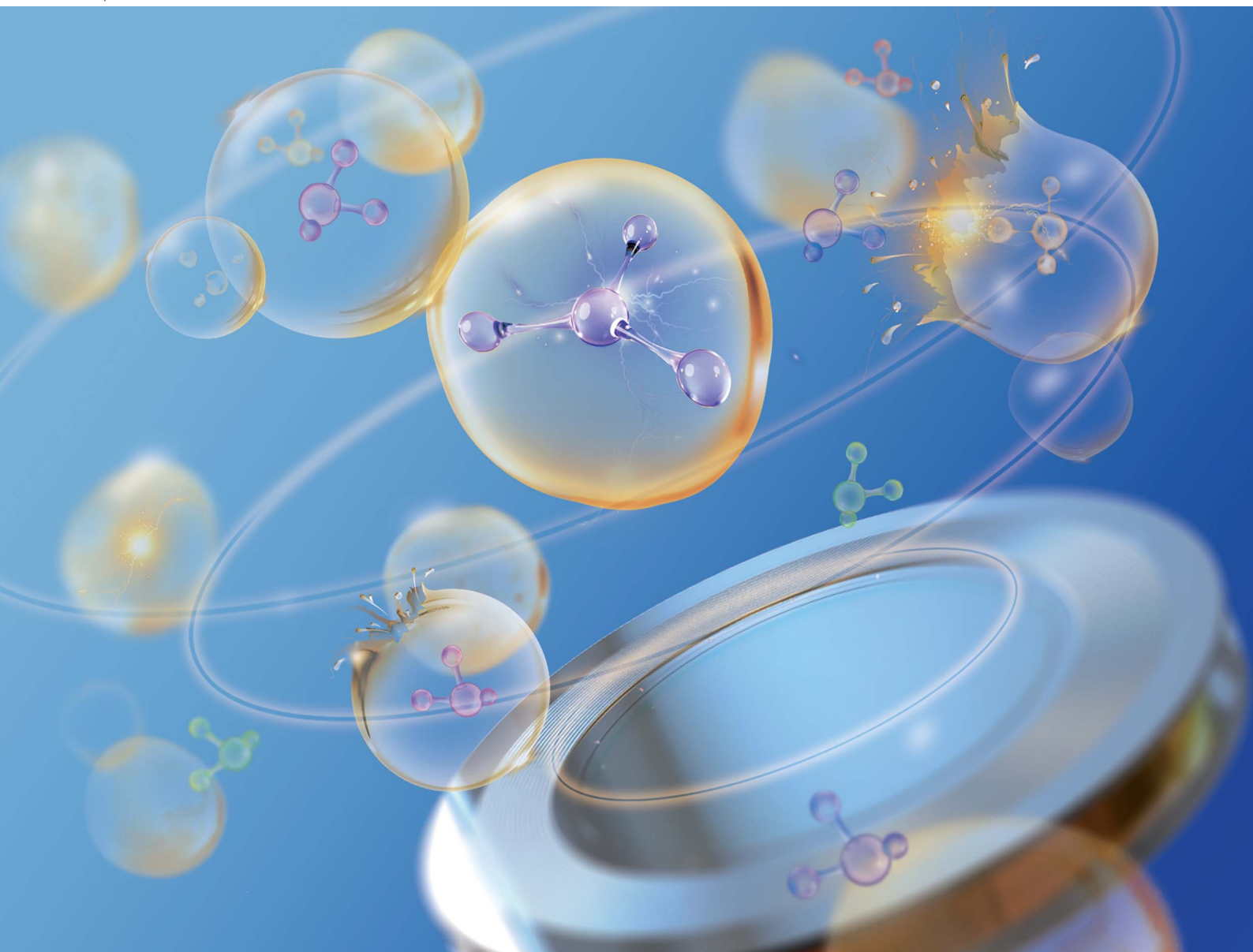


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

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Correlation of sonochemical activities measured *via*
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Correlation of sonochemical activities measured via dosimetry and an area-selective analysis of sono(chemi)luminescence†

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Ultrasonic industrial applications require theoretical support and practical guidance from a comprehensive understanding of sonochemical reaction dynamics. The influence of acoustic factors (frequency and pressure amplitude) and external parameters (liquid height) on sonochemical activity were researched. The phenomenon of sonoluminescence (SL), sonochemiluminescence (SCL) and potassium iodide (KI) dosimetry were investigated at 114 different settings. The settings included electrical loading-power of 10, 20, 30, and 40 W, 10 frequencies ranging from 22 to 2000 kHz, and reactor volumes of 200, 300, and 400 ml. A new area selection image processing technique was used to conduct a systematically quantitative analysis of SL and SCL across a broader frequency range. The sonochemical activity could be categorised into three zones based on the ultrasonic frequency (22 to 2000 kHz): $f < 200$ kHz, 200 kHz $\leq f \leq 1000$, and 1000 kHz $\leq f \leq 2000$ kHz. The Pearson and Spearman correlation coefficients were used to discuss the correlation between SL, SCL, reactive oxidant species (ROS) and hydrogen peroxide yields. The findings indicate that the influence of liquid height on cavitation activity within the reactor is mostly manifested in the power density. The ultrasonic oxidation capacity (as indicated by the yield of ROS) exhibits a strong positive relationship with SL intensity. A divergence of correlation between SL and I_3^- yield was observed. There was a lack of correlation between sonochemical activities (e.g. SCL and ROS yield). The poor correlation highlighted the importance of considering chemical mechanisms and reaction locations concerning the collapsing bubble.

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1. Introduction

The ultrasonic-driven cavitation bubbles go through generation, growth, and a quasi-collapse process, which can effectively produce extreme energies that can drive high temperatures (>800 K), high pressures (several hundred bar),¹ and generation of reactive oxidative species (ROS),^{2,3} and aqueous electrons.^{4,5} Moreover, acoustic cavitation not only involves radical chemistry but also induces significant mechanical effects, such as microjets, shock waves, and shear forces.⁶ These localized mechanical stresses resemble the action modes observed in mechanochemistry, giving acoustic cavitation the dual characteristics of sonochemistry and mechanochemistry. Therefore, sonochemistry can be regarded as a specialized form of mechanochemistry, as both share intersections in energy transfer mechanisms and reaction initiation pathways, offering new

possibilities for developing mechanically driven chemical reactions.⁷⁻⁹ These, ultrasonically-driven phenomena can contribute to degradation of organic pollutants and elimination of pathogenic microorganisms in water.^{5,10,11} Hence, acoustic cavitation is recognized as a chemical-free and green advanced water and wastewater treatment process (WWTP).^{5,12-16} Existing efforts have typically focused on the degradation and elimination of a single pollutant or microorganism under specific sonooperating conditions,^{15,17-19} or rudimentary comparison of ultrasonic processes with traditional non-ultrasonic processes. These works had already confirmed the effectiveness of ultrasonic method in green chemistry, and reported the sonopurification will be significantly influenced by cavitation effects and acoustic parameters.^{16,20} Nevertheless, such limited approaches lead to conclusions under specific parameters, which are not necessarily generally applicable and are often insufficient to be applied to optimised employment of acoustic technology.

The diameter of the transient cavitation bubble continues to grow until the energy from ultrasound can no longer be absorbed.²¹ When the bubble reaches a critical size, the force acting on it dominates, leading to a violent and quick collapse.²¹⁻²³ The fragmentation of the transient cavitation

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bubble is an important cause of acoustic cavitation phenomena and sonochemical reactions.^{24–26} Both are the significant mechanisms possibly responsible for microorganisms' destruction and chemical pollutant degradation.^{5,27–29} The sonochemical reaction and cavitation phenomena are significantly influenced by the various characteristics that affect the energy transfer within the ultrasonic system. The parameters include acoustic frequency, amplitude, energy transfer distance, as well as the viscosity of the liquid and the concentration and diffusion of gas.^{4,21,30,31} Furthermore, the dissipation and effectiveness of the ultrasonic energies are inherently problematic due to impedance-reducing energy transfer and existing secondary effects (*e.g.* flow, acoustic field attenuation, reactor corrosion, noise generation and overall mixing, *etc.*).^{14,22}

The resonance size of the bubbles is inversely proportional to the frequency applied to the acoustic field, with smaller bubbles being produced at higher frequencies.^{4,32} Due to the shortened oscillation cycle, collapse and fragmentation happen rapidly, more fragmented bubbles are produced in a short duration, the collapse intensity is reduced, and more ROS (*e.g.* hydroxyl ($\cdot\text{OH}$), hydrogen peroxide (H_2O_2)) are released out of the bubbles before undergoing any reaction.^{33–35} At lower frequencies, larger-sized bubbles are produced, collapse is strong, bubble surface instability is increased, and free radicals within the bubbles have enough time to follow reaction pathways analogous to flame chemistry.^{33,36} As a key component in organic pollutant removal during sonochemical process, the hydroxyl radical production and the afore-mentioned factors are not always positively correlated with acoustic field frequency.^{37–39} Many studies have shown the existence of optimal frequencies for ultrasound-introduced chemical processes to obtain the highest sonochemical efficiency.^{28,39–42} In general, the pressure amplitude is strongly related to the power transferred into the liquid and has effects on the number of cavitation bubbles and the strength of the collapse.⁴ Within the effective range of pressure amplitudes, higher amplitudes result in larger size bubbles.³⁶ The increase of compression and rarefaction forces can result in more violent collapse, which can contribute to sonochemical activity.^{43–45} The intensity of ultrasonic waves decrease with distance from emission source as they propagate through a medium. There are viscous interactions between liquid molecules while liquid molecules vibrate under the sonic waves.⁴⁶ Existence of reflection, refraction, diffraction and scattering of waves result in the attenuation of sound in liquid.²¹ In addition, the conversion of some amount of mechanical energy into heat can also lead to attenuation in the system, thereby affecting the sonochemical activity.^{21,47} For a plate transducer ultrasonic reaction system, the liquid height can reflect the maximum distance from the sound source.

Ultrasound has been extensively researched and employed as a valuable tool in green chemistry and WWTP, including a broad range of topics such as removing algae from water,^{48,49} water disinfection,¹³ and the degradation and removal of contaminants.^{50–52} Nevertheless, these studies have frequently been carried out with a focus on individual contaminants, and the acquired acoustic sets are highly specialised. The primary degradation process in WWTP is the presence of $\cdot\text{OH}$ in the

sonolysis system with degradation efficiency directly influenced by $\cdot\text{OH}$ concentration.⁵³ Hence, it is crucial to have a thorough and methodical comprehension of how acoustic parameters impact bubble behaviour and sonochemical activity, specifically the yield of ROS, as well as the interconnections between these factors. This information is essential for obtaining generalizable results and achieving effective ultrasound-assisted green chemistry and WWTP.

As mentioned, most studies on ultrasonic parameters have primarily investigated the impact of individual factors on the acoustic effect without considering the complex interplay of multiple variables. Thus, the presented work attempts to elucidate the impact of liquid height (reactor volume) and electric power on the ultrasonic cavitation effect throughout a wide frequency range. Three different cavitation characteristics were obtained from systematic experiments using 114 reaction settings with distinct parameter combinations. The chemical oxidation within the acoustic system was assessed by quantifying the production of ROS, including iodide oxidation radicals (IORS) and hydrogen peroxide (H_2O_2) yield. Spatial distribution and quantification of chemical activity were investigated through the sonoluminescence (SL) and sonochemiluminescence (SCL) for activity inside the bubbles or in the bulk, respectively. We present a systematic and intuitive demonstration of the dispersion of active bubbles and the efficiency of sonochemical oxidation under multi-parameters by modified hotspot maps and composite histograms. The relationship between different typical indicators of sonochemical activity was discussed by two correlation coefficients (Pearson's correlation coefficient and Spearman's rank correlation coefficient) for the first time. A new image processing method relying on luminescence area selection and greyscale value was introduced to address the issue of misleading data caused by photograph noise and the reflection of light from a glass reactor during the digitisation of SL/SCL images.

