ULTRASONIC CHARACTERIZATION OF CELL PELLET BIOPHANTOMS AND TUMORS USING QUANTITATIVE ULTRASOUND MODELS

BY

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DISSEPTION

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This dissertation investigates the ultrasonic scattering from tumors. Tumors are complex acoustic scattering media and to date there has not been an adequate acoustic scattering model that fits them well. The tumor scattering mechanisms are studied herein by a step-wise approach: starting from simple (low-concentration cell pellet biophantoms) to moderately complex (high-concentration cell pellets) and to significantly complex media (actual tumors).

The comparison between the backscattering coefficients (BSC) of lower- and higher-concentration cell pellet biophantoms suggest that high concentration of cells could lead to correlation in cell positions, causing additional scattering that may be modeled by the structure function. The structure function for the high-concentration cell pellet biophantom was isolated by comparing the BSC of the high-concentration biophantom to that of a very low-concentration biophantom where the cell spatial positions are assumed to be random. Polydisperse structure function models that take into account the polydispersity of scatterer radii are introduced and shown to be consistent with the structure function curves measured from BSC. The two-dimensional (2D) structure function is estimated from the histological sections, and shown to have a similar shape to the theoretical structure function and to that calculated from BSC.

The comparison between cell pellets and tumors of the same cell type suggests that the anatomic details in tumors must be taken into account for modeling
purposes, in addition to the scattering from cells. Also, histology studies suggest that the structure functions in tumors are slightly different from those in cell pellets: the tumor cell spatial arrangement is slightly more random compared to cell pellets. The effect of the structure function on parameter estimation is discussed. Further work is shown to be required for modeling the tumor structure function.

Additionally, the comparison between different tumor types shows that ultrasound backscattering is sensitive to unique tumor structures. The EHS tumor has a distribution of clustered cells and shows a different BSC and structure function pattern than the tumors that have a homogenous distribution of cells. A scattering model is developed to detect the clustering feature.

Overall, the dissertation improves our understanding of the acoustic scattering mechanisms in tumors, and improves the tumor scattering modeling.
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<tr>
<td>2D</td>
<td>Two-dimensional</td>
</tr>
<tr>
<td>2DZM</td>
<td>Two-dimensional impedance map</td>
</tr>
<tr>
<td>3D</td>
<td>Three-dimensional</td>
</tr>
<tr>
<td>3DZM</td>
<td>Three-dimensional impedance map</td>
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<tr>
<td>BSC</td>
<td>Backscattering coefficient</td>
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<td>CHO</td>
<td>Chinese hamster ovary</td>
</tr>
<tr>
<td>DPBS</td>
<td>Dulbecco’s phosphate buffered saline</td>
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<tr>
<td>EAC</td>
<td>Effective acoustic concentration</td>
</tr>
<tr>
<td>EHS</td>
<td>Engelbreth-Holm-Swarm</td>
</tr>
<tr>
<td>ESD</td>
<td>Effective scatterer diameter</td>
</tr>
<tr>
<td>H&amp;E</td>
<td>Hematoxylin and eosin</td>
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<tr>
<td>FF</td>
<td>Fluid-filled</td>
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<tr>
<td>MAT</td>
<td>13762 MAT B III</td>
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<tr>
<td>NS</td>
<td>Neyman-Scott</td>
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<tr>
<td>OZ</td>
<td>Ornstein-Zernike</td>
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<tr>
<td>PY</td>
<td>Percus-Yevick</td>
</tr>
<tr>
<td>QUS</td>
<td>Quantitative ultrasound</td>
</tr>
<tr>
<td>ROI</td>
<td>Region of interest</td>
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RF Radio frequency
CHAPTER 1

INTRODUCTION

1.1 Quantitative Ultrasound

Diagnostic ultrasound is a significant modality in the medical imaging world. It is safe, noninvasive, inexpensive and easily accessible. One of the most widely used techniques in conventional ultrasound is known as B-mode imaging [1]. Acoustic reflection or scattering occur when there is an acoustic impedance (defined as the sound speed times the mass density) contrast between two tissue layers. B-mode images qualitatively display the brightness of the radio-frequency (RF) echo signal, allowing for medical diagnosis. The B-mode image formation process is simple and reliable. Medical diagnosis using B-mode imaging, however, is often highly subjective and operator-dependent because of its qualitative nature.

There are many approaches to constructing quantitative ultrasonic images directly from the qualitative B-mode images. Image processing techniques such as texture analysis has been explored for many years to extract image features such as first- and second-order parameters including the mean, standard deviation, entropy, and run length parameters [2]-[4]. However, these approaches only had limited success, because the extracted quantitative parameters are system-dependent.
To obtain operator- and system-independent quantitative information from the tissue under investigation would require alternative ways of processing the RF data. One approach is to utilize the frequency-dependent information of the RF data and estimate the attenuation and backscatter coefficient (BSC) which are intrinsic properties of the tissue and are not dependent on the ultrasonic system or the operator [5], [6]. Further, acoustic scattering models that mimic tissue anatomic structures can be fitted to the BSC versus frequency curve to estimate acoustic properties of the scatterers, such scatterer size, shape, number density, and acoustic impedance. The attenuation, BSC, and model-derived acoustic parameters have been used for charactering of the eye [7], [8], prostate [9], kidney [10], heart [11], [12], blood [13]-[16], breast [17]-[20], liver [21]-[23], cancerous lymph nodes [24], and apoptotic cells [25], [26], and for evaluating disease treatment [27].

1.2 Limitations of Current QUS Models

Further success of model-based QUS techniques relies on the understanding of the tissue scattering mechanisms and the development of appropriate scattering models that match the anatomic tissues structure. To date, the scattering mechanisms in biological tissues have not been well understood, and the currently available scattering models for QUS have limitations.

First, the major scattering site has not been fully identified in many cases. Applying a simple scattering model such as the spherical Gaussian model [6] or the fluid-fill sphere model [6] to the tissue would not yield the best result. The
simple model may not fit to the BSC data, and the physical/biological meanings of model parameters are not always clear. For instance, the effective scatterer diameter (ESD) derived from many QUS models could be interpreted as the diameter of the cell nucleus, of the whole cell, or of other tissue structures, when the physical sites responsible for scattering are not identified. Although the estimated ESD may still be valuable for differentiation purposes [17], [18], the diagnostic potential is limited and will be significantly improved if the model parameter is correlated to a specific anatomic site or physiological state.

Second, major factors that may contribute to scattering have not been fully explored. It has been well established in physics that the scatterer size, shape, acoustic properties, number density, and spatial arrangement can all contribute to scattering. Yet for tissues which factors are essential to scattering and which factors are not has not been fully explored. This information is critical to scattering modeling, because essentially all models have to take into account the most important factors while ignoring the least important.

Third, many currently available models are limited to the Rayleigh region ($ka \ll 1$, where $k$ is the wave number and $a$ is the scatterer size). Although the low-frequency condition is satisfied for current clinical systems that operated in frequencies lower than 20 MHz, models that deal with higher frequencies are required with the emergence of high-frequency (>20 MHz) ultrasound [28]. In fact, studying the scattering at higher frequencies will help to understand the scattering mechanisms as well. Characterizing tissues at high frequency may yield
tissue property information that cannot otherwise be extracted from low-frequency signals.

1.3 The Proposed Study

The purpose of this dissertation is to improve our understanding of the scattering process in tissues, and eventually develop more accurate scattering models. Biological tissues are diverse and complex. This dissertation is focused on mammary tumors. Tumors are complex scattering media and to date there has not been an adequate scattering model that fits it well. Studying the scattering of tumors has medical significance: inexpensive and non-invasive tools are needed for tumor diagnosis, and QUS has the potential to provide such a tool to diagnose and classify tumors with high sensitivity and specificity.

There are several approaches to understanding tissue scattering. The three-dimensional acoustic impedance map (3DZM) [29]-[31] has been used to identify the scattering site. The 3DZMs were created by aligning serial photomicrographs of stained histologic tumor sections. Acoustic impedance values were assigned to the different stained colors. Another approach is to measure acoustic properties on various tissue/cell components. For instance, the sound speed of isolated nuclei and whole cells has been measured to study if the nucleus or the whole cell is the scattering site [32]. Furthermore, simulations have been applied to study the scattering process. Vlad et al. [33] performed two-dimensional simulations to study how the cellular size variance influences ultrasound backscatter. Saha et al. [34] performed a three-dimensional simulation to produce BSC from red blood
clusters, where the red blood cells in a cluster were stacked following the hexagonal close packing structure. More complex studies include three-dimensional simulation of the scattering for both longitudinal and shear waves [35].

The cell pellet technique is another approach to understanding tissue scattering. It has been demonstrated that a model termed the concentric-sphere model that matches the geometry of a eukaryotic cell is accurate for low-concentration cell pellet biophantoms. The biophantoms consist of live cells embedded in a plasma-thrombin supportive background [36]. The BSC increases linearly with cell concentration. The follow-up study [37] showed that the linear relationship between BSC and cell concentration does not hold when the concentration is high.

Based on the cell pellet technique, in this dissertation we will use a step-wise approach to systematically studying the scattering from tumors, i.e., to dissect the scattering by analyzing at one time each factor that may significantly contribute to scattering. To that end, we will compare the scattering from simple media (low concentrations of cells), moderately complex media (high concentrations of cells), and significantly complex media (actual tissue/tumors). The study of low-concentration cells [36] already addressed the problem of what is responsible for scattering from low-concentration cell pellet biophantoms. The comparison of various concentrations may give insight into the effect of concentration on scattering: cell concentration could affect the spatial distribution of cells. The comparison between cells and actual tissue/tumors may demonstrate what is unique in a specific tissue that contributes to scattering. The overall hypothesis is
that the scattering from tumors is dependent on three aspects: the shape and acoustic properties of tumor cells, the spatial organization of tumor cells, and the unique anatomic structure of each tumor type. The comparison among different degrees of scattering media complexity can test the overall hypothesis, and lead to improved modeling.

The overall experimental approach is to construct a variety of samples, scan the samples using high-frequency ultrasound transducers (10 – 105 MHz), fix the samples for further histology analysis, estimate the attenuation and BSC from the RF data, compare the BSC of different samples, study the histological slides, and develop and test scattering models.

The dissertation is organized as follows: Chapter 2 introduces basic scattering theory and reviews acoustic scattering models applied to biological media. Chapter 3 compares the scattering from low- and high-concentration cell pellet biophantoms, and develops polydisperse structure function models to address the effect of cell spatial distribution. Chapter 4 presents the BSC results of various tumor types, compares the tumor BSC to the cell pellet BSC when such a comparison is possible, and shows how unique tumor structures can affect scattering. Chapter 5 introduces a histology-based method for estimating the structure function, which provides independent evidence supporting the analytical structure functions developed in Chapter 3, and provides guidelines for developing scattering models for tumors. Chapter 6 addresses the issue of how to develop scattering models that take into account unique tumor structures.
This chapter reviews the acoustic scattering theories related to soft biological tissues. The aim is to predict the backscattering coefficient (BSC). The scatterers are assumed to be discrete particles. Concepts of form factor and structure function are introduced. The effect of scatterer size polydispersity is also discussed.

