically and sonophysically effective for the sono-decomposition if cellulose dissolved in [C4mim][CH3COO].

**Funding:** This research received no external funding.

**Data Availability Statement:** No data was generated.

**Acknowledgments:** The author acknowledges the support given by the Ministry of Higher Education and Scientific Research in Algeria in the frame of the Project with Social-Economical Impact (PISE 2023) entitled “Green Processes for Sanitation”.
Conflicts of Interest: The authors declare no conflict of interest.

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