2. Materials and methods

2.1 Chemicals

All chemical reagents were direct used in the studies without undergoing further purification after acquisition. Luminol (5-amino-2,3-dihydro-1,4-phthalazinedione, purity $\geq 98.0\%$), ammonium molybdate tetrahydrate ($(\text{NH}_4)_6\text{Mo}_7\text{O}_{24}$, purity $\geq 99.0\%$), and sodium hydroxide (NaOH , purity $\geq 98.0\%$) were acquired from Sigma-Aldrich. Potassium iodide (KI , purity $\geq 99.0\%$) was acquired from Fisher Scientific Ltd. All solutions were prepared using distilled water from Milli-Q water (18.2 M Ω).

2.2 Sonoreactor configuration

The acoustic reaction system comprises a cooling-jacketed glass cylinder with an internal diameter 67 mm and an interchangeable plate transducer (Fig. 1). The glass reactor was custom-designed and made by the University of Southampton's Scientific Glassblowing Service. Cooling water was supplied *via* the reactor jacket by a 20 L Cole-Palmer PolyScience MX open



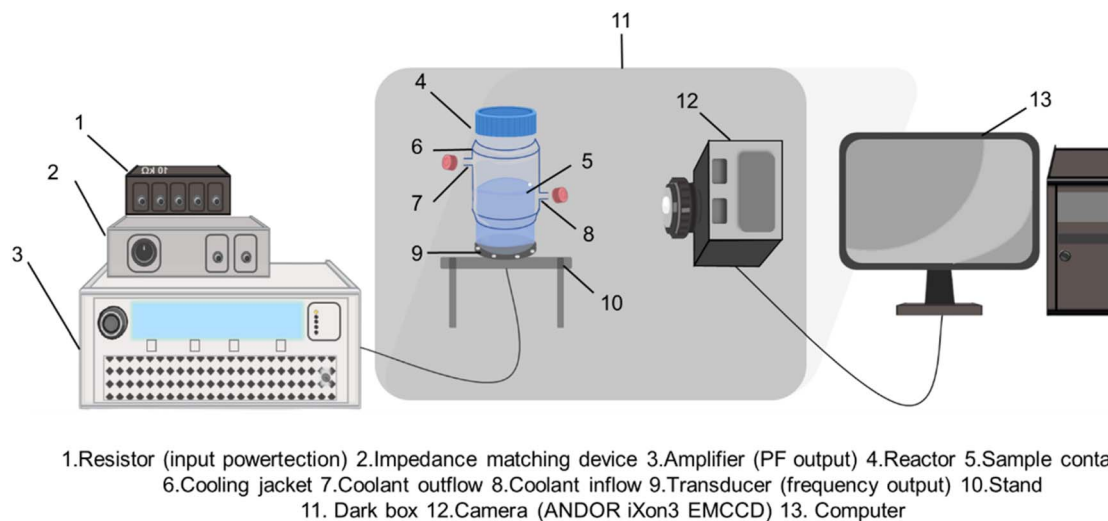


Fig. 1 Schematic diagram of the ultrasonic system and shooting system.

bath circulation system. All the plate transducers of a given frequency (22, 44, 98, 128, 200, 400, 500, 760, 1000, and 2000 kHz) were purchased from Honda electronics Co. Ltd. Each transducer comprised a 50 mm PZT piezo-electric ceramic, connected to a 100 mm diameter stainless steel vibration plate. The transducer was powered by the T&C Power Conversion AG 1006 amplifier. This amplifier model enables monitoring forward power, loading-power (10, 20, 30, and 40 W), and corresponding reflected power values. The amplifier operated in load power mode, automatically boosting power to offset any reflected power. The frequency was adjusted to minimise the resistance between the amplifier and transducer to decrease the reflected power values. A low-frequency RF impedance transformer from T&C Power Conversion functioned as a step-up transformer (SUT) and assisted in this process.

2.3 Experimental and analytical methods

2.3.1 Sono(chemi)luminescence intensity and image analysis. The spatial distribution and intensities of the SL and SCL generated during the ultrasonic irradiation were recorded for each experimental condition by ANDOR iXon3 EMCCD camera (Fig. 1). The images were taken entirely in a dark enclosure to prevent interference from ambient light. All ultrasonic parametric settings were carried out at three different tested solution volumes: 200, 300 and 400 ml. SL was conducted at ambient temperature using Milli-Q water as the cavitating medium,⁵⁴ with the camera's exposure duration set to 20 s and the electron multiplier (EM) gain level at 50. SCL was performed at room temperature using a 0.1 M NaOH solution with 1 mM luminol.^{30,38} The camera applied an exposure duration (s)/EM gain level of 4/4 for SCL since the considerably higher brightness generated by the luminol. The same brightness/contrast settings are applied to the images to make the comparison possible. SL/SCL images were taken after 30 s of sonication. Image-catching for every setting was repeated 3 times.

The camera associated ANDOR software (ANDOR SOLIS for Imaging X-7827) was used as per conventional methods to

perform pixel analysis on the entire captured photograph.^{4,29,55} In the subsequent section, this method will be referred as the 'entirety method' as its quantitative data is based on the whole picture. We introduced another image processing method for SL/SCL, based on a MATLAB program, to minimise subjectivity in noisy pixel data processing and eliminate the impact of reflections from glass instruments on the data (Fig. 2). The self-amending codes are provided in the ESI (Appendix A).† An analysis-threshold setting was adopted to better exclude irrelevant regions.⁵⁶ Luminous areas are introduced based on binary images (Fig. 2(c)).⁵⁷ The efficiency of the acoustic chemical activity in the reactor can be quantified to a certain degree through the grayscale value in the selected area (SA). The novel approach will be referred to as the 'selective method'. A comparison of entirety method and selective method was conducted.

The effective luminescence area proportion was calculated from the area data by the selective method, and this outcome used to indicate the collapse effectiveness of each reactor (eqn (1)).

$$P_{\text{eff}} = \frac{A_{\text{SA}}}{A_{\text{TR}}} \times 100\% \quad (1)$$

where, P_{eff} is the proportion of effective area in the total reactor cross-section. A_{SA} is total pixels in the brightness region selected by the area selection method and A_{TR} is the pixels for the total reactor cross-section (see Fig. 2 for examples).

The average luminescence intensity inside the active region was calculated *via* eqn (2).

$$I_{\text{SA}} = \frac{\text{GS}_{\text{SA}}}{A_{\text{SA}}} \quad (2)$$

where, I_{SA} is the average luminescence intensity in the effective reaction area and GS_{SA} is the total greyscales value in the effective area.

2.3.2 Iodide dosimetry and ROS measurements. The sonochemical oxidation was quantified by iodide dosimetry. IORS, such as $\cdot\text{OH}$, O radicals, superoxide anion ($\cdot\text{O}_2^-$) and



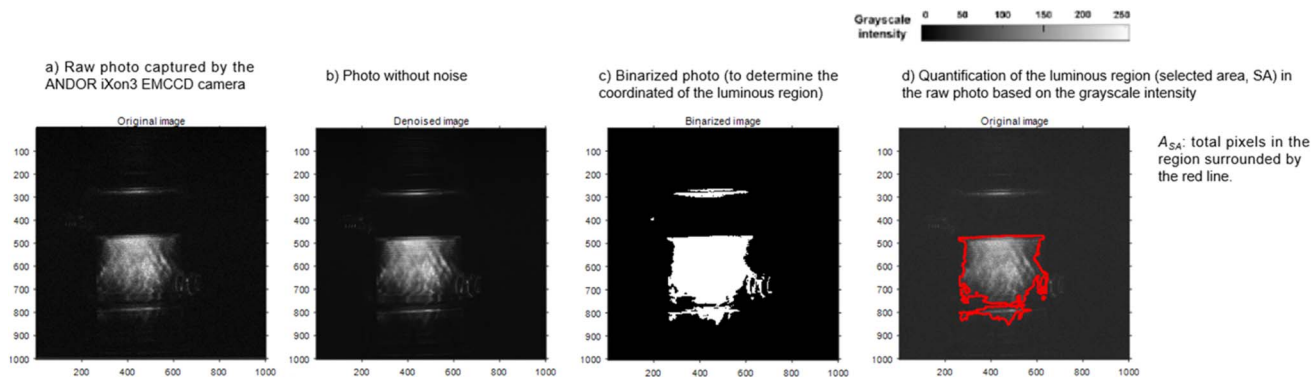


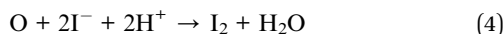
Fig. 2 Image processing based on (a) raw image, (b) photo without noise (denoised), (c) binarized photo, and (d) selected area in red.

hydroperoxyl radical ($\cdot\text{HO}_2$) and so on, and H_2O_2 are the primary ROS generated during sonication and play a key role in acoustic chemistry.

The oxidation of KI under ultrasound is generally considered to be the result of the following reaction with OH radicals:



However, it is worth noting that KI as a chemical probe is limited in its selectivity for $\cdot\text{OH}$. Several works pointed out that the primary oxidant in the sonochemical reaction may not always be the $\cdot\text{OH}$ radical, but sometimes the O radical,⁵⁸ and pointed out that the O radical may react in the KI oxidation reaction:^{59,60}



Additionally, when the subject is studied under an open system in an air environment,⁴ the limited selectivity results from the generation of nitrous acid from dissolved nitrogen during sonication, which oxidises iodide ions.^{61,62} Taking into account the potential of the ultrasonic oxidation and the experimental atmosphere, the determined yield of I_3^- was used to signify the overall IORS production, excluding H_2O_2 introduced to the KI solution by ultrasonic irradiation and without pointing solely to $\cdot\text{OH}$.

The generation rate of ROS during the ultrasonic process adheres to a zero-order reaction kinetics model, meaning the data point obtained after 10 minutes of irradiation is sufficient to determine the rate constant.^{29,60} Here, for the measurement of the relative concentration of IORS, 0.1 M KI solution was irradiated under the same conditions as for SL/SCL, and 2 ml samples were taken and tested by UV-vis (Thermo Scientific Evolution 201 UV-visible spectrometer) at 350 nm wavelength after each 10-min irradiation. IORS yield was calculated using the Beer–Lambert Law with a cuvette length of 1 cm and a molar absorptivity coefficient (ϵ) of $26\,303\text{ L mol}^{-1}\text{ cm}^{-1}$.¹⁷ Each experiment was repeated at least 3 times.

H_2O_2 produced during sonication reacts slowly with the iodide in the KI solution; thus a catalyst is required for reaction (eqn (5)) to occur^{28,63}



Hence, 0.5 mM ammonium molybdate was used as the catalyst in 0.1 M KI solution to account for the generation of total ROS (H_2O_2 and IORS). The measurement and calculation principles align with the pathway to obtain the IORS yields in pure KI under the acoustic process. The H_2O_2 yield could be derived by subtracting the IORS yield in pure KI from the total ROS.²⁸

2.3.3 Correlation analysis. The correlation coefficient is a statistical metric that quantifies the degree and direction of the linear association between two variables.⁶⁴ The most often used correlation coefficients are the Pearson correlation coefficient and the Spearman's rank correlation coefficient.

The Pearson correlation coefficient, commonly represented as r , is a statistical metric that quantifies the magnitude and direction of a linear association between two variables (eqn (6)).⁶⁵

$$r = \frac{\sum (x_i - \bar{x})(y_i - \bar{y})}{\sqrt{\sum (x_i - \bar{x})^2 \sum (y_i - \bar{y})^2}} \quad (6)$$

where, x_i, y_i = the i -th observation. \bar{x}, \bar{y} = the average value of variable x and variable y .

The variable ' r ' ranges from -1 to 1 . A value of $r = 1$ implies a linear increase that follows a strong positive correlation. A value of $r = -1$ shows a strong negative correlation. A value of $r = 0$ shows the absence of any linear correlation. The Pearson correlation coefficient is appropriate for quantifying linear associations.