2.1 Backscattering Coefficient

Consider a plane wave of unit amplitude incident on a scattering volume $V$ that contains $N$ discrete scatterers. The total scattered field far from the scattering volume behaves as a spherical wave (Equation (4) of [6]):

$$p_s(r) = \frac{e^{ikr}}{R} \sum_{j=1}^{N} \Phi_j(K)e^{ikr_j}, \quad (2.1)$$

where $r$ is the observation position with respect to the origin, $R = |r|$, $r_j$ is the position of the $j$th scatterer, $k$ is the propagation constant ($k = \omega/c$ where $\omega$ is the angular frequency and $c$ is the propagation speed). The factor $\Phi_j(K)$ is the scattering amplitude of the $j$th scatterer and describes the spatial frequency.

dependence of the scattered pressure; $\Phi_j$ is a function of the scattering vector $K$ whose magnitude is given by $|K| = 2k \sin(\theta/2)$, where $\theta$ is the scattering angle ($\theta = \pi$ for backscattering). $\Phi_j$ is dependent on the scatterer size, shape and acoustic properties.

The differential cross section per unit volume $\sigma_d$ (i.e., the power scattered into a unit solid angle observed far from the scattering volume divided by the product of the incident intensity and the scattering volume) may be expressed as

$$\sigma_d(K) = \frac{R^2 I_r}{V I_0} = \frac{1}{V} \left| \sum_{j=1}^{N} \Phi_j(K) e^{iK \cdot r_j} \right|^2,$$

(2.2)

where $I_r$ and $I_0$ denote the scattering intensity and incident intensity, respectively, and $\left| \right|^2$ represents the squared modulus of the quantity.

BSC is defined as the differential cross section per unit volume to backscattering direction ($|K| = 2k$). The frequency-dependent BSC is an intrinsic property of tissue, describes how effectively the tissue backscatters ultrasound, and can be obtained experimentally by analyzing the RF data.

2.2 Uncorrelated Scatterers

If the scatterers are spatially uncorrelated (which often occurs when the scattering volume contains a sparse concentration of scatterers without clustering), the phase terms $e^{iK \cdot r_j}$ in Equation (2.2) may be assumed to be
uncorrelated. The differential cross section per unit volume for this case is denoted as $\sigma_{d,\text{incoherent}}$, and may be expressed as

$$\sigma_{d,\text{incoherent}}(K) = \frac{1}{V} \sum_{j=1}^{N} |\Phi_j(K)|^2,$$

(2.3)

which becomes

$$BSC_{\text{incoherent}}(2k) = \frac{1}{V} \sum_{j=1}^{N} |\Phi_j(2k)|^2,$$

(2.4)

for backscattering.

2.2.1 Exact solutions

For a simple scatterer shape, the exact solution of the scattering amplitude $\Phi_j(2k)$ may be calculated precisely by solving the wave equation. Commonly used scatterer shapes include the fluid sphere filled in fluid [38], solid cylinders and spheres [39], and two-fluid-concentric spheres in fluid [40]. The exact solutions have complex expressions with multiple parameters. The simplest case, the fluid sphere in fluid, requires at least five parameters: the scatterer radius, the speed of sound and mass density of the scatterer, and the speed of sound and mass density of the background.

Figure 2.1 gives an example of what the BSC versus frequency curve looks like for the fluid-filled spheres and the concentric spheres using the exact solutions. Note that the spatial frequency $k$ is converted to temporal frequency in MHz, and monodisperse scatterers are assumed.
2.2.2 Approximate solutions

Alternatively, a simpler expression of the scattering amplitude $\Phi_j(K)$ may be calculated from the spatial autocorrelation function, if the scatterers have acoustic property values (density, $\rho$, and compressibility, $\kappa$) very close to those of the background medium, and Equation (2.3) becomes (Equation (12a) of [6]),

$$\sigma_{d,\text{incoherent}}(K) = \frac{k^4 V_s}{16\pi^2} \bar{n} \gamma_0^2 \int_{-\infty}^{\infty} b_\gamma(\Delta r)e^{-iK \cdot \Delta r} d^3 \Delta r,$$  

(2.5)

where $\bar{n}$ is the scatterer number density (i.e., average number of scatterers per unit volume), $V_s$ is the average scatterer volume, $\gamma_0^2$ is the mean-square variation in acoustic impedance per scatterer, and $b_\gamma(\Delta r)$ is the spatial autocorrelation function of the scattering volume and depends on the scatterer shape. The integral $\int_{-\infty}^{\infty} b_\gamma(\Delta r)e^{-iK \cdot \Delta r} d^3 \Delta r$ is the form factor, which approaches unity when the frequency approaches zero.

For backscattering, Equation (2.5) becomes

$$BSC_{\text{incoherent}}(2k) = \frac{k^4 V_s}{16\pi^2} \bar{n} \gamma_0^2 F(2k),$$  

(2.6)

where $F(2k)$ is the form factor. Common form factors are [6]

$$F_1(2k) = \left[ \frac{3j_1(2ka)}{2ka} \right]^2 \text{ (fluid sphere)}$$  

(2.7)

$$F_2(2k) = \left[ j_1(2ka) \right]^2 \text{ (spherical shell)}$$  

(2.8)
The BSC expressions in Equations (2.6) – (2.9) have fewer parameters and simpler forms than the exact solution does. The simpler expressions are applicable under the condition that the acoustic property contrast is low between the scatterer and the background. Figure 2.2 compares the fluid-filled sphere BSC calculated using the exact solution using the Anderson model [38] to that calculated using the low-contrast approximation using Equations (2.6) and (2.7). As the impedance contrast between the sphere and the background increases, the approximate solution becomes less accurate. Monodisperse spheres (radius = 7 µm) are assumed for calculation.

2.2.3 The effect of size distribution

Typical BSC versus frequency curves have peaks and dips, and the positions of the peaks and dips are primarily determined by the scatterer size for a given sound speed. If the scatterers are polydisperse in size, a distribution of scatterer size will result in smoother peaks and dips. Figure 2.3 shows the polydisperse fluid-filled sphere BSC calculated using $BSC_{\text{dist}}(2k) = \int_{0}^{\infty} BSC(2k, x) f(x) dx$, where $BSC_{\text{dist}}(2k)$ is the BSC of the polydisperse spheres, $x$ is the spherical radius, $f(x)$ is the probability density function of the spherical radius, and $BSC(2k, x)$ is the BSC of monodisperse spheres that have a radius $x$. The quantity $BSC(2k, x)$ was calculated using the exact solution (the Anderson model [38]) and the low-contrast approximation (Equations (2.6) and (2.7)), respectively. The sphere radius is assumed to have a uniform distribution between 6 and 8 µm. The
size distribution makes the sharp dip (around 80 MHz) appear smoother comparing Figure 2.2 and Figure 2.3. Further, the discrepancy between the exact solution and the low-contrast approximate is decreased when the scatterers are polydisperse compared to monodisperse. This result suggests that the low-contrast approximate result may be used for polydisperse scatterers if fewer parameters are desired for the model.

2.3 Correlated Scatterers

The spatial positions of scatterers are often correlated. The cell positions are strongly correlated if the concentration is high. Spatial correlation in scatterer positions can also occur at low concentrations, for example, when there is clustering.

2.3.1 Structure function

If the scatterers are spatially correlated, we cannot proceed to Equation (2.3) from Equation (2.2). Dividing Equation (2.2) by Equation (2.3) yields the three-dimensional (3D) structure function:

\[ S(K) = \frac{\sigma_d(K)}{\sigma_{d, \text{incoherent}}(K)}, \quad (2.10) \]

which is

\[ S(2k) = \frac{BSC(2k)}{BSC_{\text{incoherent}}(2k)} \quad (2.11) \]

for backscattering.
The structure function defined in Equations (2.10) and (2.11) is a quantity describing the effect on scattering caused by the pattern of the spatial arrangement of the scatterers. If we assume that the scattering amplitudes $\Phi_j(K)$ are identical for all the scatterers, Equation (2.2) may be simplified as

$$\sigma_d(K) = \bar{n} |\Phi_j(K)|^2 \frac{1}{N} \left( \sum_{j=1}^{N} e^{\mathbf{kr}_j} \right) \left( \sum_{j=1}^{N} e^{-\mathbf{kr}_j} \right),$$

(2.12)

where $\bar{n} = \frac{N}{V}$ is the number density of the scatterers. By substituting Equation (2.3) and Equation (2.12) into Equation (2.5), and making appropriate simplifications, the structure function may be expressed as

$$S(K) = \frac{1}{N} \left( \sum_{j=1}^{N} e^{i\mathbf{kr}_j} \right) \left( \sum_{j=1}^{N} e^{-i\mathbf{kr}_j} \right),$$

(2.13)

where the structure function is determined by the scatterer positions, and is not dependent on the scattering amplitude $\Phi_j(K)$.

If the exact position of each scatterer is known, the structure function can be calculated deterministically. Note that Equation (2.13) may also be expressed as

$$S(K) = \frac{1}{N} \left| \sum_{j=1}^{N} e^{-i\mathbf{kr}_j} \right|^2,$$

(2.14)

which means that the structure function is simply the squared modulus of the Fourier transform of the scatterer positions. Equation (2.14) is used in Chapter 5 to estimate the structure function of a tissue from a 2D histology slide.
If the exact positions of the scatterers are unknown, the structure function has to be determined statistically. In this case, the structure function can be calculated from the statistical distribution (e.g., pair correlation function) of the scatterer positions. Equation (2.13) is mathematically equivalent to [41]

$$S(K) = 1 + \overline{n} \int [g(r) - 1] e^{-Kr} \, dr,$$

(2.15)

where the structure function is expressed in terms of the pair-correlation function $g(r)$, a quantity related to the probability of finding two scatterers separated by the distance $r$. With Equation (2.15), the structure function can be interpreted as the 3D Fourier transform of the total correlation function $h(r) = g(r) - 1$. The total correlation may be obtained by solving the set of equations formed by the Ornstein-Zernike (OZ) integral equation [42] and a closure relation. The OZ equation splits the total correlation $h(r)$ into the direct correlation $c(r)$ and the indirect correlation by the equation

$$h(r) = c(r) + \overline{n} \int_0^\infty h(|r - r'|) c(|r'|) \, dr'.$$

The closure relation couples the same quantities $h$ and $r$. Depending on different models of closure relation, the structure function will be different.

2.3.2 Structure function models

A number of structure function models have been successfully used in ultrasonic tissue characterization, particularly in blood characterization. The Percus-Yevick (PY) structure function [43] is typically used to model the structure of dense medium that does not have clustering. To address the clustering effect, the structure function derived from the Neyman-Scott (NS) point process was used for blood characterization [44]. Additionally, a second-order Taylor
approximation of structure function [45] was developed to achieve a mathematically simple expression.

**A) Percus-Yevick approximation**

The PY approximation is a commonly used closure relation valid for a random distribution of non-overlapping spheres (the sphere positions are not truly random because they are non-overlapping). With PY closure, an analytical expression of the structure function has been obtained for backscattering [46]-[48]:

\[
S_{PY}(2k) = \frac{1}{1-nC(2k)} ,
\]

\[
C(2k) = -32\pi a^3 \int_0^1 s^3 \frac{\sin(4kas)}{4kas} \left( \alpha + \beta s + \gamma s^3 \right) ds ,
\]

\[
\eta = \frac{4\pi a^3}{3} \bar{n}, \quad \alpha = \frac{(1+2\eta)^2}{(1-\eta)^4}, \quad \beta = -\frac{6\eta(1+\eta/2)^2}{(1-\eta)^4}, \quad \gamma = \frac{\eta(1+2\eta)^2}{2(1-\eta)^4} ,
\]

where \(a\) is the sphere radius, \(s\) is a dummy variable of integration, and \(\eta\) is the sphere volume fraction.