Considering, the intricate nonlinear characteristics of the ultrasound process may result in complex nonlinear interactions among numerous acoustic measurements, rather than straightforward linear ones.^{66,67} Spearman's rank correlation coefficient, symbolised as ρ , not supposing a linear relationship between variables⁶⁸ may be more appropriate for elucidating the correlations among SL, SCL, and ROS, as well as their responses to variations in acoustic circumstances. The derived monotonicity results may elucidate complex connections in sonochemical processes and offer direction for experimental improvement. The calculation of this metric relies on the



ranking of variables rather than the raw data, making it suited for non-linear yet monotonic connections (eqn (7)).

$$\rho = 1 - \frac{6 \sum d_i^2}{n(n^2 - 1)} \quad (7)$$

where, d_i = the disparity in the rankings of the two variables for the i -th observations. n = the number of observation data.

The ρ ranges from -1 to 1 . A value of $\rho = 1$ implies a complete positive correlation, meaning that the variables have identical ranks. A value of $\rho = -1$ implies an exact negative correlation, meaning that the variables have precisely opposite rankings. A value of $\rho = 0$ implies the absence of a monotonic relationship between the two variations. Spearman's rank correlation coefficient is computed from the ranks of the observations and is unaffected by the data's distribution. It is less affected by extreme values and is appropriate for non-normally distributed data.

3. Results and discussion

3.1 Validation of the novel image processing method (selective method)

The intensity of SL emission is related to the highest temperature achieved by a collapsing cavitation bubble, and associated luminescent species.⁶⁹ The intensity of SCL emission is caused by the reaction between luminol molecules and superoxide anion radicals (O_2^-) or $\cdot\text{OH}$.^{4,70} Bubble characteristics are often assessed by measuring the emission intensity of SL and SCL.^{28,71,72}

Initially, the ANDOR camera and its accompanying software, ANDOR SOLIS for Imaging X-7827 was used for all processing. The software was configured to a scale mode ranging from 293 to 3159 to accurately reflect all pixel values in the photo. The image size, measured in pixels, is 1005 pixels wide and 1002 pixels high. No further adjustments were made. The SL and SCL photographs' total counts obtained from ANDOR system was exhibited under all testing ultrasonic input parameters (Fig. 3). The data collection indicates higher values at ultrasound frequencies below 200 kHz (excluding 200 kHz).

Fig. 4 shows the qualitative comparison of SL and SCL intensity and spatial distribution at 300 ml reactor volume (images for the 200 and 400 ml are presented in Fig. S1 and S2,[†] respectively). The systematic photos clearly illustrate that both SL and SCL demonstrate greater intensity at ultrasonic frequencies ranging from 200 to 760 kHz. This outcome is incongruous with the direct digitalization of images given *via* the ANDOR software. The brightness digitization process is distorted by the noise signals found in the image with the lower real brightness, leading to misleading results.

Four main factors cause noise in photographs captured under low-light conditions, commonly seen in normal photography.^{73,74} (1) The camera's increasing photosensitivity (ISO). To capture more accurate data graphs and more detailed information, the ISO of the camera is typically increased. However, this also amplifies noisy signals while enhancing small signals. (2) The signal-to-noise ratio (SNR) diminishes. When there is less light, the intensity of the image weakens, causing an increase in the visibility of random noise. (3) The emergence of thermal noise. Thermal noise occurs

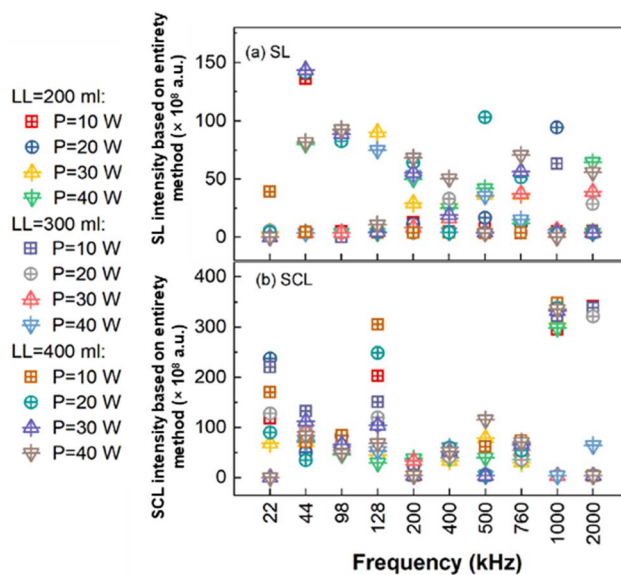


Fig. 3 The luminescence (a) SL and (b) SCL intensities measured from complete photo (entirety method).

when the camera sensor's increased temperature causes electrons to move randomly and more vigorously, especially during longer exposure durations or while employing high ISO settings. (4) Amplification effects intensify the noise in image regions that have a weak signal, such as dark areas and shadows.

The camera's sensitivity remains the same during the shooting process. The ANDOR camera, as a high-performance device, is equipped with an advanced cooling system that efficiently controls the sensor's temperature, thus reducing the generation of thermal noise. Accordingly, the main reason for deceiving noise data in SL and SCL images is the poor signal-to-noise ratio and the amplification effect of low light emission.

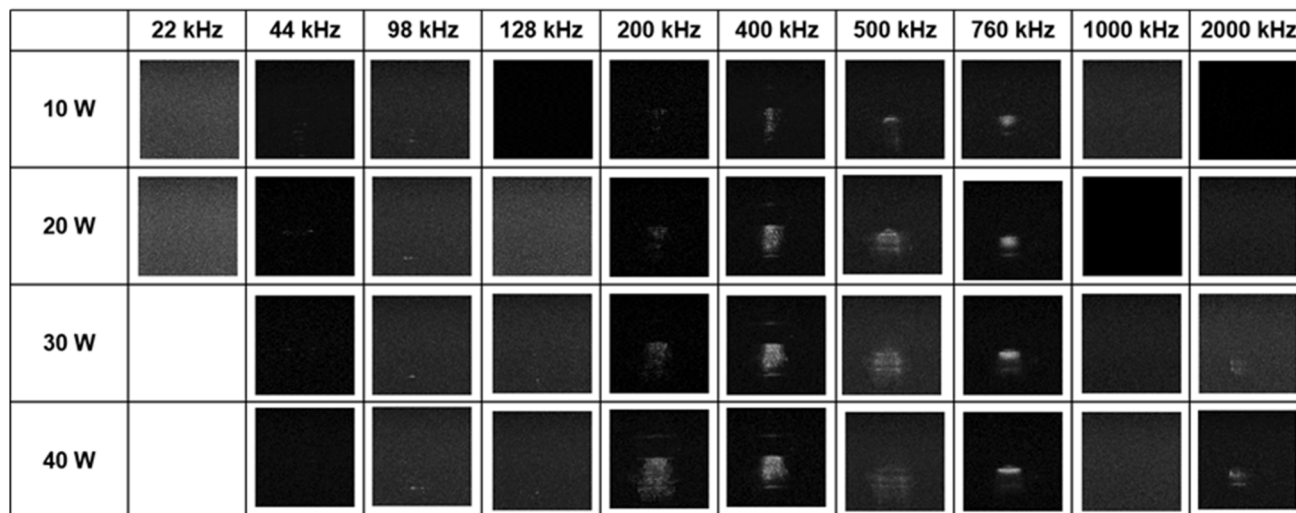
We sought to introduce a methodology that could enhance the accuracy of quantitative results through noise reduction. Based on the reference method,^{56,57} we define the grayscale value as 0 to 255 to reflect brightness differences. Furthermore, this method starts with intelligent pre-noise reduction of the complete image through MATLAB built-in programme statements (Fig. 2(b)).⁷⁵ Then the appropriate range of luminance in the image is selected based on the binarised image (Fig. 2(c and d)). The specified area captured the grayscale values, theoretically improving the object's quantitative precision.

The images' pixel values obtained from the ANDOR approach and the selective method were compared for all testing conditions in SL and SCL (Fig. 5). In contrast to the data collected through the original methodology, the region selection method's findings show greater digital values above 200 kHz, indicating a higher intensity of both SL and SCL at frequencies under these sonic conditions. This matches the findings of visuals taken within an enclosed dark box (Fig. 4, S1 and S2[†]). Empirical evidence suggests that selective methods are more reliable for quantitative studies across a broad range of parameters.

To evaluate the suitability of the selective method, a subset of SCL images were selected to extract digital values and conduct a more detailed analysis (loading-power 40 W, volume 400 ml,



(a) 300 ml - SL



(b) 300 ml - SCL

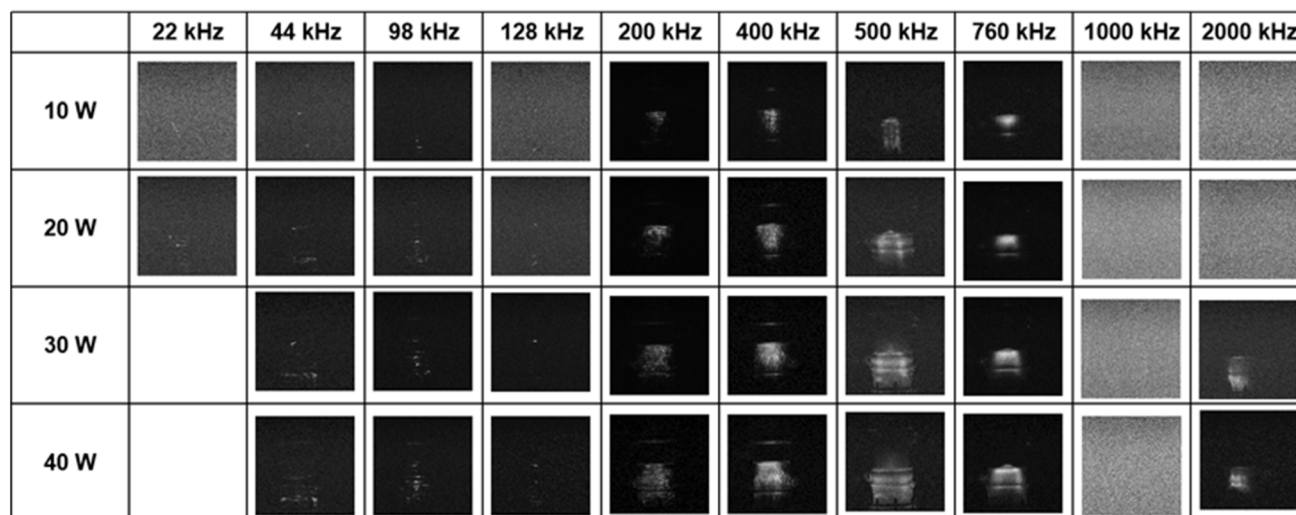


Fig. 4 Sono-/sonochemi-luminescence intensity distribution for all testing frequencies and powers at 300 ml reactor volume: (a) SL and (b) SCL.

Fig. 6 and 7). The displayed images are subjected to colour inversion to enhance visibility, where greater pixel values correlate to darker regions of the image (Fig. 7). This pathway could also potentially enhance our perception of the original image's signal to noise ratio. The SCL emission intensities at 40 W and 300 ml showed notable differences at low (44, 98, and 128 kHz), medium (200, 400, 500, and 760 kHz), and high (1000 and 2000 kHz) frequencies using separate approaches. According to the findings provided by the entirety method, the intensity of SCL emission at low frequencies seems comparable to that at medium frequencies (Fig. 7). Nevertheless, the findings obtained by applying the selective method better match the actual scenario (Fig. 6(a–g) and 7 (red)). The subset of SCL images, consisting of colour-reversal format photographs, illustrates that the light emission images at five distinct frequencies (44, 98, 128, 500, and 1000 kHz) exhibit elevated

levels of greyscale in the empty region (Fig. 6). The presence of disruptive signals in the silent sections led to errors in the quantification process of the data from the whole photo, both at low and high frequencies (Fig. 7 (blue)). The distortion was especially noticeable at 1000 kHz (Fig. 6(h) and 7). Thus, it is reasonable to conclude that the results obtained through the selective method demonstrate acceptable reliability.

When high-quality luminescence photographs can be acquired, typically at mid-frequency ($200 \text{ kHz} \leq f \leq 760 \text{ kHz}$), the quantitative data collected through both procedures demonstrated an identical variation tendency (Fig. 5 and 7). The selective method can enhance accurate data acquisition in a broader variety of experimental settings by circumventing misleading signals. Thus, for the following study of emission intensity for SL and SCL, quantitative analyses uses image digitation through the selective method.



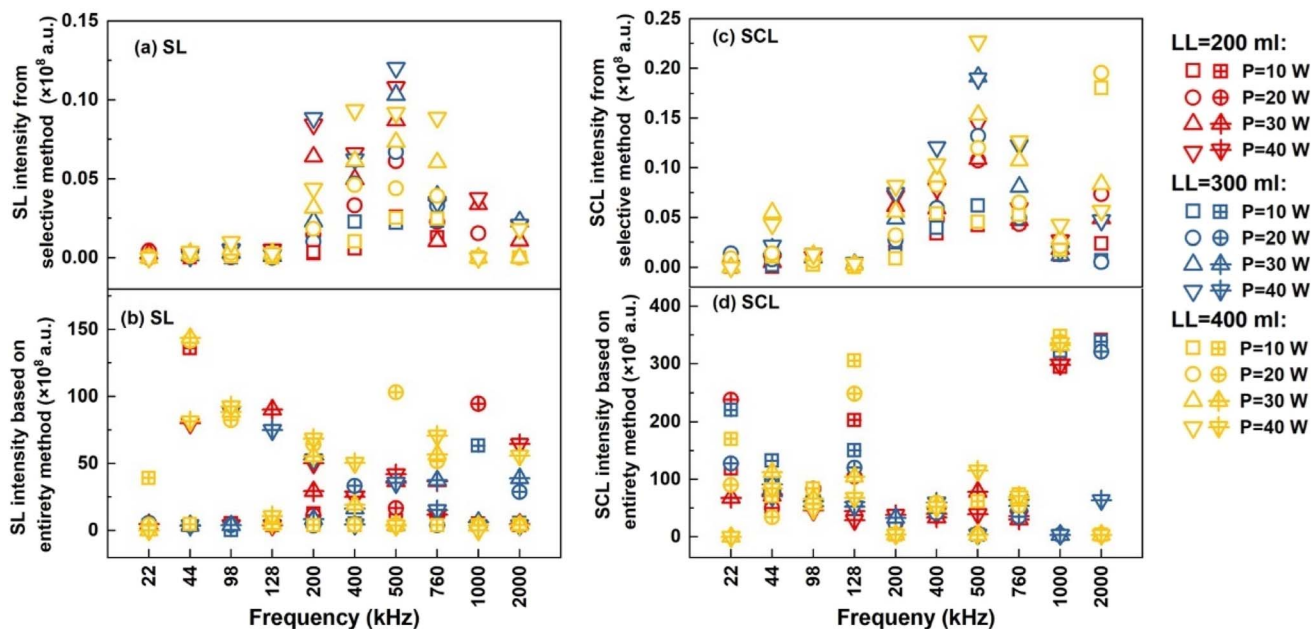


Fig. 5 Image digitalization date of sono(chemical) luminescence intensity from the selective method ((a) SL and (c) SCL) and entire method ((b) SL and (d) SCL).

3.2 SCL, SL and ROS measurements

The subsequent section presents the findings of the SCL/SL activity and KI dosimetry. All the powers mentioned in the results are loading-powers (P_L). The loading-power density (PD_L) were calculated through eqn (8)

$$PD_L = \frac{P_L}{V} \quad (8)$$

where, V is the reactor volume (L). The values of PD_L can be found in the appendix (Table S1[†]), and the calorimetric powers tested results can be found in the appendix (Table S2[†]).

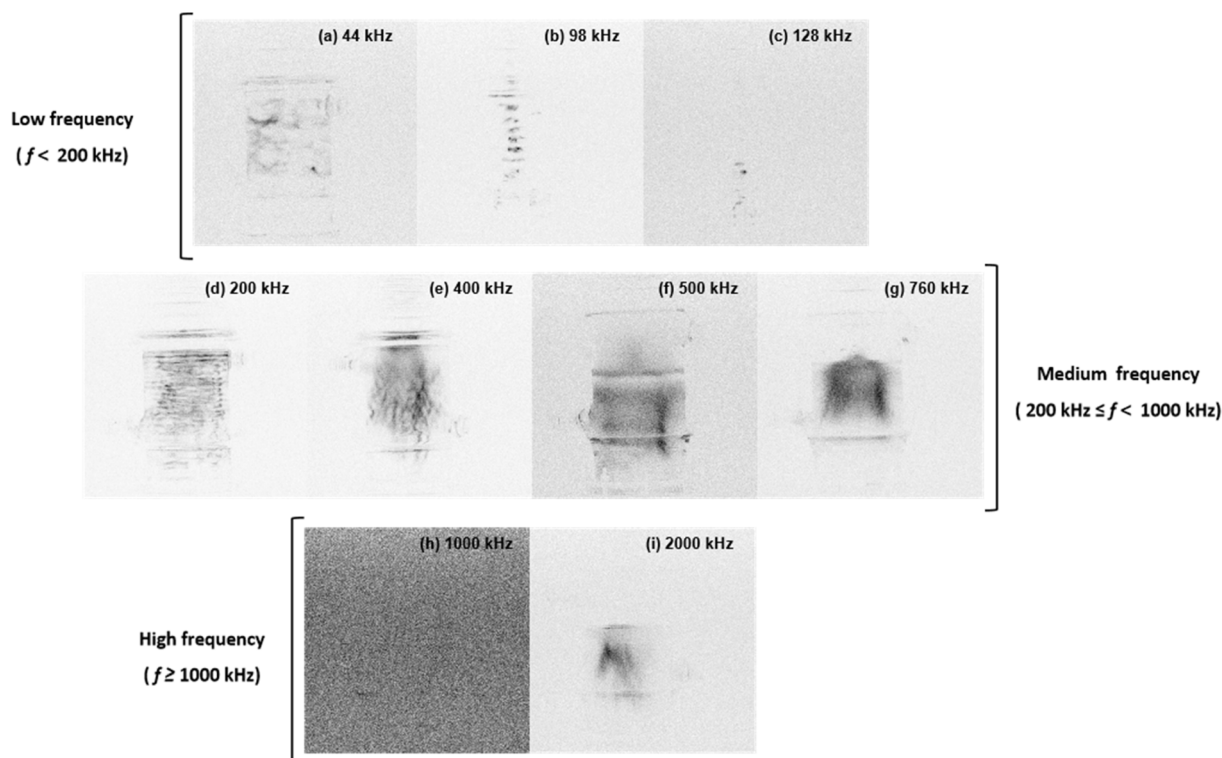


Fig. 6 SCL emission at 40 W loading-power, 400 ml reactor volume, and frequencies with (a–c) $f < 200$ kHz, (d–g) $200 \text{ kHz} \leq f < 1000$ kHz, and (h and i) $f \geq 1000$ kHz. Image colour-reversal.



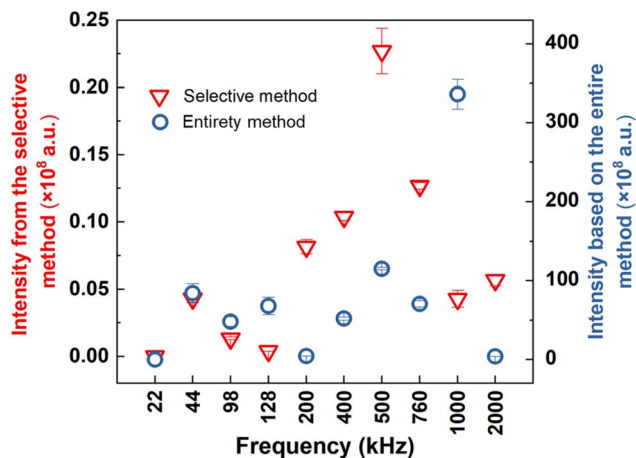


Fig. 7 Extracted digital value from the selective (red) and entirety (blue) methods for SCL intensity under 40 W and 400 ml.

3.2.1 Sono(chemi)luminescence intensity measurement.

The overall brightness emission intensities of SL and SCL were evaluated over 114 operating conditions. The combined effect of

frequency, power and volume on the sonochemical activity with SL/SCL intensities are demonstrated by bubble-modified heatmap (Fig. 8 and 9). The modified heatmap circle size represents the proportion of the effective luminous area, while the average luminous intensity is represented by the hue of the circles. The findings of the ultrasound luminous reactions show greater sono(chemi)luminescence at frequencies ranging from 200 to 760 kHz, namely at middling-setting sonic frequencies, compared to other frequency settings (Fig. 8 and 9). This signifies that a greater number of bubbles experience stable and transient cavitation within this range.

Frequencies 400 and 500 kHz seem to be the most effective for SL. These frequencies of irradiation not only generate the most intense luminescence but also encompass a broad, effective area of sonoluminescence. This phenomenon could be attributed to the resonance conditions of the bubbles being well-matched to the properties of the acoustic waves at these frequencies.⁷⁶ As a result, there is an efficient conversion of energy and a broad variety of cavitation effects. The increase in power density results in an increasing influx of energy into the bubble core, which benefits SL activities. The positive impact of

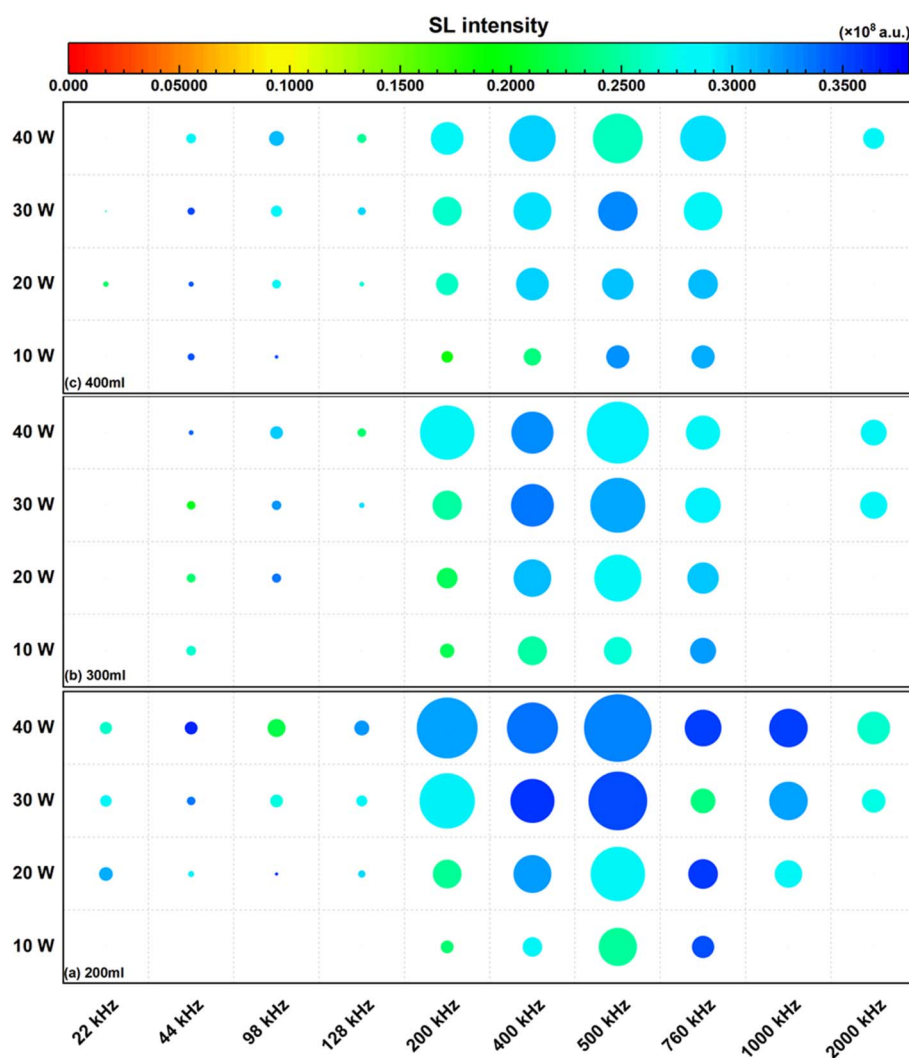


Fig. 8 Relationship between SL intensity with frequencies, power and reactor volume ((a) 200 ml, (b) 300 ml and (c) 400 ml).