Figure 2.4 shows a comparison among the PY structure functions at various volume fractions (similar results can be found in [47] and [48]. The PY structure function approaches unity as the concentration becomes extremely low. A peak starts to appear around \(ka = 2\) as the concentration increases. The peak becomes relatively sharp as the concentration becomes considerably large. This sharp peak is not observed from tissue data, because the scatterers in tissue are polydisperse in nature, whereas the PY structure function given in Equations (2.16) – (2.18) is
only valid for identical scatterers. Structure functions that are based on the PY closure relation but take into account the scatterer polydispersity are developed in Chapter 3, and applied to the high-concentration cell-pellet biophantoms.

**B) Neyman-Scott point process**

The NS process is a random point process commonly used to model clusters of points [44]. The NS process applied in [44] assumes that the spatial distribution of cluster centers follows a Poison distribution, and the spheres within the cluster follow a Gaussian distribution (more spheres in the cluster center). The resulting structure function is given by

\[
S_{NS}(2k) = 1 + (W - 1)e^{-2(ka)^2/\Delta^2} ,
\]

(2.19)

\[
W = n_c + (\sigma_c^2 / n_c) ,
\]

(2.20)

\[
\Delta = \frac{1}{\sqrt{2}} (n_c - 1)^{1/D_f} ,
\]

(2.21)

where \(n_c\) and \(\sigma_c^2\) are the mean and variance of the number of scatterers per cluster, \(D_f\) is the fractal dimension that morphologically characterizes the growth process of the aggregates (\(D_f \approx 3\) for compact spherical aggregates), \(W\) is the packing factor which increases when the number of cells per aggregate grows, and \(\Delta\) is the size factor which decreases when the spatial dimension of the cluster increases.

Figure 2.5 plots the NS structure functions calculated using four sets of parameters. The NS structure function approaches unity when the average number
of scatterers per cluster is one. This observation means that the NS structure function only models the clustering effect, but does not take into account the concentration effect on scatterer position correlation. Therefore, it only works for low scatterer concentrations (volume fraction less than 5%). As the number of scatterers per cluster increases, the structure function increases in the low-frequency range \((ka < 0.5)\), whereas it remains unity at the high-frequency range.

**C) Second-order Taylor approximation**

A second-order Taylor approximation of structure function was given in [45] as

\[
S_{Taylor}(2k) \approx W - \frac{12}{5} D^2 (ka)^2, \tag{2.22}
\]

where \(W\) is the packing factor, and \(D\) equals the aggregate radius divided by the scatterer radius. \(W\) and \(D\) have been shown to increase during blood aggregation [45].

Figure 2.6 plots the structure functions calculated by Equation (2.22) using four sets of parameters picked from Table V of [45].

**2.4 Chapter Summary**

The scattering from soft tissue is affected not only by the properties of individual scatterers (e.g., scatterer size, shape, acoustic impedance contrast), but also by the scatterer position correlation. The scatterer positions can be significantly affected by clustering, high concentration, etc. Success has been achieved in literature using the structure function to address the scatterer position
correlation issue. However, limitations also exist in available structure function models.

2.5 Figures

![Figure 2.1 Comparison of calculated BSC versus frequency curves for the concentric sphere model and the single fluid-filled sphere model using the exact solutions. The parameters used for the concentric sphere (blue solid) are: inner sphere radius $a_{\text{in}} = 4 \ \mu\text{m}$, outer shell radius $a_{\text{out}} = 7 \ \mu\text{m}$, inner sphere impedance $Z_{\text{in}} = 1.6 \ \text{Mrayl}$ (density: 1.03 g/cm$^3$; sound speed: 1550 m/s), outer shell impedance $Z_{\text{out}} = 1.55 \ \text{Mrayl}$ (density: 1.01 g/cm$^3$; sound speed 1535 m/s), background impedance $Z_{0} = 1.5 \ \text{Mrayl}$ (density: 1 g/cm$^3$; sound speed 1500 m/s), and number density $n = 2 \times 10^7 \ \text{cm}^{-3}$. The parameters used for the larger fluid-filled sphere (dotted magenta) are: radius $a = 4 \ \mu\text{m}$, sphere impedance $Z = 1.55 \ \text{Mrayl}$ (density: 1.01 g/cm$^3$; sound speed 1535 m/s), background impedance $Z_{0} = 1.5 \ \text{Mrayl}$ (density: 1 g/cm$^3$; sound speed 1500 m/s), and number density $n = 2 \times 10^7 \ \text{cm}^{-3}$. The parameters used for the smaller fluid-filled sphere (dashed red) are the same as the larger one except that the sphere radius is set to be 4 $\mu\text{m}$.](image-url)
Figure 2.2 Comparison of monodisperse fluid-filled sphere BSC calculated using the exact solution (blue solid) to that calculated using the low-contrast approximate (red dashed). The sphere radius is 7 µm, the number density is \( n = 2 \times 10^7 \text{ cm}^{-3} \), and the background impedance is 1.5 Mrayl (density: 1 g/cm\(^3\); sound speed 1500 m/s). The impedance contrast between the sphere and the background increases from (a) to (c): the sphere impedance is 1.52 Mrayl (density: 1.01 g/cm\(^3\); sound speed 1505 m/s) for (a), 1.6 Mrayl (density: 1.03 g/cm\(^3\); sound speed 1550 m/s) for (b), and 1.68 Mrayl (density: 1.05 g/cm\(^3\); sound speed 1600 m/s) for (c).
Figure 2.3 Comparison of polydisperse fluid-filled sphere BSC calculated using the exact solution (blue solid) to that calculated using the low-contrast approximate (red dashed). The parameters are the same as Figure 2.2, except that the sphere radius is assumed to following a uniform distribution between 6 and 8 µm.
Figure 2.4 Comparison among PY structure functions at five different volume fractions: 1%, 10%, 50%, 60%, and 74%, computed using Equations (2.16) – (2.18).
Figure 2.5 Comparison among NS structure functions for four parameter sets: (a) \( n_c = 1, \sigma_c^2 = 0 \), (b) \( n_c = 5, \sigma_c^2 = 4 \), (c) \( n_c = 15, \sigma_c^2 = 4 \), and (d) \( n_c = 5, \sigma_c^2 = 40 \), using Equations (2.19) – (2.21). \( D_j \) is assumed to be 3 for all four curves.
Figure 2.6 Comparison among structure functions calculated by second-order Taylor approximation using four parameter sets: (a) $W = 1.4$, $D = 2.4$, (b) $W = 6.8$, $D = 5.3$, (c) $W = 0.5$, $D = 1.1$, and (d) $W = 0.2$, $D = 0.7$. 
CHAPTER 3
THE SCATTERING OF HIGH-CONCENTRATION BIOPHANTOMS

This chapter addresses the problem of dense media scattering using structure functions. The effect of scatterer polydispersity on the structure functions is investigated. Structure function models based on polydisperse scatterers are theoretically developed and experimentally evaluated against the structure functions obtained from cell pellet biophantoms.

3.1 Background and Introduction

A previous study [36] demonstrated that a concentric-sphere model that matches the geometry of a eukaryotic cell is accurate for low-concentration cell pellet biophantoms that consist of live cells embedded in a plasma-thrombin supportive background. The study [36] also showed that the ultrasonic backscatter coefficient (BSC) increases linearly with cell concentration. The follow-up study [37] showed that the linear relationship between BSC and cell concentration does not hold when the concentration is high. There was no adequate model that could apply to the high-concentration cell pellet biophantoms. In the meantime,

modeling such dense media is of practical significance because the cell concentration is high in real tissues such as mammary tumors.

Given the acoustic scattering theory reviewed in Chapter 2, it is hypothesized that higher concentration causes stronger spatial correlation in the cell positions. The structure function could be useful to address this problem.

The structure function has been applied in a number of areas of ultrasonic scattering. To give a more comprehensive review than what have been mentioned in Chapter 2, the concept of structure function was introduced for the first time by Twersky [41], [49] to model ultrasonic scattering. The structure function was used to model the differential cross section per unit volume for a random distribution of identical scatterers [41] and for a mixture of similarly shaped but differently sized particles [49]. In the field of QUS techniques for tissue characterization, Franceschini et al. [15] recommended a method to model the scattering from densely packed cells in tumors using BSC models that take into account the structure function. They performed experiments on concentrated tissue-mimicking phantoms and showed the superiority of the BSC models that take into account the structure function in comparison with other classical BSC models that do not account for the structure function. Vlad et al. [33] performed two-dimensional simulations to study the difference in the backscattering coefficient between the particle distribution with uniform and heterogeneous sizes. They also made a comparison with the Percus-Yevick packing factor – the low-frequency limit of the PY structure function. In [33] and another study [50], particle size variance
was shown to be affecting the structure function and BSC behavior in the case of highly concentrated scattering medium.

Although structure function has seen many applications in ultrasonic scattering, none of the available structure function models in the field of medical ultrasound work well for high-concentration cell pellet biophantoms. Most models were developed for specific conditions and have limitations. For example, the PY structure function starts to show a sharp peak (Figure 2.4) when the concentration becomes relatively high (volume fraction > 50%). Such a sharp peak is not physically realistic. The NS model deals with clustering and does not take into account the concentration issue. Therefore, a new structure function model is needed.

To best model the structure function and demonstrate its effect, it is desirable to separate the effect of the form factor. Therefore, instead of modeling the BSC that is affected by both the structure function and the form factor, we focus only on modeling the structure function in this chapter. Specifically, we will develop analytical structure function models for randomly distributed scatterers that are polydisperse in size, and evaluate the models against the cell pellet biophantoms both forwardly and inversely. More specifically, we will estimate the BSCs of cell pellet biophantoms at two concentrations, a very low concentration where the cells are randomly distributed, and a very high concentration where the cells are closely packed to mimic the condition of a tumor. The structure function that is related to the spatial distribution of cell positions will be isolated by comparing the BSCs of the two concentrations. The
theoretical structure function will be calculated from three models and compared to the experimentally estimated values. The inverse problem will also be explored to generate cell size estimates. Three distinct cell lines will be studied to demonstrate repeatability.

A number of advantages exist in the models developed and the approach used in this chapter: (1) By studying only the structure function rather than the BSC, the effect of spatial scatterer position correlation on scattering is separated and thus can be better studied. (2) The developed structure function models will be able to elucidate the effects of scatterer size distribution on scattering. For instance, the polydisperse structure function models suggest that the size distribution could affect the BSC by affecting not only the incoherent scattering component (which is well established), but also the structure function. (3) The models have minimal dependence on the form factor, which is a significant advantage because identifying the best form factor for a tissue is challenging. (4) The models are in analytical forms and have a limited number of variables, which makes the inverse problem easier to solve. (5) The study is strengthened by evaluating the models in a forward manner using high-frequency (the center of the frequency band is around $ka = 2$) experimental data from biophantoms that mimics tumors.