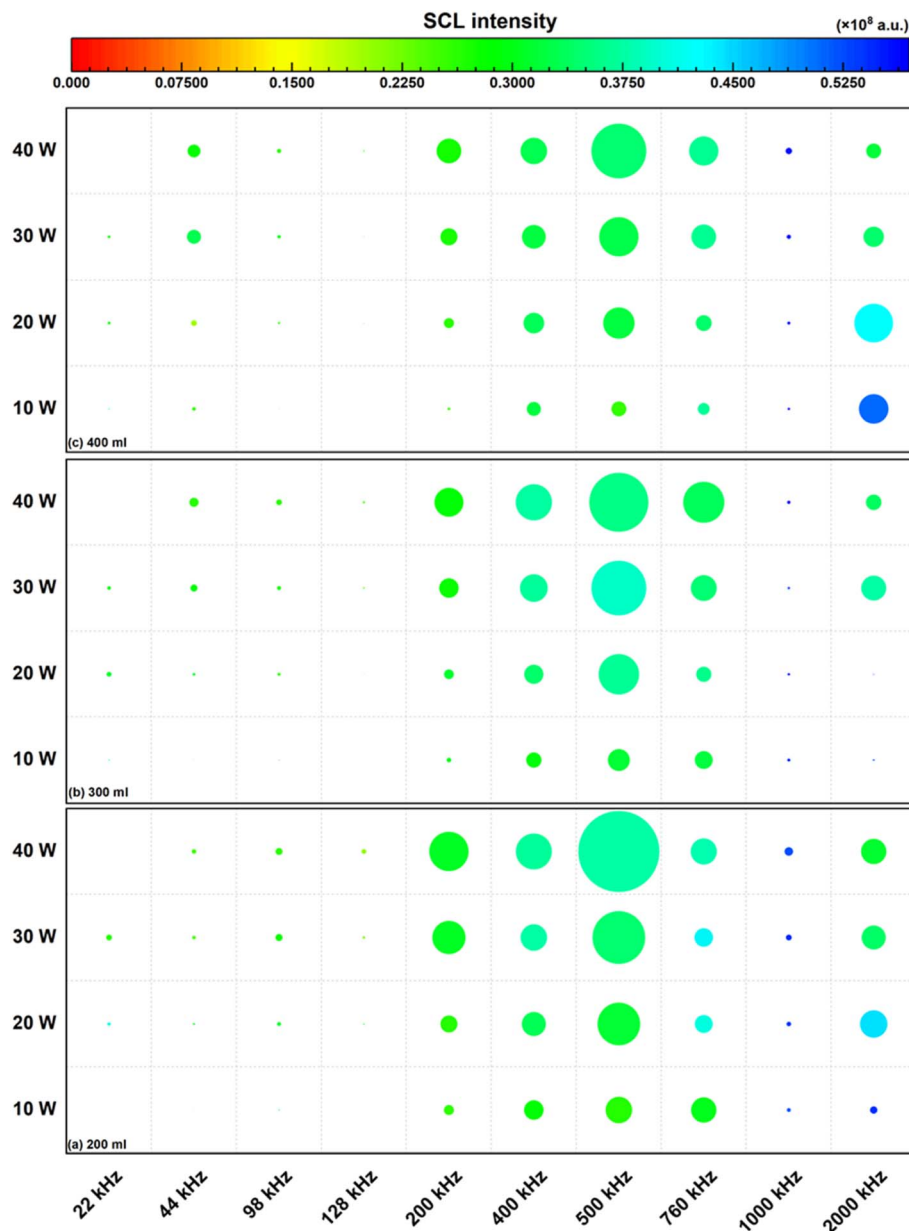


Fig. 9 Relationship between SCL intensity with frequencies, power and reactor volume ((a) 200 ml, (b) 300 ml and (c) 400 ml).

power on overall SL intensity becomes more significant at frequencies of 200 kHz compared to lower frequencies. One reason is that as the irradiation frequency increases, the number of bubbles in the reactor also increases. Additionally, the boundary or attenuation effects in cavitation caused by excessive energy input should not be ignored.¹ The adverse impact of extra high-power loads is demonstrated by the average luminous intensity or effective luminous area of the SL at lower frequencies. In a 44 kHz system, an increase in power will invariably result in a drop of at least one unit in the effective luminous area or average brightness of the SL. In a 98 kHz system, the average emission intensity of the SL consistently diminishes with increasing power. The data for effective luminous area are presented in the appendix (Tables S3 and S4†).

The frequency of 500 kHz is the optimal choice for SCL in these conditions, since it consistently demonstrates a significant intensity and luminescent area across various liquid heights and power settings (Fig. 9). Under conditions of high-power density, frequencies of 400 and 760 kHz also demonstrate favourable outcomes. The SCL frequency that have high brightness and active area are in the medium frequency (200 kHz $\leq f \leq$ 760 kHz) range, gaining similar outcomes to SL. Augmenting power can greatly amplify the intensity of SCL and expand the area of luminescence. The increasing of SCL overall intensity with increasing power is typical.⁷⁷ No attenuation from the extra power input, distinguish to those shown in the SL results, were detected at frequencies ranging from 22 to 128 kHz. In general, increasing the volume can result in a wider area



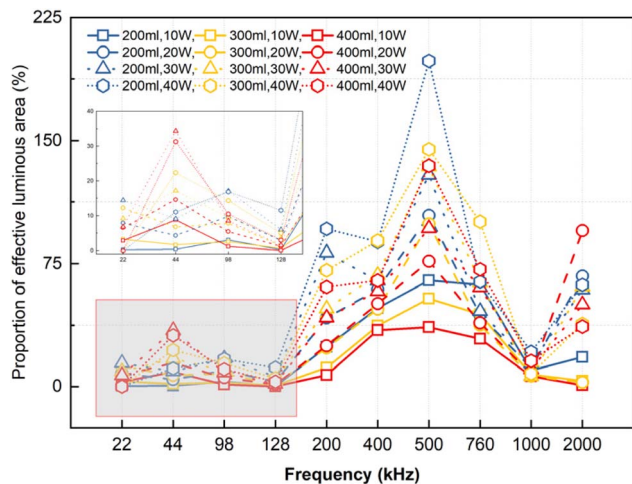


Fig. 10 The variation of effective SCL area with frequency, power, and volume.

of luminescence and enhance the overall SCL intensity (Fig. 5(c)). Nevertheless, as the volumes increase, the proportion of the SCL area to the entire reactor decreases (Fig. 10). Throughout the entire test frequency range, as the liquid level height increases, the SCL region consistently remains close to

the liquid surface, exhibiting elongation and narrowing in the vertical direction. A representative set of SCL images is shown in Fig. 11. As the solution volume increases, the SCL zone moves farther away from the bottom of the reactor, resulting in a decrease in the proportion of the luminescence region in the total reactor. The same regular situation was observed in the tests of SCL under other conditions (Fig. 4(b), S1(b) and S2(b)†).

There are a few noteworthy findings from SCL results. The effective area of SCL at 44 kHz increases with increasing liquid height (Fig. 10 and 11(a–c)). This differs from the results at other frequencies. At liquid levels of 200 and 300 ml in the reaction system, the SCL exhibited a nearly complete distribution of effective luminescence across the reactor at a frequency of 500 kHz (Fig. 11(d and e)). In the travelling wave field, the resonant bubbles with SCL characteristics were directed towards the liquid surface by radiation force.⁶⁶ The radiation force also induces deformation at the liquid–air contact. At low intensity, the deformation resembles a concentric annulus (Fig. 11(a–c) and (g–i)); at high intensity, an auditory fountain manifests as droplets erupt forcefully (Fig. 11(d–f)).^{78,79} Consequently, under specific working conditions, the effective area of the SCL exceeds 100% (Fig. 10). These findings suggest that each frequency of ultrasound will have an associated ideal solution height to maximise efficiency and provide a more uniform sonication process.

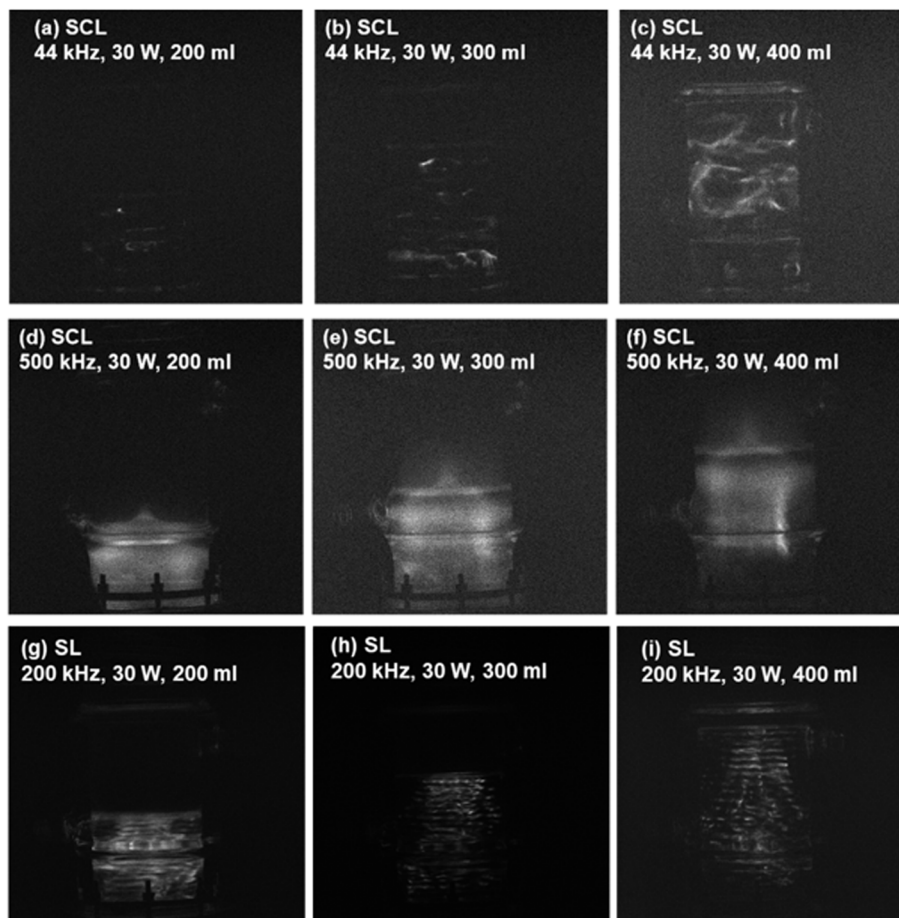


Fig. 11 SCL distribution in different solution volumes under 44 kHz (a–c), 500 kHz (d–f) and 200 kHz (g–i).



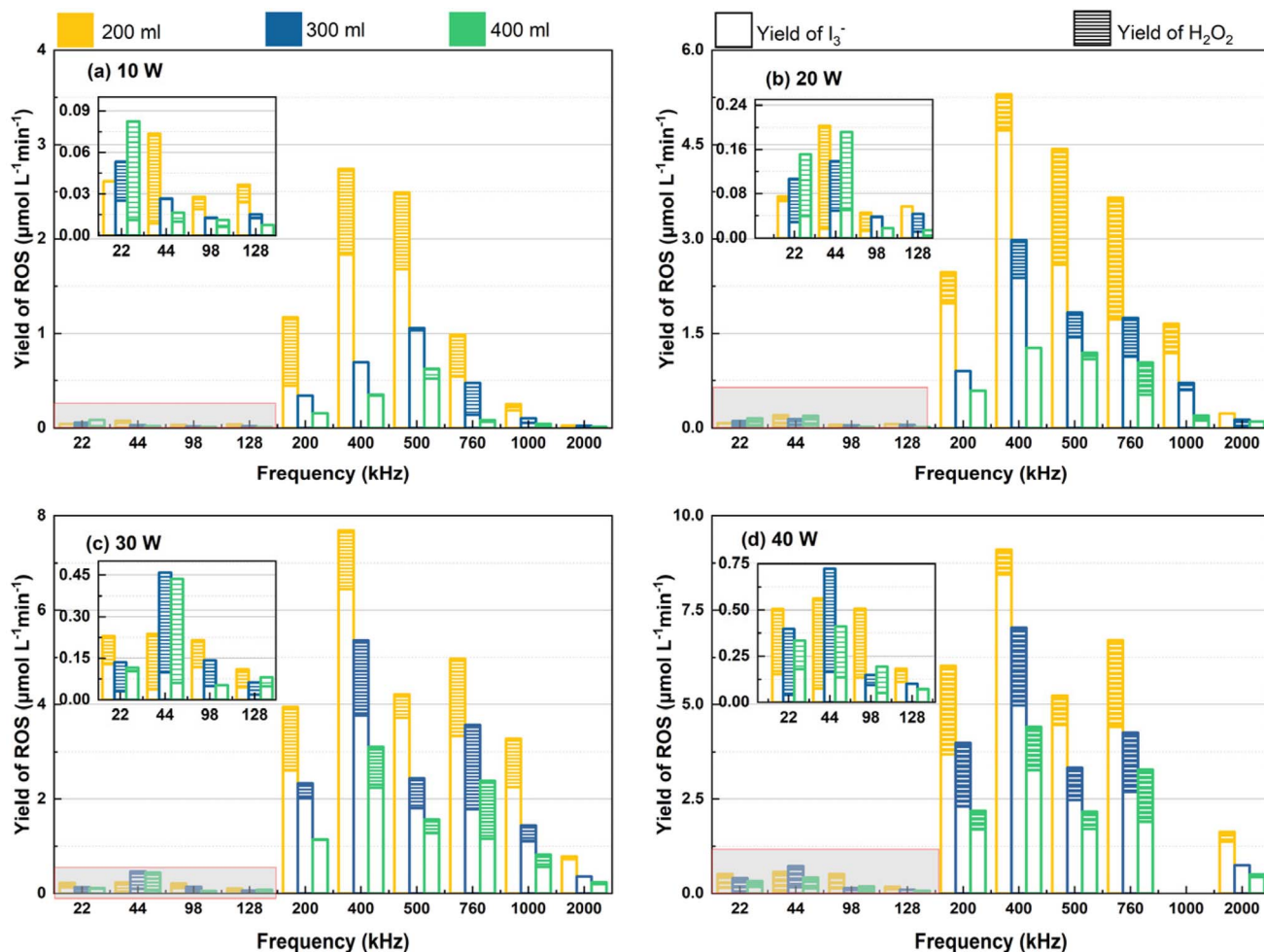


Fig. 12 The dosimetry results for quantification of IORS, H_2O_2 and total ROS with different frequencies and solution volume: (a) 10 W, (b) 20 W, (c) 30 W and (d) 40 W.

Based on the findings from both SL and SCL, alterations in the luminescent area had a more significant influence on the total sono(chemi)luminescent activity and efficiency levels compared to changes in the average luminescence intensity (Fig. 5(a and c), and Tables S3 and S5[†]). According to these results, we classify the area of sono-active bubble behaviour into three separate ranges of ultrasonic frequencies: $f < 200$ kHz, $200 \text{ kHz} \leq f \leq 760$ kHz, and $1000 \text{ kHz} \leq f \leq 2000$ kHz, in this reactor set-up.

3.2.2 KI dosimetry. The yield of I_3^- was calculated and used to reflect the yield of IORS during ultrasonic irradiation due to the limited $\cdot\text{OH}$ selectivity of KI. The total oxidation behaviour was characterized through the total and individual yield of IORS and H_2O_2 (Fig. 12). Recent studies have shown that reducing agents (such as H^+ , H_2 , and HO_2) may also spontaneously form during sonication and influence the overall reaction system.⁸⁰ Given that reducing agents may compete with I^- for ROS, the I_3^- yield measured in KI dosimetry reflects a net oxidation outcome rather than an absolute measure of IORS or ROS concentration.

The overall production of ROS is greater at 400 and 500 kHz, whatever the operating conditions. Nevertheless, the application of greater amplitude (power ≥ 20 W) at a frequency of 760

kHz significantly enhances the reaction system's oxidising capability. The oxidant ability almost aligns with the three distinct frequency intervals discussed in sono(chemi)luminescence. The systems with medium frequency ($200 \text{ kHz} \leq f \leq 760$ kHz) ultrasound can generate higher concentrations of ROS than low and high-frequency irradiated systems.

At low frequencies ($22 \text{ kHz} \leq f \leq 128$ kHz), the primary ROS generated are H_2O_2 (Fig. 12). However, as the system frequency reaches 200 kHz, the proportion of IORS in the overall ROS rises. The ROS (mainly $\cdot\text{OH}$) are dispersed through two pathways into the liquid. The majority of ROS are released into the medium through the collapsing bubble, while some diffuses into the solution through the bubble wall.²⁵ Bubbles generated with low-frequency irradiation have larger critical dimensions due to extended expansion durations and compared to those formed at higher frequencies.⁸¹ This phenomenon enables bubbles formed at low frequencies to reach higher temperatures and to vaporise a greater number of water molecules before collapsing.^{82,83} The $\cdot\text{OH}$ that is confined within the bubble for a longer duration experiences a recombination reaction, leading to a greater H_2O_2 proportion of a lower total ROS in liquids exposed to low-frequency ultrasound (Fig. 13).



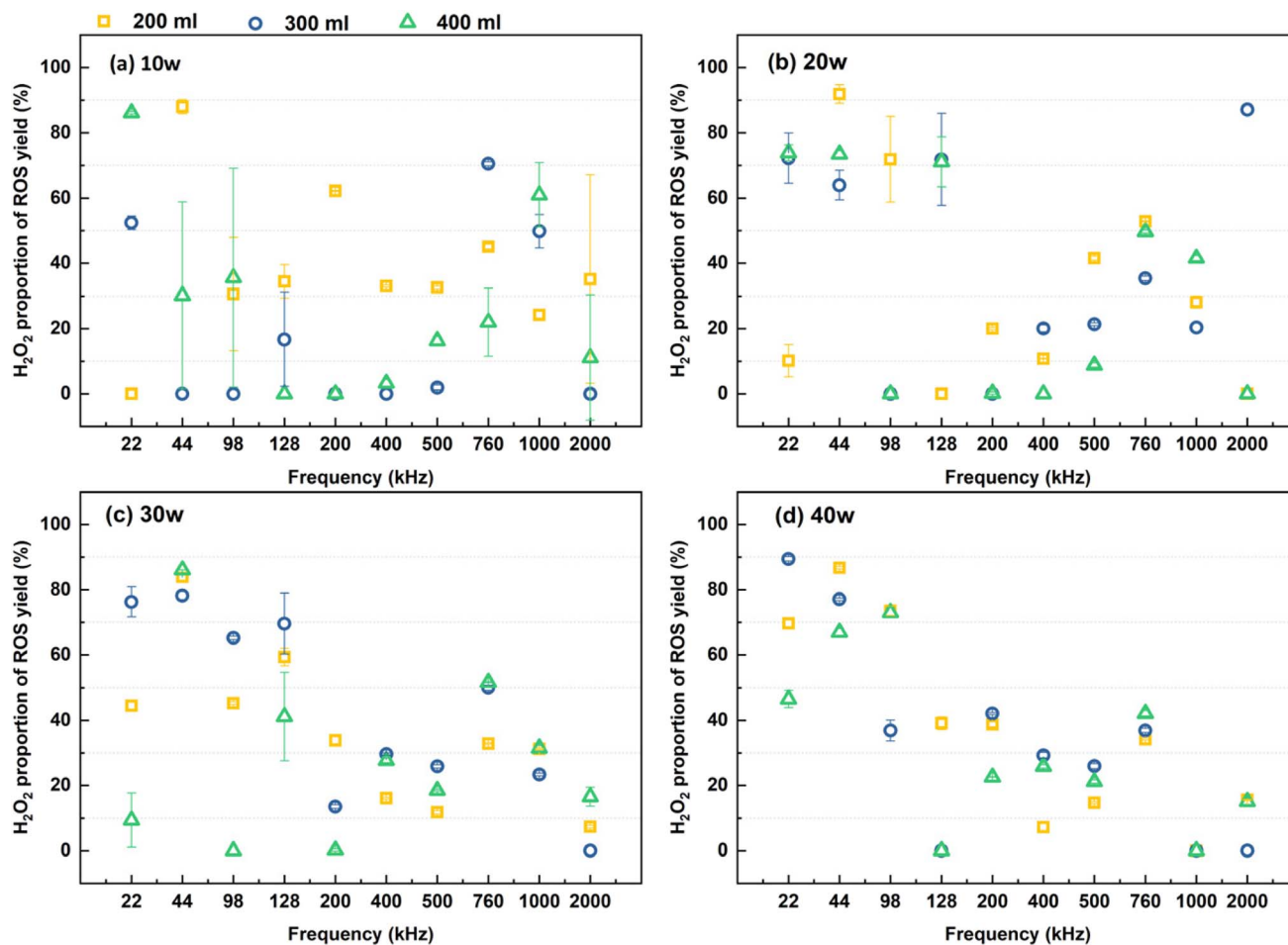


Fig. 13 H_2O_2 proportion of total ROS yield with different frequencies and solution volume: (a) 10 W, (b) 20 W, (c) 30 W and (d) 40 W.

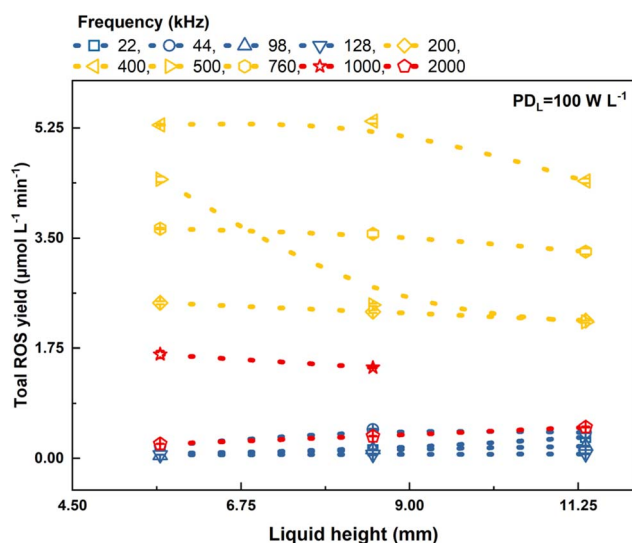


Fig. 14 Correlation between liquid height and ROS yield at different frequencies with same power density (100 W L^{-1}).