3.2 Polydisperse Structure Functions

We will start with the PY structure function model introduced in Chapter 2. The sharp peak (Figure 2.4) occurs in the structure function curve at high
concentration because the scatterers were assumed to be identical spheres when Equations (2.16) – (2.18) were derived. Intuitively, identical spheres could lead to a spatial arrangement that is more periodic than polydisperse spheres would under the high-concentration situation. Taking into account the polydispersity in sphere size could reduce the sharp peak in the PY structure function. Two polydisperse structure function models are developed to achieve this goal. For comparison purposes, the PY structure function introduced in Chapter 2 is also called the Monodisperse Model in this chapter, and will be compared to the polydisperse models.

3.2.1 Polydisperse Model I

In this model, the scatterers are assumed to be non-overlapping spheres that are polydisperse in size but monodispersive in scattering amplitudes $\Phi_j(K)$. Note that the assumption of monodispersive scattering amplitude is unrealistic if the system is polydisperse in size, because the scattering amplitude is a function of scatterer size. We make the monodispersive scattering amplitude assumption simply as a mathematical approximation such that Equation (2.15) will still hold, and the structure function will be determined solely by the pair-correlation function. As such, the structure function may be written in terms of partial structure functions $H_y(2k)$ as given by Blum and Stell [51] using PY closure as in [52]

$$S(2k) = 1 + \bar{n} \int_0^\infty \int_0^\infty H_y(2k)f(x_i)f(x_j)dx_i dx_j ,$$

(3.1)

where $f(x)$ is the probability density function of the sphere radius, $x$. 

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The structure function has an analytical expression following Equation (3.1) if
the sphere size follows a $\Gamma$ (Schulz) distribution with a probability density function [52]

$$f_z(x) = \frac{1}{z!} \left( \frac{z+1}{a} \right)^{z+1} x^z e^{-\frac{(z+1)x}{a}}, \quad z = 0, 1, 2, \ldots,$$

(3.2)

where $a$ is the mean of the radius, and $z$ is the Schulz width factor which measures the width of the distribution (a greater $z$ representing a narrower distribution). The $\Gamma$ distribution has been widely used to model polydisperse biological systems, and is an ideal distribution to use for cells.

The analytical expression of the structure function for Polydisperse Model I is listed in Appendix A. The structure function is expressed as a function of mean sphere radius $a$, Schulz width factor $z$, wave number $k$, and sphere volume fraction $\eta$. The structure functions at various Schulz width factors are shown in Figure 3.1. When $z \rightarrow \infty$, the Polydisperse Model I yields the same result as that of Monodisperse Model, which could serve as a code sanity check. As the polydispersity of sphere radius increases (i.e., $z$ decreases), the peak of the structure function curve reduces accordingly.

3.2.2 Polydisperse Model II

Polydisperse Model I does show decreased peak values compared to the Monodisperse Model for high concentrations as expected. However, an unrealistic assumption was made during the development of Polydisperse Model I: the scatterers were assumed to be polydisperse in size but monodisperse in scattering amplitudes $\Phi_j(K)$. This assumption allows for convenient mathematical
approximation to be made. To investigate if the assumption is reasonable, Polydisperse Model II is developed and compared to Polydisperse Model I.

For Polydisperse Model II, the scatterers are assumed to be non-overlapping spheres that are polydisperse in both size and scattering amplitude, and the sphere size is assumed to follow a $\Gamma$ distribution. As a result of the polydispersity in scattering amplitude, the scattering amplitude cannot be factored out in Equation (2.2). Therefore, Equations (2.13) and (2.15) are not valid any more. To derive the structure function expression for this case, we express the BSC as

$$BSC(2k) = \bar{n} \int_{0}^{\infty} |\Phi_j(2k)|^2 f(x_j)dx_j$$
$$+ \bar{n} \int_{0}^{\infty} \Phi_j(2k)\Phi_j(2k)H_y(2k)f(x_j)f(x_j)dx_jdx_j$$

Equation (3.3) is a modification of Equation (1) of [53]. Similar expressions may also be found in [49]. The first integral $\bar{n} \int_{0}^{\infty} |\Phi_j(2k)|^2 f(x_j)dx_j$ of Equation (3.3) represents the quantity $\text{BSC}_{\text{incoherent}}(2k)$. The second integral of Equation (3.3) represents the excess scattering caused by the spatial correlation in scatterer positions. Substituting Equation (3.3) into Equation (2.11) yields the structure function for Polydisperse Model II:

$$S(2k) = 1 + \frac{\int_{0}^{\infty} \int_{0}^{\infty} \Phi_j(2k)\Phi_j(2k)H_y(2k)f(x_j)f(x_j)dx_jdx_j}{\int_{0}^{\infty} |\Phi_j(2k)|^2 f(x_j)dx_j} .$$

(3.4)

This structure function is dependent on the scattering amplitude $\Phi_j(2k)$. Therefore, a specific form of scattering amplitude is needed to evaluate Equation (3.4). The scattering amplitude that is used herein is derived from the fluid-filled
sphere form factor in Equation (2.7) for which the integrals in Equation (3.4) have analytical expressions. The resulting expression for the structure function is listed in Appendix B. The structure function is expressed as a function of the mean sphere radius $a$, Schulz width factor $z$, wave number $k$, and sphere volume fraction $\eta$. The structure functions for Polydisperse Model I and Polydisperse Model II are compared at various degrees of polydispersity (Figure 3.2). The peak at around $ka = 2$ in the structure function curve is lower for Polydisperse Model II than for Polydisperse Model I when the scatterers are polydisperse (Figure 3.2(a)-(c)), suggesting that the monodisperse scattering amplitude assumption made for Monodisperse Model I does affect the structure function result. However, the discrepancy between the two polydisperse models tends to be reduced as the polydispersity is reduced. As a code sanity check, Polydisperse Model I and Polydisperse Model II generate identical results when the scatterers are essentially monodisperse (Figure 3.2(d)).

3.3 Experimental and Data Reduction Methods

3.3.1 Biophantom construction

The cell pellet biophantoms were composed of a known number of cells clotted in a mixture of bovine plasma (Sigma-Aldrich, St. Louis, MO) and bovine thrombin (Sigma-Aldrich, St. Louis, MO). Three cell lines, Chinese hamster ovary [CHO, American Type Culture Collection (ATCC) #CCL-61, Manassas, VA], 13762 MAT B III (MAT, ATCC #CRL-1666) and 4T1 (ATCC #CRL-2539), were used to create the cell pellet biophantoms. The three cell lines were chosen because: (1) they represent normal and tumor cell lines (normal cells:
CHO, tumor cells: MAT and 4T1), and (2) they represent different cell sizes (see Figure 3.3 for measured cell radius histograms and the corresponding Schulz distribution fit). Two cell concentrations (Table 3.1) were realized for each cell line, with each concentration having two to three independent replicates of biophantoms.

The detailed procedure of constructing cell pellet biophantoms is as follows. The cells were cultured in an ATCC-recommended medium along with 8.98% of fetal bovine or calf serum (Hyclone Laboratories, Logan, UT) and 1.26% of antibiotic (Hyclone Laboratories, Logan, UT). A Reichert Bright-Line® hemacytometer (Hausser Scientific, Buffalo, NY) was used to count viable cells to yield the number of cells per known volume. Equal volumes of the dye Trypan Blue (Hyclone Laboratories, Logan, UT) and cell suspension were gently mixed by pipetting and then added to the counting chambers of the hemacytometer. Trypan Blue was used to differentiate nonviable cells (stained as blue cells) from viable cells (displayed as bright cells). At this point, the cells had an average of over 90% viability. A known number of viable cells was placed in a 50-mL conical centrifuge tube (Corning® Incorporated, Corning, NY), and spun in a 4 °C centrifuge at 2500 rpm for 10 min, and the supernatant was removed. Then, 90 µL of bovine plasma were added to the cell sediment in the centrifuge tube, which was then vortexed. Next, 60 µL of bovine thrombin were added, and the mixture was lightly agitated to coagulate and form a biophantom. The biophantom was transferred onto a planar Plexiglas® plate, and submerged in Dulbecco’s
Phosphate Buffered Saline (DPBS) (Sigma-Aldrich, St. Louis, MO) for ultrasonic scanning.

3.3.2 Experimental setup and BSC estimation method

The biophantoms were ultrasonically scanned using three single-element, weakly focused transducers (20-MHz transducer IS2002HR, from Valpey Fisher Cooperation, Hopkinton, MA; 40- and 80-MHz transducers from NIH High-frequency Transducer Resource Center, University of Southern California, Los Angeles, CA; see Table 3.2). The total frequency range covered was 11 to 105 MHz.

The transducers were interfaced with a UTEX UT340 pulser/receiver (UTEX Scientific Instruments Inc., Mississauga, ON, Canada) that operated in the pitch-catch mode. A 50DR-001 BNC attenuator (JFW Industries Inc., Indianapolis, IN) was connected to the pulser to attenuate the driving pulse to avoid transducer saturation. An RDX-6 diplexer (Ritec Inc., Warwick, RI) was used to separate the transmitted and received signals because only the transmitted signal needed to be attenuated. The received RF signals were acquired using a 10-bit Agilent U1065A-002 A/D card (Agilent Technologies, Santa Clara, CA) set to sample at 1 GHz. The transducers were moved using a precision motions control system (Daedal Parker Hannifin Corporation, Irwin, PA) that has a linear spatial accuracy of 1 µm. The biophantoms were placed on the Plexiglas® plate during ultrasound scans. The scans were performed in a small tank filled with DPBS at room temperature (Figure 3.4).
Attenuation and BSC measurements were performed for each sample. The attenuation was determined to allow for attenuation compensation during the BSC estimation process. An insertion-loss broadband technique [54] was used to estimate the attenuation. The insertion loss was determined by comparing the power spectra of the echoes reflected off the top surface of the Plexiglas® with and without the sample being inserted in the ultrasound path. The transducer focus was positioned at the Plexiglas® surface when the signal was being recorded. The effect of DPBS attenuation was compensated for when the biophantom attenuation was estimated from the insertion loss. The attenuation (dB/cm) of a sample was generated by averaging the attenuation obtained from 36 independent locations laterally across the sample.

The BSC scanning procedure started with acquiring the reference signals from the DPBS-Plexiglas® interface whose pressure reflection coefficient at room temperature is known (= 0.37). The reference signals were acquired at the set of axial positions that covered the –6 dB depth of focus with a step size of a half wavelength. Next, a raster scan on the biophantom sample was performed with a lateral step size of one beam width. The transducer focus was positioned in the sample during the scan. The scan covered a sufficient length both axially and laterally to make sure that a sufficient number of regions of interest (ROIs) could be acquired and processed. Eleven equally spaced slices were imaged for each sample, and the number of A-lines per sample varied depending on the transducer frequency and sample size. The BSC was estimated from the RF data using a planar reference method [55] to remove equipment-dependent effects. To generate
a BSC versus frequency curve for a sample scanned by a single transducer, (i) a BSC estimate was made for each ROI based on the gated RF echo data from that ROI, (ii) a mean BSC was estimated for each of the 11 slices by averaging the BSCs from all the ROIs within that slice, and (iii) the 11 mean BSCs were averaged.

3.3.3 Histology processing

Immediately after scanning, the sample was placed into a histology processing cassette and fixed by immersion in 10% neutral-buffered formalin (pH 7.2) for a minimum of 12 h for histopathologic processing. The sample was then embedded in paraffin, sectioned, mounted on glass slides and stained with hematoxylin and eosin (H&E) for further evaluation by light microscopy (Olympus BX–51, Optical Analysis Corporation, Nashua, NH, USA).