The system's oxidative characteristics are significantly enhanced by increasing power, and this improvement is primarily shown in the total ROS generation (Fig. 12). The

liquid's level affects the energy distribution and propagation path of sound waves within the ultrasonic system using the plate transducer. The correlation between ROS yield and variations in liquid height at a power density of 100 W L^{-1} is shown in Fig. 14. As the liquid height increases, the yield of ROS exhibits a marginal decline in the medium frequency (200 to 760 kHz) and a rise in the low-frequency range (22 to 128 kHz). This alteration trend is associated with the frequency-dependent cavitation threshold.⁸⁴ In contrast to the influence of volume on power density or energy distribution, the impact of liquid height on energy transfer is comparatively less. Within a similar reactor, the restricted influence of height on sonochemical activity was previously observed.⁵

3.3. Correlation between sonochemical/luminescent activities

Visualisation of the distribution of activity, and active bubbles are achieved through sono(chemi)luminescence, and chemical processes measured *via* SCL and ROS. Through a comprehensive analysis of the impact of ultrasonic frequency, power, and liquid height on sono(chemi)luminescent activity and ROS yield, we have determined that adjusting these acoustic parameters is essential in building an efficient acoustic system.



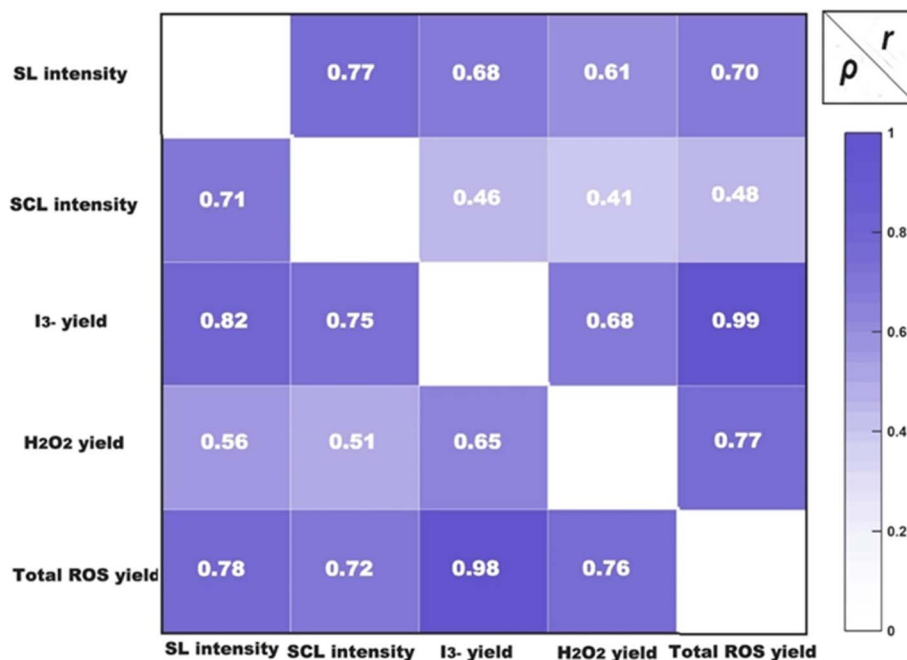
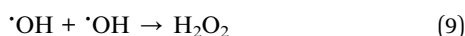


Fig. 15 The correlation coefficients between I₃⁻ yield, H₂O₂ yield, total ROS yield, SL intensity and SCL intensity.

The activity of SL and SCL and the production of ROS of the sonicated reaction system were increased in the middle-frequency range, particularly at 400 and 500 kHz. The cavitation effect was optimised, resulting in a substantial increase in the chemical activity of the reaction system, particularly when operating at high power and with low liquid levels. To better understand these phenomena, an analysis will be conducted on the interrelationships among these sonochemical features.

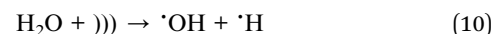
3.3.1 Correlation between ROS (IORS and H₂O₂) yield with SL/SCL intensity. The potential relationships between ROS (IORS and H₂O₂) yield with SL and SCL were examined by calculating the correlation coefficients (Pearson product-moment correlation coefficients (r) and Spearman's rank correlation (ρ)) (Fig. 15). The yield of IORS was reflected by the yield of I₃⁻.

The correlation coefficient evaluation revealed an extremely strong positive association between the yield of I₃⁻ and the total generation of ROS, *i.e.* $\rho_{\text{total ROS yield vs. I}_3^- \text{ yield}} = 0.98$, and $r_{\text{total ROS yield vs. I}_3^- \text{ yield}} = 0.99$. This indicates that most of the ROS generated in the reaction cell from ultrasonic irradiation could be measured using potassium iodide dosimetry, even without the use of a catalyst. Hydrogen peroxide (H₂O₂) is a significant by-product of the secondary reaction involving the $\cdot\text{OH}$ follow the reaction as



The amount of H₂O₂ generated depends on the amount of $\cdot\text{OH}$ released and the efficiency of the subsequent recombination reaction.⁸⁵ Typically, the powerful oxidising ability of the hydroxyl radical causes its yield to substantially impact the efficiency of chemical reaction, including but not limited to

pollutant degradation/water disinfection achieved *via* ultrasound treatments.^{86–88} In water mediated reaction, the bulk of IORS are mainly explained by the hydroxyl radical yield, which is based on the theory of molecular sonolysis, denoted ' $\cdot\text{OH}$ ', as^{28,89}



The positive correlation between changes in I₃⁻ yield and changes in SL intensity was stronger than that of SCL, but the change was not strictly monotonically linear, *i.e.* $\rho_{\text{total ROS yield vs. SL intensity}} = 0.78 > \rho_{\text{total ROS yield vs. SCL intensity}} = 0.72$, and $r_{\text{total ROS yield vs. SL intensity}} = 0.70 > r_{\text{total ROS yield vs. SCL intensity}} = 0.48$. These results explain why some studies acknowledge a strong link between ROS yield and SL intensity,⁵⁴ whereas other investigations provide results that do not align with this observed trend.^{4,28} Different test parameters always produce different findings, which is why this inherent correlation is always predicated on those differences. Based on the systematic test results of the present investigation, it can be concluded that there is a positive relationship between ROS yield and SL in most circumstances. Nevertheless, it is important to thoroughly investigate the presence of exceptional circumstances.

3.3.2 Disparities between ROS yield with SCL intensity. While SCL and KI dosimetry are both sonochemical processes that can occur at lower energy (than SL), their relationship is not linear nor strong monotonic (Fig. 15 and 16). The coefficient results, $r_{\text{I}_3^- \text{ yield vs. SCL intensity}} = 0.46$, $r_{\text{H}_2\text{O}_2 \text{ yield vs. SCL intensity}} = 0.41$ and $\rho_{\text{H}_2\text{O}_2 \text{ yield vs. SCL intensity}} = 0.51$, reflect this disparity.

The disparities could be attributed to the mechanism of sonochemistry, which is directly influenced by the rate at which the active species is transferred to the solution.⁹⁰ The abrupt implosion of cavitation bubbles undergoes non-linear growth



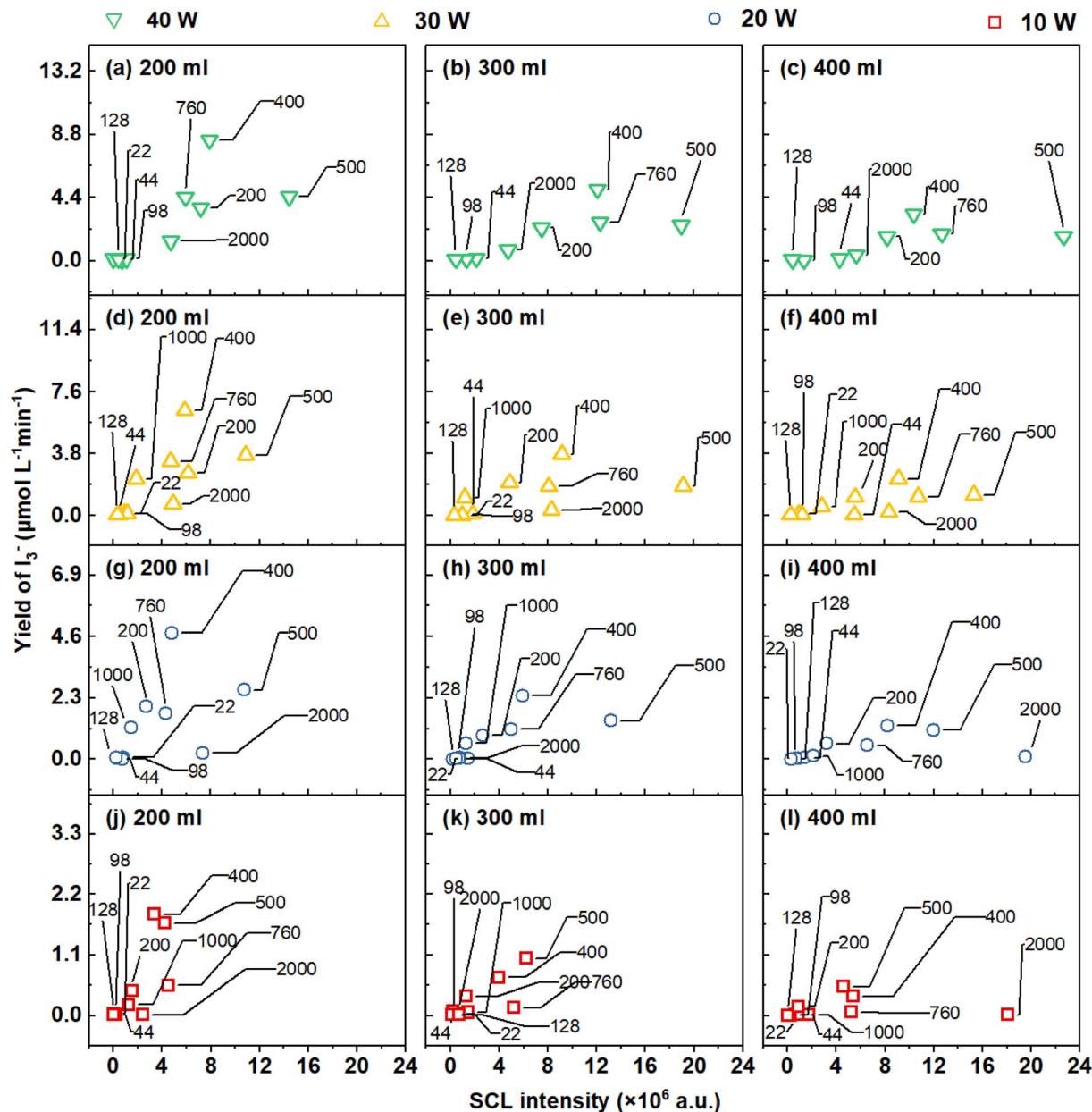


Fig. 16 The correlation between I_3^- and SCL generated during sonication (testing frequencies are labelled (kHz)): (a–c) 40 W, (d–f) 30 W, (g–i) 20 W and (j–l) 10 W.

during the ultrasonic process, which can result in extreme scenarios where heat and mass transmission deviate from the linear model.^{54,90} Additionally, the luminescence mechanism of SCL is formed by a chemical reaction between luminal molecules and $\cdot O_2^-$ and $\cdot OH$ radicals produced within the cavitation bubble.⁷⁰ The luminol molecule comprises polar and nonpolar elements. Polar elements, such as amino and carbonyl groups, render the molecule hydrophilic, whereas nonpolar components, like the benzene ring, tend to avoid polar water molecules and prefer interaction with nonpolar atmospheres. At the gas–liquid interface, the hydrophobic segment of the luminol molecule may orient toward the gas phase. This amphiphilic molecule may autonomously move to the gas–liquid boundary, reducing the exposure of the hydrophobic portion to the liquid

while maintaining the interaction of the hydrophilic portion with the liquid phase.⁹¹ Luminol's activities follow amphiphilic molecular characteristics, indicating an affinity for the bubble's surface.^{92,93} Therefore, we theorise that the sonochemical activity triggered by SCL is primarily concentrated in the vicinity of the bubble surface.⁹⁴ The complex variations in acoustic pressure cause intricate non-linear characteristics in the waveform and affect the dynamics of the gas–liquid interface through the absorption and release of energy by cavitation bubbles.^{95,96} However, KI dosimetry measures the response of ROS (mainly $\cdot OH$ and H_2O_2) by assessing the formation of iodine complexes in the entire solution. Dosimetry results demonstrate the oxidation that sonication imparts into the entire reaction.



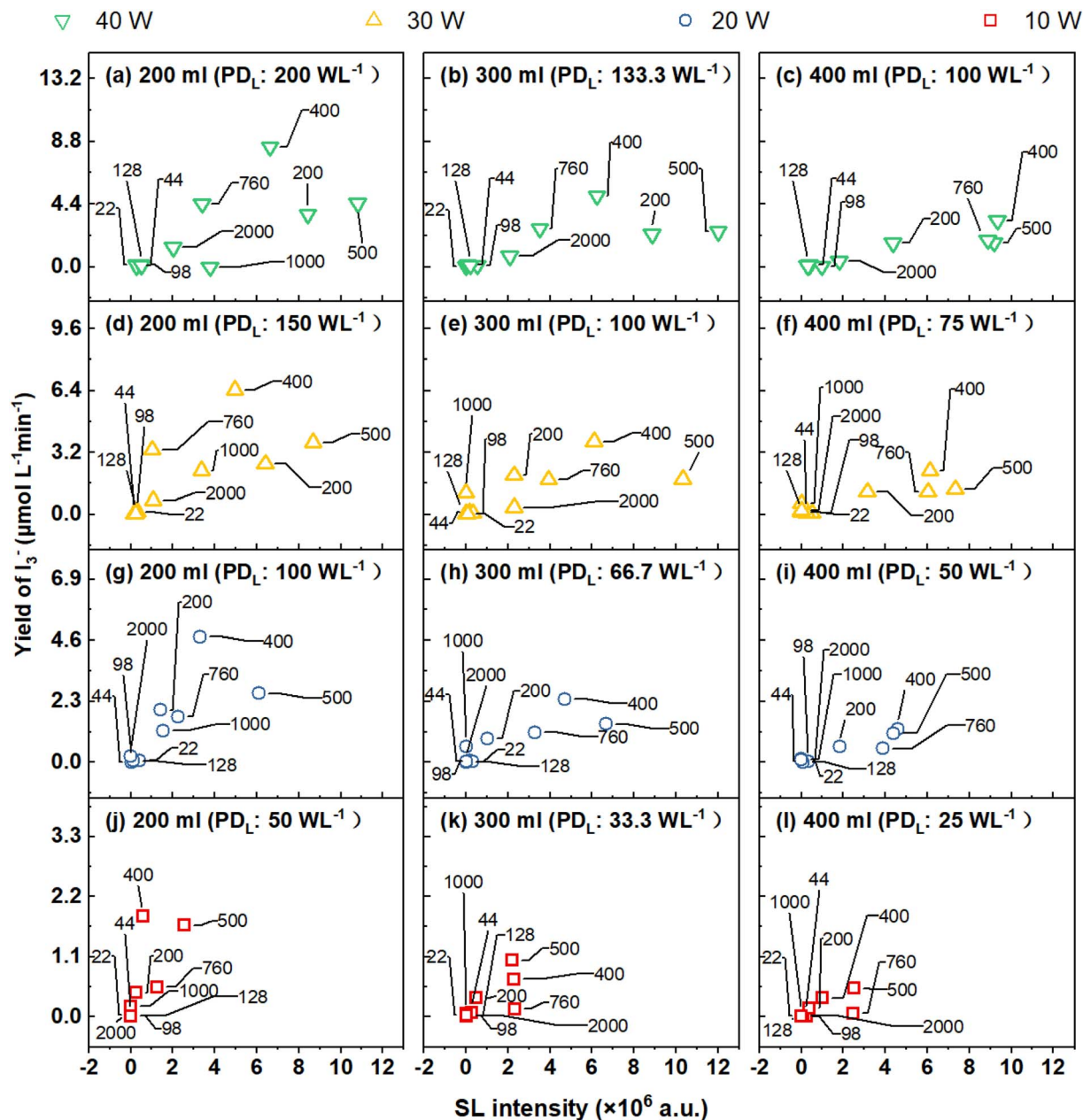


Fig. 17 The correlation between I_3^- and SL generated during sonication (testing frequencies are labelled (kHz)): (a–c) 40 W, (d–f) 30 W, (g–i) 20 W and (j–l) 10 W.

Besides the variation in the reaction's location, SCL and KI dosimetry may exhibit distinct excitation conditions. In Fig. 16, each row depicts a system with the same power input but decreasing power densities and increased volume (left to right). As the liquid height increased/power density decreased, SCL intensity increased (x -axis) rose dosimetry yields declined (y -axis). This suggests that with an ultrasonic system, SCL is more likely to happen than dosimetry.

3.3.3 Correlation between I_3^- yield and SL intensity. Across all tested cases, generation of I_3^- reaches a maximum value as the intensity of SL increases. For example, in Fig. 17(e and l) SL intensity increase to the maximum at 400 and 500 kHz, respectively. Correspondingly, the I_3^- yield reaches a peak value

under the same power and liquid height. Yet, beyond this point, the correlation of I_3^- yield and SL is not observed, further increases in SL intensity occur when there is a decline in I_3^- production. Interestingly, in general circumstances (with a constant volume and power for the reaction), the point where the trends of SL and I_3^- yield diverge is observed at an acoustic frequency of 400 kHz (Fig. 17). However, when a lower power (10 W) and a larger volume of reactor (≥ 300 ml) are employed, the point of divergence of trends is observed at 500 kHz (Fig. 17(h and l)).

The production of both ROS and SL are affected by the number of active bubbles and collapse intensity in multi bubble system, which is complex. Both are influenced by the



interaction between the multi-parametric effects, *i.e.* driving effects, physical force and so on.²² The maximum temperature of the active bubbles is thought to be the primary determinant of SL intensity⁵⁴ whereas the average temperature inside the bubble might represent the primary factor influencing the generation of most radicals represented by 'OH'.⁹⁷ Once the temperature exceeds the optimal temperature for radical yield, the oxidation reactions will be diminished through secondary processes, such as nitrogen oxidation or super-oxidation reaction.⁹⁸ Power density seems to be an important factor influencing the SL and I_3^- yield's positive correlation divergence point. In our settings, the oxidative characteristics of the reaction induced by ultrasound at a power density of 50 W L^{-1} result in I_3^- yields practically identical at 400 and 500 kHz (Fig. 17(i and j)). This slight difference may stem from variations in axial direction energy attenuation attributable to the liquid level's height.⁹⁹ In the studied system, the divergence points of the SL and I_3^- correlations consistently occur at 400 kHz for $PD_L > 50 \text{ W L}^{-1}$ (Fig. 14(a–h)), and at 500 kHz for $PD_L < 50 \text{ W L}^{-1}$ (Fig. 14(k and l)). The effect of the liquid height appears to be more reflected in the impact of the reactor volume on power density. 500 kHz systems have greater quasi-acoustic and acoustic streaming compared to 400 kHz system.^{22,66} This liquid flow pattern, which facilitates bubble de-conglomeration and oscillation and enhances free radical release, has the potential to facilitate the sonochemical activity under low pressures for cavitation.^{89,100}

3.3.4 Interrelationship between bubbles with different properties in the same system. SL is a measurement of the energy state of bubbles, which is determined by the physical effects caused by temperature and pressure within the bubbles. It serves as a fundamental indicator of the level of cavitation activity during sonication. The outcomes of SL have a high positive correlation with those of SCL and dosimetry, which is confirmed through the correlation coefficients, *i.e.* $\rho_{\text{total ROS yield vs. SL intensity}} = 0.78$, $\rho_{I_3^- \text{ yield vs. SL intensity}} = 0.82$ and $\rho_{\text{SCL intensity vs. SL intensity}} = 0.71$. Dosimetry and SCL are distinct sonochemical reaction mechanisms. The SL and dosimetry correlation analyses indicate various sonochemical processes have distinct responses with respect to changing bubble properties (Fig. 9 and 12).

The ultrasonic system can be categorized into three main regions for chemical reactions, inside the bubble core, at the interfacial region and in the bulk solution. The intense environmental conditions in the bubble core play a crucial role in the breakdown of gas molecules, generating free radicals.²⁵ Simultaneously, a portion of the generated radicals will also experience reforming reactions (Fig. 12). This result is confirmed by the high correlation between ROS generation and SL intensity. A high intensity of SCL indicates that more reactions occur at the interface between the gas and liquid phases. However, this reactivity observed at the gas–liquid interface is not consistent with the reactivity of the entire solution (bulk). At 500 kHz, the SL output demonstrates optimal results, indicating effective cavitation oscillation and collapse within the system. The equivalent outcomes of the SCL corroborate this. Nevertheless, the dosimetry data did not exhibit consistency, and the results revealed a lower oxidation of iodine ions at 500 kHz. The dosimetry data

demonstrated effective iodide oxidation in the system with 400 or 760 kHz, unlike the optimal SCL (*i.e.* at 500 kHz). According to the preceding discussion, systems that favour one activity (SCL/SL/ROS *etc.*) may not be beneficial for another as previously observed.⁴ Although such interactions must be based on the adequate cavitation level within the specified system. Therefore, the sonochemical activity that drives SCL is likely different from the ultrasonic oxidation by ROS.

4. Conclusions

Varying parameter configurations can result in variations in the bubble properties of the reaction system. Sono(chemi)luminescence and KI dosimetry allow for the visualization and quantification of these changes and reflect the acoustic process. This work introduces a novel method for digitizing sono(chemi)luminescence images, which allows for the quantification of SL and SCL data across a broader range by selectively targeting regions.

The extensive experimental data and correlation coefficients were used to examine the variations in the behaviour of bubbles created by ultrasound under different operating settings. Based on the results of three types of sonochemical characteristics (SL, SCL and ROS), the sonic reactivity is categorized into three distinct zones according to the frequency of irradiation (low-frequency: $22 \text{ kHz} \leq f \leq 128 \text{ kHz}$, medium-frequency: $200 \text{ kHz} \leq f \leq 760 \text{ kHz}$ and high-frequency: $1000 \text{ kHz} \leq f$). It has been verified that acoustic systems exposed to intermediate frequencies ($200 \text{ kHz} \leq f \leq 760 \text{ kHz}$) typically exhibit greater total cavitation intensity and reactivity.

The results of independent cavitation characteristics gained under varied operating parameters were used for the computation of correlation coefficients (r and ρ) to explore their relationship. The distinction between the varieties of sonochemical reactions was validated through the low coefficient value, like $r_{I_3^- \text{ yield vs. SCL intensity}} = 0.46$ and $r_{H_2O_2 \text{ yield vs. SCL intensity}} = 0.41$. This difference can be attributed to variation in the reaction's location, and alternate reaction energies and mechanisms.

The Spearman's rank correlation coefficient of 0.78 between total ROS generation and SL intensity suggests a robust positive correlation between these two cavitation characteristics. The $\rho_{I_3^- \text{ yield vs. SL intensity}} = 0.82 > \rho_{H_2O_2 \text{ yield vs. SL intensity}} = 0.56$ indicates that this positive connection mostly arises from the positive relationship between I_3^- yield and SL intensity. IORS are the primary ROS generated by ultrasonic irradiation, as corroborated by the correlation coefficient values ($r_{I_3^- \text{ yield vs. ROS}} = 0.99$ and $\rho_{I_3^- \text{ yield vs. ROS}} = 0.98$). This positive correlation consistently diverges at a point dependent on the system parameters, which in our experimental configuration may occur at 400 or 500 kHz, based on each system's power density.

Data availability

Data will be made available on request.

Conflicts of interest

There are no conflicts to declare.



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