3.3.4 B-spline fit and structure function estimation

Two concentrations (Table 3.1) were studied for each cell line: the higher concentration was chosen to be as high as possible to mimic the cell concentration in tumors, and the lower concentration was chosen to be sufficiently low such that the structure function can be assumed to be unity, while still high enough to ensure sufficient signal-to-noise ratio in the backscatter data. Based on these conditions, the structure function for the higher concentration may be obtained experimentally by

\[
S(f) = \frac{\tilde{n}_H BSC_H(f)}{\tilde{n}_H BSC_L(f)}, \quad (3.5)
\]
where $\bar{n}_l$ and $\bar{n}_h$ represent the number density for the lower and the higher concentrations, respectively, $BSC_L(f)$ and $BSC_H(f)$ represent the BSC for the lower and the higher concentrations, respectively.

There were a number of BSC versus frequency curves obtained from multiple transducers and multiple realizations for each concentration of each cell line. A B-spline fit was performed on these curves to generate a single fitted curve that covered the entire frequency range (11 – 105 MHz) for a concentration of a cell line. The fitted BSC values were used for structure function estimation using Equation (3.5).

The B-spline is a commonly used smoothing spline for large data sets. The advantage of a smoothing spline is that the resulting curve is not required to pass through each data point. The resulting B-spline curve is a linear combination of $M$ B-spline basis functions, where $M$ is the degrees of freedom, and the B-spline basis functions are spaced at different locations to provide local shape control. In this dissertation, we fit cubic B-splines with five degrees of freedom, giving us five B-spline basis curves at five equally spaced locations in the frequency range. The best-fit B-spline is then a linear combination of five B-spline basis functions:

$$bs(f) = \sum_{i=1}^{5} \beta_i b_i(f),$$  \hspace{1cm} (3.6)

where $b_i(f)$ is the $i$th B-spline basis function, and $\beta_i$ is the corresponding coefficient of each basis function to control the shape locally. The calculation of

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and the least square estimation of $\beta_i$ are performed using custom programs developed in MATLAB® (The Mathworks Inc., Natick, MA).

3.4 Results and Discussions

3.4.1 BSC estimates and B-spline fit

The attenuation-compensated BSC estimates for the biophantoms are shown in Figure 3.5 for each cell line. The BSC curves of all realizations were plotted to show the degree of measurement uncertainty and/or the uncertainty in concentration control. Overall, multiple realizations had consistent BSC results.

The BSC behaviors in Figure 3.5 reveal significant information about the structure function. The BSC shape is significantly different between the lower and higher concentrations for all three cell lines. This observation confirms that it is necessary to consider the structure function for the higher concentration condition. The BSC magnitude appears to be similar between the lower and higher concentrations at lower frequencies ($f < 30$ MHz), whereas the difference in BSC magnitudes of the two concentrations start to increase at higher frequencies ($f \sim 60$ MHz). A physical interpretation of this behavior is that the effect of cell position correlation on scattering for the high-concentration case is destructive at frequencies lower than 30 MHz, and is constructive (or less destructive) at around 60 MHz. This interpretation is consistent with the shape of the theoretical structure functions presented in Figures 2.4, 3.1, and 3.2: the structure functions are lower than unity at lower $ka$ values, and are peaking at around $ka = 2$. Furthermore, a peak at around 60 MHz, and a dip at around 90
MHz are observed for every BSC curve. The peak and dip behavior for the lower-concentration case is explained by form factors (e.g., fluid-filled sphere, concentric spheres) that match the geometry and acoustic impedance distribution of individual cells. The peak for the higher concentration is sharper compared with the lower concentration. None of the commonly used form factors for soft materials could yield such a sharp peak, indicating that other factors such as the structure function might contribute to the sharp peak.

3.4.2 Experimental and theoretical structure functions

The experimental structure function (Figure 3.6) for Concentration 2 (see Table 3.1) was determined using Equation (3.5) assuming the structure function was unity for Concentration 1 as discussed in details in Section 3.4.5. The theoretical structure functions (including the two polydisperse models developed in this chapter and the Monodisperse Model (PY) reviewed in Chapter 2, Figure 3.6) for Concentration 2 were calculated using the three structure function models. For the theoretical calculation, the volume fraction was assumed to be 74% for Concentration 2. The values of parameters $a$ and $z$ used for theoretical calculation were the same as the Schulz distribution fit results presented in Figure 3.3. To convert from $k$ to $f$, a propagation speed of 1540 m/s was assumed throughout this chapter.

Figure 3.6 shows that the theoretical structure functions from the three models have a peak-dip pattern consistent with that of the experimental structure function. The positions of the peaks and dips are well aligned between the theoretical and experimental curves. However, the exact magnitude of the major peak varies.
between the theoretical and experimental curves. The Monodisperse Model shows the highest peak, which extends well above 7 and is clipped in Figure 3.6. Polydisperse Model I shows a lower peak, and Polydisperse Model II shows the lowest peak among all three models. Relative to the magnitude of the peak, Polydisperse Model II has the best agreement to the experimental curve, and therefore seems to be the most accurate model out of the three.

Although the theoretical curves of the two polydisperse models show agreement with the experimental curves, the agreement is not perfect. A perfect agreement is not expected, because the scattering of cells is so complex that many factors could contribute to scattering. The structure function only models one factor, the spatial correlation of cell positions, and shows that this factor is important. Other factors, such as multiple scattering, might explain in general why the polydisperse models do not perfectly agree with experimental data. That being said, we try herein to explain, within the framework of structure function, a number of observed discrepancies between the models and the experimental data. The polydisperse models seem to work better for CHO and MAT than for 4T1. This observation might be attributed to the fact that 4T1 has the highest degree of polydispersity among all the three cell lines. A higher degree of polydispersity leads to a smoother peak in the structure function. A smooth peak is easier to be shifted due to measurement errors than a sharp peak. Another noticeable difference between the theoretical and experimental curves is that the peak of the theoretical curves is higher than that of the experimental curves. There could be a number of explanations for this difference. We might have underestimated the
polydispersity of cells. We have considered only the polydispersity in cell size, but not the polydispersity in cell shape. Experimental errors might as well contribute to the difference. For instance, if the attenuation was underestimated for the higher concentration, then the BSC might be underestimated consequently, resulting in an underestimated experimental structure function. Also, the volume fraction of 74% might have uncertainty. If the actual volume fraction was slightly deviated from 74%, then the theoretical structure function in Figure 3.6 would be slightly different as well.

3.4.3 Inverse problem

The usefulness of Polydisperse Models I and II is demonstrated herein via solving the inverse problem: estimating the mean radius from experimental structure functions.

The mean radius $a$ and the Schulz width factor $z$ were the unknowns in the inverse problem. The volume fraction was assumed to be known $a$ priori ($\eta = 74\%$). The two unknowns were estimated by fitting the theoretical structure function $SF_{\text{theo}}(f)$ to the experimental structure function $SF_{\text{exp}}(f)$. Specifically, we perform an exhaustive search procedure for values of $(a, z) \in [4 \, \mu m, 12 \, \mu m] \times [5, 100]$ to minimize the cost function:

$$C(a, z) = \sum_i \| SF_{\text{theo}}(f_i) - SF_{\text{exp}}(f_i) \|^2$$

over the frequency range 11–105 MHz.
The results of the search show that a unique global minimum always exists for the CHO, MAT, and 4T1 cell pellets for Polydisperse Model I and Polydisperse Model II. A typical logarithm of the cost function $C(a,z)$ is shown in Figure 3.7(a). The mean radius estimates and the Schulz width factor estimates are shown in Figure 3.7(b) and Figure 3.7(c), respectively. Overall Polydisperse Models I and II yield relatively accurate mean radius estimates, with a maximum percentage error of 13.5% for Polydisperse Model I and 6.7% for Polydisperse Model II for all three cell lines evaluated. As expected, Polydisperse Model II provides slightly better size estimates than does Polydisperse Model I. The Schulz width factor estimates are not as accurate as the mean radius estimates. Both polydisperse models underestimate the Schulz width factor, i.e., overestimate the degree of polydispersity in cell size, possibly because the polydispersity in cell shape might also contribute to scattering and could decrease the estimated Schulz width factor value. It is not surprising that Polydisperse Model II yields a better Schulz width factor estimate than does Polydisperse Model I, because Polydisperse Model II takes into account the polydispersity in scattering amplitude to some extent, whereas Polydisperse Model I does not. Although the Schulz width factor $z$ is underestimated by the models, the estimated $z$ values are accurate in relative terms: 4T1 has the lowest $z$ values, both measured and estimated, and MAT has the highest $z$ values, both measured and estimated.

The fitted structure function curves (Figure 3.8) show good agreement with the experimental curves in terms of peak positions. This observation is consistent with the relatively good accuracy in size estimates, because the position of the
peak is mainly determined by the cell size. Polydisperse Model II appears to have better-fitted curves than Polydisperse Model I in terms of agreement in the peak magnitude (Figure 3.8). This observation might explain why Polydisperse Model II has better Schulz width factor estimates, because the peak magnitude is presumably related to the Schulz width factor more than to the mean radius.

3.4.4 Comparison to Gaussian and fluid-filled sphere BSC models

To test whether fitting the structure function curves could yield better mean cell radius than fitting BSC curves, we fit two commonly used BSC models, the spherical Gaussian (Equations (2.6) and (2.9)) and the fluid-filled sphere model (Equations (2.6) and (2.7)), to the high-concentration BSC curves presented in Figure 3.5. Both BSC models take into account only the geometry and acoustic impedance profile of the cells, but not the spatial correlation of cell positions. The detailed estimation procedure can be found in [18]. The estimated effective scatterer radius from the two BSC models is compared with the estimated mean cell radius from the two polydisperse structure function models (Figure 3.9). The two polydisperse structure function models show advantage in terms of estimating the cell radius. They yield relatively accurate mean cell radius estimates, whereas the two BSC models do not. One might argue that the effective scatterer size estimates from the two BSC models might correspond to the size of cell nucleus. In fact, this argument pointed out a significant disadvantage of the two BSC models: it is difficult to relate the effective scatterer size estimates to real tissue anatomy. It is not clear if the effective scatterer size estimates relate to the cell radius, the nucleus radius, or anything else. This ambiguity does not exist in the
polydisperse structure function models. The estimated mean scatterer radius from the polydisperse structure function models can only be related to the cell radius, because the models describe the spatial correlation of scatterer positions which is affected by the cells as opposed to the nuclei.

3.4.5 The theoretical structure function for Concentration 1

A basic assumption for the experimental structure function curves presented in Figure 3.6 is that the structure function is unity for Concentration 1. This subsection investigates if the assumption is reasonable.

We start with calculating the theoretical structure function curves for Concentration 1 predicted by Monodisperse Model, Polydisperse Model I, and Polydisperse Model II (Figure 3.10), and compare them to unity. The volume fraction values in Table 3.1 (2.7% for CHO, 3.4% for MAT, and 6.1% for 4T1) and the size distribution parameters in Figure 3.3 are used for theoretical structure function calculation. At frequencies above 40 MHz, Figure 3.10 shows no noticeable difference between unity and the theoretical structure function curves for Concentration 1. Slight (compared with Concentration 2) but noticeable difference appears at the lower frequency end. Overall, the unity assumption of structure function for Concentration 1 appears to be reasonable, which may be further demonstrated by comparing the difference between size parameters estimated with and without the unity assumption.

Size parameters were estimated in Section 3.4.3 by fitting the theoretical structure function curves to the experimental curves, under the unity assumption. If the unity assumption does not hold, then the accurate way of fitting the data to
an equation would be to fit the ratio of the theoretical structure function of Concentration 2 to Concentration 1, to the “experimental” structure function curves presented in Figure 3.6. Therefore, Equation (3.7) should be revised as

\[
C(a, z) = \sum_i \frac{SF_{\text{theo,conc2}}(f_i)}{SF_{\text{theo,conc1}}(f_i)} - SF_{\text{exp}}(f_i) \right|^2,
\] (3.8)

where the subscripts conc1 and conc2 represent Concentration 1 and Concentration 2, respectively. The cell size parameters \((a\) and \(z\)) estimated using this approach (Equation 3.8) appear to be sufficiently close to those estimated using the unity assumption (Table 3.3), suggesting that the unity assumption is reasonable.

The above analysis also suggests that the frequency matters for determining at what concentration levels the structure function can be assumed to be unity. A previous study [48] at lower frequencies \((ka < 0.5)\) showed that the structure function cannot be assumed to be unity for concentrations greater than 2.5%. The frequency range of the study in this chapter extends up to \(ka = 3\). For such a broad frequency range, a volume fraction of around 6.1% still seems to be sufficiently low for assuming a unity structure function. For high concentrations such as 74% volume fraction, however, the structure function is significant for both lower and higher frequencies.

3.4.6 Theoretical implications of the polydisperse structure function models

The central problem we are trying to address is to elucidate the mechanism(s) of ultrasonic scattering at high concentrations and to model the scattering. The
results seem to support the hypothesis that the scattering at high concentrations is determined by both the scattering from individual scatterers \[ \Phi_i(2k) \] and the correlation of scatterer positions (the structure function). Without taking into account the contributions from the correlation of scatterer positions, it proves extremely difficult to interpret the BSC data for high-concentration media. The sharp peak in the BSC of high-concentration biophantoms (Figure 3.5) cannot be explained alone by the fluid-filled sphere model or the more complex concentric spheres model that has worked well for the low-concentration case. Neither the fluid-filled sphere model nor the concentric spheres model provides a satisfactory fit to the high-concentration BSC data. Force-fitting those models to the data does not yield reasonable size estimates either. On the other hand, the structure function models could explain the data better and yields relatively accurate size estimates. This fact indicates the important role of correlation of scatterer positions on scattering.

The models also demonstrate the significant role of polydispersity on structure function. The Monodisperse Model, which does not take into account the polydispersity of scatterers, does not fit the data very well, although it could qualitatively explain the peaks in the experimental structure functions. Two types of treatment to the polydispersity issue have been considered: Polydisperse Model I assumes polydispersity in scatterer size and monodispersity in individual scattering amplitudes \( \Phi_i(2k) \), whereas Polydisperse Model II assumes polydispersity in both scatterer sizes and scattering amplitudes. Both polydisperse
models show acceptable results, with Polydisperse Model II having slightly better performance than Polydisperse Model I.

3.4.7 Practical usefulness of the polydisperse structure function models

The structure function provides additional new information about tissue structure, independent of the information provided by BSC and attenuation. From the structure function, we may be able to estimate tissue properties such as the mean scatterer radius and the Schulz width factor. In the future, parameters such as the Schulz width factor could potentially be explored for tissue characterization using high-frequency ultrasound. For instance, the Schulz width factor could be used for detecting cell death, as an earlier work [33] has shown that the cellular size variance increases after cell death.

The structure functions expressed in the models have a limited number of parameters. The structure function is not dependent on the acoustic property contrast between the scatterer and the background. For instance, the acoustic impedance contrast between the scatterer and the background is not affecting the structure function. Neither is mass density an issue to be considered in the models. The limited number of parameters makes it more likely to find a unique global minimum in the inverse problem.

The experimental setup in this chapter requires the measurement of biophantoms of two concentrations, with the lower concentration serving the reference purpose. This was designed primarily for dissecting the BSC, demonstrating the role of scattering position correlation on scattering, and quantifying that role in terms of structure function. It is difficult to directly
implement this setup in clinical settings. However, with the models developed through this setup, we are progressing toward accurately modeling the BSC from high-concentration scattering media by combining the structure functions with appropriate form factors. If that were successful, then the requirement of measuring a low-concentration biophantom as a reference would be eliminated.

3.5 Chapter Summary

The correlation of scatterer positions has significant contributions to the scattering of dense media. This contribution could be modeled by the structure functions. The polydispersity of the scatterer size has a significant effect on the structure functions, and should be taken into account in structure function models. The two polydisperse structure function models developed in this chapter could lead to improved modeling of scattering from dense media.
3.6 Figures

Figure 3.1 Comparison among structure functions at five different Schulz width factor values: 5, 10, 50, 100, and $10^5$, computed by Polydisperse Model I. The structure function at $z = 10^5$ computed by Polydisperse Model I (solid line) is identical to that computed by Monodisperse Model (circles). The volume fraction is assumed to be 74% for all the curves.
Figure 3.2 Comparison between structure functions of Polydisperse Model I (dotted line) and Polydisperse Model II (solid line) at (a) $z = 10$, (b) $z = 50$, (c) $z = 100$, and (d) $z = 10^7$. The volume fraction is assumed to be 74% for all the curves.
Figure 3.3 Measured cell radius distribution and Schulz distribution fit for (a) CHO cells, (b) MAT cells, and (c) 4T1 cells. The size of live cells was measured by light microscope using a procedure detailed in [36]. The normalized histograms were generated from results of 500 CHO, 200 MAT, and 200 4T1 cell size measurements. The mean radius is 6.7, 7.3, and 8.9 µm for CHO, MAT, and 4T1 cells, respectively. The fitted Schulz width factor $z$ is 51.9, 65.8, and 31.9 for CHO, MAT, and 4T1 cells, respectively.
Figure 3.4 Diagram of the experimental setup for attenuation and BSC measurements.
Figure 3.5 BSC versus frequency for (a) CHO, (b) MAT, and (c) 4T1 cell pellet biophantoms measured using three transducers with center frequencies at 20, 40, and 80 MHz, respectively. The B-spline curve is also displayed for each concentration.
Figure 3.6 Comparison between experimental structure functions (solid gray lines) and theoretical structure functions (dotted lines: Monodisperse Model, dashed lines: Polydisperse Model I, solid dark lines: Polydisperse Model II) for high-concentration (a) CHO, (b) MAT, and (c) 4T1 cell pellet biophantoms.
Figure 3.7 (a) A representative 2D cost function map: the logarithm of the cost function $C(a, z)$ for Polydisperse Model II obtained from experimental structure function for CHO. (b) A comparison between the mean radii estimated by Polydisperse Models I and II and direct light microscope measures. (c) A comparison between the estimated and directly measured parameter $z$. The legends in (c) are the same as in (b).
Figure 3.8 A comparison between experimental and best-fit structure function curves for (a) CHO, (b) MAT, and (c) 4T1.
Figure 3.9 A comparison between the estimated effective scatterer radius from two BSC models (the spherical Gaussian and the fluid-filled sphere model) and the estimated mean cell radius from the two polydisperse structure function models.
Figure 3.10 Theoretical structure function curves for Concentration 1 predicted by (a-b) Monodisperse Model, (c) Polydisperse Model I, and (d) Polydisperse Model II, along with the curves for Concentration 2 for comparison. (b) is a zoomed-in version of (a).
### 3.7 Tables

Table 3.1 Summary of the cell concentrations for cell pellet biophantoms of the three cell lines. The cell concentration is represented by number density in million cells/mL (Mcell/mL) and volume fraction (%).

| Cell line | Concentration 1 | | Concentration 2 | | |
|-----------|-----------------|---|-----------------|---|
|           | Number density  | Volume fraction | Number density  | Volume fraction |
|           | (Mcell/mL)      | (%)           | (Mcell/mL)      | (%)           |
| CHO       | 20              | 2.7           | 556             | 74            |
| MAT       | 20              | 3.4           | 442             | 74            |
| 4T1       | 20              | 6.1           | 244             | 74            |

Table 3.2 Transducer information and characteristics.

<table>
<thead>
<tr>
<th>Center frequency (MHz)</th>
<th>–10 dB bandwidth (MHz)</th>
<th>Wavelength at center frequency (µm)</th>
<th>f-number</th>
<th>–6 dB depth of field (mm)</th>
<th>–6 dB beam width (µm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>11–33</td>
<td>75.0</td>
<td>3.0</td>
<td>4.0</td>
<td>230</td>
</tr>
<tr>
<td>40</td>
<td>26–65</td>
<td>37.5</td>
<td>3.0</td>
<td>2.4</td>
<td>113</td>
</tr>
<tr>
<td>80</td>
<td>49–105</td>
<td>18.8</td>
<td>3.0</td>
<td>1.2</td>
<td>56.4</td>
</tr>
</tbody>
</table>
Table 3.3 Comparison between cell size parameters estimated with and without the assumption that the structure function curve for Concentration 1 is unity. The numbers in parentheses represents estimated values without the unity assumption.

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Estimated mean radius $a$ (µm)</th>
<th>Estimated Schulz width factor $z$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Polydisperse Model I</td>
<td>Polydisperse Model II</td>
</tr>
<tr>
<td>CHO</td>
<td>6.4 (6.4)</td>
<td>7.0 (7.0)</td>
</tr>
<tr>
<td>MAT</td>
<td>6.6 (6.7)</td>
<td>7.1 (7.1)</td>
</tr>
<tr>
<td>4T1</td>
<td>7.7 (7.9)</td>
<td>8.3 (8.3)</td>
</tr>
</tbody>
</table>
This chapter presents high-frequency BSC results for various tumor types to provide a database for understanding the scattering from tumors and modeling tumor structures. Complementing the BSC data from the tumors is high-concentration cell pellets of the same cell lines when available. Additionally, the high-frequency attenuation results are also presented. The comparison between the tumor and cell pellet BSC for the same cell line will provide insight into understanding the scattering from tumors.

4.1 Introduction

Chapter 3 has discussed modeling the scattering of high-concentration cell pellet biophantoms. To progress from moderate complexity to significant complexity, this chapter deals with real tumors. To take advantage of the modeling results presented in Chapter 3, a comparison between the cell pellet and tumor of the same cell line is highly desirable. Such a comparison will help understanding the scattering mechanism(s) in tumors.

The concentration of the biophantoms is chosen to be the highest to mimic the tumor: the biophantoms are made of densely packed cells without supportive materials. The dense cell pellet has actually been used as a model of real tissues/tumors for high-frequency QUS studies due to its simplicity of preparation and ease of implementation [19], [56], [57]. However, the cell pellet and tumors are different in many ways, which could make their ultrasound backscatter differ from each other. The tumors have more complex components and structures than cell pellets. There are cells, complex extracellular matrix and vasculatures in a tumor, whereas there are only cells in a cell pellet. The scattering from the tumor will be even more complex if there are regions of necrosis or apoptosis.

In addition to the comparison between cell pellets and tumors of the same cell type, the comparison among different tumor types will also reveal useful information on how different tumor structures may scatter ultrasound differently. For this reason, we have scanned several different tumor types. Doing so also has other benefits. It will enrich the database for high-frequency tumor scattering data to the scientific community. It will also provide an opportunity to demonstrate the powerfulness, as well as potential limitations, of QUS techniques for tumor classification.

The results of five tumor types will be presented in this chapter: spontaneous fibroadenoma, carcinoma (4T1, ATCC #CRL-2539), adenocarcinoma (MAT [13762 MAT B III], ATCC #CRL-1666), malignant fibroblast sarcoma (LMTK, ATCC #CCL-1.3), and sarcoma (EHS [Englebreth-Holm-Swarm], ATCC #CRL-2108). The last four are injected tumors. Three tumor types (MAT, 4T1, LMTK)
have tumor cells that can be used to construct a cell pellet. Each of the five tumor types may have some unique structures. The spontaneous fibroadenoma consists of a large amount of connective tissues. The 4T1, MAT, and LMTK tumors have a homogenous distribution of tumor cells, with minimal extracellular matrix. However, their cell radii are slightly different; their growth rates are different; and each may have its unique structure as well. The EHS tumor is known to have a heterogeneous distribution of tumor cells, with abundant extracellular materials. The EHS cells are arranged in groups usually containing less than 20 cells per group.

With abundant available data, this chapter will present (1) the comparison between 4T1 tumors and cell pellets, MAT tumors and cell pellets, and LMTK tumors and cell pellets, respectively, in terms attenuation and BSC curves; (2) the EHS tumor results in comparison to 4T1; and (3) the fibroadenoma results. The reason for including attenuation data in comparison is that attenuation is related to the macro-molecular components of the media, and comparing attenuation can help understanding the tissue components.

4.2 Materials and Methods

4.2.1 Cell pellet biophantom construction

The cell pellet biophantoms used in this chapter were composed of densely packed cells without any supportive background materials. Three tumor cell lines, the MAT, 4T1, and LMTK, were used to create the cell pellets, each cell line having three to five independent replicates of cell pellets.
The cell culture and harvest procedures were the same as described in Section 3.3.1. The cell pellet biophantom construction procedure was slightly different. A known number of viable cells was placed in a 50-mL conical centrifuge tube and spun in a 4 °C centrifuge at 2500 rpm for ten minutes, and the supernatant was removed. Thereafter, the 50-mL conical tube was centrifuged again at 4 °C and 2500 rpm for three minutes, and the supernatant was removed. A heated surgical blade was used to cut off the top portion of the 50-mL centrifuge tube 1 cm above the remaining cell sediment. A spatula was used to collect the cell sediment and place it on a planar Plexiglas® plate. After a ten minute wait, the cell sediment and Plexiglas® plate were carefully submerged in Dulbecco’s Phosphate Buffered Saline (DPBS) (Sigma-Aldrich®, MO, USA) for ultrasonic scanning.

4.2.2 Animal use, cell injection and tumor sample preparation

The MAT, 4T1, LMTK, and EHS cells were injected into Fischer 344 rats, BALB/c mice, athymic nude mice, and C57BL/6 mice, respectively, to grow into tumors. All animals were purchased from Harlan® Laboratories, Inc. (Indianapolis, IN, USA). Sprague-Dawley rats that had spontaneous fibroadenoma were also purchased (the fibroadenoma has to be confirmed by histological slides after the tumors were scanned). To excise the tumors, the animals were anesthetized via CO₂. The excised tumors were trimmed such that the thickness did not exceed 2 mm. This was done to reduce total attenuation and achieve a good signal-to-noise ratio for the insertion-loss attenuation measures. The excised tumors were placed on a planar Plexiglas® plate. The tumors and the Plexiglas® plate were submerged in DPBS for ultrasonic scanning. Eventually
thirteen MAT, fifteen 4T1, thirteen LMTK, thirteen EHS, and nine fibroadenoma samples were successfully excised, scanned and analyzed.

The experimental protocol was approved by the Institutional Animal Care and Use Committee of the University of Illinois and satisfied all campus and National Institutes of Health rules for the humane use of laboratory animals. Animals were housed in an Association for Assessment and Accreditation of Laboratory Animal Care (Rockville, MD, USA)-approved animal facility and provided food and water *ad libitum*.

4.2.3 Ultrasound scanning procedure, BSC estimation method and histology processing

See Section 3.3.2 for ultrasound scanning procedure, and the attenuation and BSC estimation methods. See Section 3.3.3 for the histology processing method. All the samples were scanned using the 20-, 40-, and 80-MHz transducers (Table 3.2), except that the fibroadenoma samples were scanned using only the 20- and 40-MHz transducers due to the extremely high attenuation of the samples.

4.3 BSC and Attenuation Results and Discussion

4.3.1 Tumors versus cell pellets

*A) BSC results*

The attenuation-compensated BSC estimates from both cell pellets and tumors are shown in Figures 4.1, 4.2, and 4.3 for 4T1, MAT, and LMTK, respectively. The BSC curves are presented in a fashion that the curve from each individual realization of each sample type is shown. This type of presentation allows one to infer how large the measurement uncertainty is. For instance, the five MAT cell
pellet curves in Figure 3.2 appear to have the narrowest distribution. This distribution can serve as an indication of the upper limit of measurement uncertainty. One may also note that the tumors have a much wider distribution in BSC curves than cell pellets. Such a wide distribution in tumors can then be reasonably attributed to large inter-tumor variation rather than measurement uncertainty if we consider the upper limit of measurement uncertainty indicated by the MAT cell pellet data. Overall, it is observed that tumors have a wider BSC distribution than that of the cell pellets for both cell lines, because the cell pellets are well-controlled biophantoms, whereas tumors are less controlled and thus have larger variance in nature. Also shown in the BSC plots are the BSC shape trends for each sample type represented by the B-spline curves.

To compare tumors against cell pellets, Figures 4.1 – 4.3 are examined separately. Figure 4.1 shows that, for the 4T1 cell line, the tumors and cell pellets share similar BSC values over the entire frequency range under investigation except around 65 MHz where the BSC of the cell pellets is slightly higher than that of the tumors. There appears to be a slight but noticeable peak at 65 MHz for the cell pellet BSC, whereas the tumor BSC curves are flatter. The peak in cell pellets has been attributed to the structure function as analyzed in Chapter 3. The fact that the peak does not appear in the tumor curve indicates that the structure function in tumors might be different. For instance, the tumor cell arrangement might happen to be more random than in a cell pellet. Nevertheless, the overall BSC shapes and magnitude are similar for 4T1 cell pellets and tumors. This similarity is not observed for the MAT, or LMTK cell lines, however. Figure 4.2
shows two distinct BSC shape trends for the MAT cell pellets and tumors, and Figure 4.3 shows two distinct BSC shape trends for the LMTK cell pellets and tumors. Before attempting to explain such distinction between MAT cell pellets and tumors, and between LMTK cell pellets and tumors, the attenuation results and histopathologic evaluation of cell pellets and tumors are presented first.

B) Attenuation results

The attenuation (dB/cm) results are presented in Figures 4.4 – 4.6 as a function of frequency. In Figure 4.4, the 4T1 cell pellet attenuation curves are not readily separable from those of the 4T1 tumors. However, the MAT cell pellet attenuation curves are completely separated from the MAT tumor attenuation curves (Figure 4.5). The attenuation curves almost overlap for LMTK cell pellets and tumors (Figure 4.6).

C) Histology

The H&E stained histological slides are shown in Figures 4.7, 4.8, and 4.9 for 4T1, MAT, and LMTK, respectively.

No apparent difference is observed between the histological images of the 4T1 cell pellets and 4T1 tumors (Figure 4.7). The microscopic features of the 4T1 cells in these two types of preparations are similar. The extracellular matrix is scant and minimal necrosis is seen.
However, for the MAT cell line, interesting observations are made. First, various degrees of necrosis are observed in the MAT tumors, and the necrotic regions are dispersed over the tumor. Second, there are a few blood vessels with intra-luminal red blood cells, and lymphocytes.

No apparent difference is observed between the histological images of the LMTK cell pellets and LMTK tumors from Figure 4.9. The LMTK cells have similar sizes and shapes in cell pellets and in tumors. The extracellular matrix is scant and minimal necrosis is seen. The higher magnification image (100x) in Figure 4.10, however, shows slightly more cellular/nuclear orientation in the tumor than in the cell pellet.

**D) Discussion**

The 4T1 cell pellets and tumors have been shown to be similar: they have similar histopathologic features and similar ultrasonic characteristics (specifically BSC and attenuation). The similar ultrasonic characteristics are likely caused by the similar histopathologic features. In terms of histopathologic features, the 4T1 cells have similar sizes and shapes in the cell pellet form and in tumors. Also, the 4T1 cell pellets and tumors have similar components: the cell pellets are composed of tumor cells only, and the tumors are mainly composed of tumor cells with little extracellular matrix. In terms of ultrasonic characteristics, it is observed that the 4T1 cell pellets and tumors have similar BSC (except for the 65-MHz peak) and attenuation estimates. The observed similarity in BSC could be interpreted by the hypothesis that the 4T1 cell pellets and tumors have similar
scattering sites (whether being cell nuclei or whole cells or a combination of both) given their similar components. The observed similarity in attenuation could be analyzed in two aspects: the scattering and the absorption (note: attenuation is the combined effect of both scattering and absorption). The scattering from 4T1 cell pellets and tumors has been shown to be similar in terms of BSC. The absorptions are speculated to be similar based on the observation that 4T1 cell pellets and tumors have similar components and on the fact that acoustic absorption occurs primarily at the macromolecular level. With similar BSCs and absorptions, it makes sense that the attenuation estimates are found to be similar for 4T1 cell pellets and tumors. Therefore, the similarity in ultrasonic characteristics is a result of the similarity in histopathologic features. The 4T1 cell line represents an example where the tumor has relatively simple scattering structures such that the scattering from the tumor could be similar to that from the cell pellet.

In contrast to 4T1, the MAT cell pellets and tumors show different BSC and attenuation estimates, which can be attributed to their different histopathologic features. The main difference between MAT cell pellets and tumors histologically is that regions of necrosis were found in MAT tumors but not in MAT cell pellets. The scatterers are significantly different in necrotic regions than other regions where tumor cells are intact. There are no scatterers that are as big as tumor cells or cell nuclei in necrotic regions. The necrotic areas consist of fragmented cytoplasm and nuclei rather than intact nuclei and cytoplasm with definite shape. This fragmented cellular material is of variable size. Therefore, necrosis can be an important factor causing the BSC estimates to be significantly different in the
MAT tumors versus cell pellets. In addition to BSC, the absorptions are likely to be different as well, due to necrosis in MAT tumors. With different BSCs and absorptions, it makes sense that the attenuation estimates are different for MAT cell pellets and tumors. Therefore, MAT is an example where the tumor is more complex than the cell pellet and the anatomic structures such as necrosis are playing a role in scattering.

The situation for LMTK is different from the above two cases. The LMTK cell pellets and tumors have similar histopathologic features and similar attenuation results. The similarity in attenuation suggests that the macromolecular components are similar in the two types of media. The BSC behaviors are dramatically different, however. The BSC values are closer to each other for LMTK cell pellets and tumors at lower frequencies (<25 MHz). This might be explained by the observation that the histopathogic features are similar under the 40x microscope image. The higher degree of cellular/nuclear orientation seen in the LMTK tumor on the 100x image might be a reason for the BSC discrepancy between the LMTK cell pellets and the tumors at higher frequencies. Again, the structure function might be playing a role here. A higher degree of cellular orientation could mean the cell positions are more random.

The comparison between tumors and cell pellets of the same cell lines has provided insights into the scattering in tumors: (1) the 4T1 result is encouraging in the sense that it demonstrates the scattering from tumors of homogeneous morphology can be as similar as the scattering from the cell pellets of the same cell lines. The scattering model theories for such tumors can be developed by
studying the cell pellets; (2) the MAT result improves our understanding of tumor scattering in the sense that it demonstrates the important role of tumor anatomic details on scattering for tumors with heterogeneous morphology and complex scattering structures; (3) the LMTK result reveals the complexity of tumor modeling: even though the cellular components and cellular size are similar, the BSC could still be different. Further study of the tumor scattering process is required.

4.3.2 EHS tumor results

The attenuation-compensated BSC estimates from EHS tumors are shown in Figure 4.11 in comparison to 4T1 tumor results. The two tumor types show different BSC shapes: the EHS BSC is higher than the 4T1 BSC at low frequencies, and lower than the 4T1 BSC at higher frequencies. Comparing the B-spline trend lines for the two tumor types shows that the EHS has significantly more scattering at around 25 MHz than 4T1.

The attenuation (dB/cm) results for EHS tumors are presented in Figures 4.12 in comparison to 4T1 tumor results. EHS shows significantly lower attenuation values compared to 4T1.

An H&E stained histological slide for the EHS tumor is shown in Figure 4.13. Compared to the 4T1 tumor (Figure 4.7b), the EHS tumor has a similar cell size and shape, but different spatial organization. The EHS tumor cells are groups into spherical clusters, with a cluster radius of approximately 20 µm. With this unique structure, the EHS BSC curves turned out to be significantly different than the
4T1 curves. The excessive scattering at 25 MHz seems to have come from the spherical clustering structure: 25 MHz corresponds to a \( ka \) value of 2 for a 20-\( \mu \)m radius.

In addition to the unique spherical clustering structure, the EHS tumor appears to have more extracellular matrix than any other tumor types discussed in this chapter so far. Extracellular matrix has different macromolecular components than tumor cells, which might be the reason why the EHS attenuation is significantly lower than the 4T1 tumor attenuation.

The EHS results demonstrate how detailed tumor structure could affect ultrasound backscatter characteristics. Scattering models that appropriately takes into account these structure details will have great potential in tumor characterization.

### 4.3.3 Spontaneous fibroadenoma results

Four rats that had spontaneous fibrous tumors were evaluated, and three of them were diagnosed from histology to have fibroadenoma. The BSC and attenuation results from those three rats are presented in this subsection. Unlike the MAT, 4T1, or LMTK tumors, each of the spontaneous fibroadenoma tumors had a much larger size (greater than 10 mm in diameter). Therefore, three samples were trimmed from each tumor at different locations, and a total of nine samples from three tumors (animals) were scanned and evaluated.
The BSC and attenuation results for the three animals are shown in Figure 4.14 and Figure 4.15, respectively. The BSC figure shows that Animal 1 has a significantly higher BSC in the tumor than Animal 2 does. Both Animal 1 and Animal 2 have a steeper BSC versus frequency slope than Animal 3 does. The attenuation figure shows that both Animal 1 and Animal 2 have a much higher attenuation than Animal 3 does. Also, the within-animal variance in attenuation is much less than the cross-animal variance. Histology analysis found that the tumors for Animals 1 and 2 were mostly infarcted and hyalanized, whereas the tumor for Animal 3 was not (Figure 4.16). The histopathologic feature suggests that high-frequency ultrasound is sensitive to different disease conditions for the same disease, as long as the tissue structure changes with disease conditions.

4.3.4 Discussion on the *ex vivo* condition

The tumor results that have been discussed in this chapter were obtained from *ex vivo* scans. Working on *ex vivo* tumors has both advantages and disadvantages. The excised tumor samples allow for accurate attenuation estimation using the insertion-loss technique. Also, the excised tumor samples can be trimmed such that no skin layer is present in the acoustic path, which eliminates the problem of compensating for transmission loss. Furthermore, working on *ex vivo* samples allows for the study of high-frequency data. It is not feasible to scan the tumors *in vivo* using high-frequency (e.g., 80 MHz) single-element transducers. The disadvantage of working on the *ex vivo* data is that the *ex vivo* condition could be different from the *in vivo* condition. For instance, there is blood supply *in vivo*,
but not ex vivo, which could cause the scattering to be different. Therefore, the ex vivo BSC results do not necessarily represent the in vivo BSC results.

To get a sense of how close the ex vivo BSC results are to the in vivo results, the ex vivo BSC results for the 4T1 and MAT tumors discussed in Section 4.3.1 are compared to the in vivo BSC results from another study [58]. In [58], thirteen 4T1 tumors and eight MAT tumors were scanned in vivo using three imaging systems and five transducers covering the frequency range 3–22 MHz. This frequency range has some overlap with the bandwidth of the 20-MHz transducer used for the ex vivo scans in this dissertation. Therefore, the B-spline fits of the MAT and 4T1 tumor BSC results from the 20-MHz transducer were compared to the B-spline fits of the in vivo BSC results from [58]. Figure 4.17 shows the result of this comparison. The in vivo and ex vivo BSC results were similar for the 4T1 tumors. The in vivo MAT tumor BSC appears to be slightly higher than the ex vivo MAT tumor BSC.

4.4 Chapter Summary

This chapter summarizes the high-frequency attenuation and BSC results from various tumor types. The comparison between tumors and cell pellets of the same cell types provides a start point for accurately model tumor scattering. The fact that MAT and LMTK each has different BSCs between the tumor and the cell pellet suggests a closer analysis between the cell pellet and the tumor is required to proceed with tumor scattering modeling. The EHS and fibroadenoma results provide more examples of how unique tumor structures can affect the BSC, which presents opportunities for tumor modeling as well. More detailed analysis of the
tumor structures, structure functions, and modeling will be discussed in Chapters 5 and 6.

4.5 Figures

Figure 4.1 BSC versus frequency for five 4T1 cell pellets (blue) and fifteen 4T1 tumor samples (red). The two B-spline curves show the trend shapes for the cell pellets and tumors, respectively.
Figure 4.2 BSC versus frequency for five MAT cell pellets (blue) and thirteen LMTK tumor samples (red). The two B-spline curves show the trend shapes for the cell pellets and tumors, respectively.
Figure 4.3 BSC versus frequency for three LMTK cell pellets (blue) and thirteen LMTK tumor samples (red). The two B-spline curves show the trend shapes for the cell pellets and tumors, respectively.
Figure 4.4 Attenuation (dB/cm) versus frequency for five 4T1 cell pellets (blue) and fifteen 4T1 tumor samples (red).
Figure 4.5 Attenuation (dB/cm) versus frequency for five MAT cell pellets (blue) and thirteen MAT tumor samples (red).
Figure 4.6 Attenuation (dB/cm) versus frequency for three LTMK cell pellets (blue) and thirteen LMTK tumor samples (red).
Figure 4.7 Optical microscope images (40x) of H&E stained (a) 4T1 cell pellet, and (b) 4T1 tumor slides. Scale bars represent 50 µm.
Figure 4.8 Optical microscope images (40x) of H&E stained (a) MAT cell pellet, and (b) MAT tumor slides. Scale bars represent 50 µm.
Figure 4.9 Optical microscope images (40x) of H&E stained (a) LMTK cell pellet, and (b) LMTK tumor slides. Scale bars represent 50 µm.
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Figure 4.11 BSC versus frequency for EHS tumors (red) in comparison with 4T1 tumors (blue). The two B-spline curves show the trend shapes for the EHS (gray) and 4T1 (black), respectively.
Figure 4.12 Attenuation (dB/cm) versus frequency for EHS tumors (red) in comparison with 4T1 tumors (blue).
Figure 4.13 Optical microscope images (40x) of an H&E stained EHS slide. The scale bar represents 50 µm.
Figure 4.14 BSC versus frequency for nine fibroadenoma samples from three animals (rats) evaluated using the 20- and 40-MHz transducers.
Figure 4.15 Attenuation (dB/cm) versus frequency for nine fibroadenoma samples from three animals (rats) evaluated using the 20- and 40-MHz transducers.
Figure 4.16 Optical microscope images (40x) of H&E stained rat fibroadenoma slides: (a) infarcted and hyalanized, from Animal 1, and (b) not infarcted or hyalanized, from Animal 3. Scale bars represent 50 µm.
Figure 4.17 Comparison between the *in vivo* and *ex vivo* B-spline BSC curves for 4T1 and MAT tumors. The *in vivo* results were adapted from Figure 2 of [58], where the blue solid is the B-spline fit to the BSC results obtained from thirteen 4T1 tumors scanned *in vivo* using three imaging systems and five transducers, and the pink solid is the B-spline fit to the BSC results from eight MAT tumors scanned *in vivo* using the same imaging systems and transducers. 90% confidence interval is displayed for the *in vivo* results. The *ex vivo* B-spline curves (blue dashed for 4T1 and red dashed for MAT) were obtained from BSC results acquired by the 20-MHz transducer (Figures 4.1 and 4.2). No confidence interval is displayed for the *ex vivo* results for clarity.
CHAPTER 5

ESTIMATING STRUCTURE FUNCTION FROM HISTOLOGICAL TISSUE SECTIONS

This chapter discusses the estimation of structure function from histological tissue sections. The 2D structure functions are estimated for high-concentration cell pellets and tumors. The structure function estimated from the cell pellet histological tissue sections is related to the experimental and theoretical structure functions discussed in Chapter 3. The estimated structure function for tumors is compared to that for cell pellets to give insight into how tumor structures are affecting acoustic scattering. The structure functions related to unique structures are discussed. Also mentioned in this chapter is the 2D pair correlation function extracted from the histological tissue sections to help understanding the structure function.

5.1 Background and Introduction

In Chapter 3, the structure function has been estimated for high-concentration cell pellets from the BSC data. The estimated structure function curve has been attributed to the possibility that the high-concentration condition makes the spatial arrangement of cell positions less random than it would be for low concentrations. Two polydisperse structure function models have been developed to describe the effect of high concentration on cell spatial arrangement and the structure function. The models theoretically predict a structure function curve that agrees well with
the experimental one, particularly in terms of the peak position. However, the
models are not direct evidence to support the hypothesis that the structure
function curves estimated from the BSC curves are a result of the cell spatial
position correlation.

To provide direct evidence, we plan to examine the histological sections of the
samples. Ideally, if the 3D cell positions are known, the exact structure function
can be calculated using Equations (2.13) – (2.14). Unfortunately, determining the
3D cell positions is very difficult and may not be feasible. A less ideal alternative
would be to study the cell positions in 2D and determine the 2D structure function
$S_A(2k)$, where the subscript $A$ represents 2D (area). Precisely relating the 2D
structure function estimated from a thin section of finite thickness to the actual 3D
structure function is a difficult stereological problem, and to date there is no
generally applicable solution. However, studying the 2D structure function may
provide a qualitative comparison between the cell position correlation and the
backscattered data.

In addition to the structure function, the 2D pair correlation function $g_A(r)$ can
be determined from the tissue sections as well. The pair correlation function
provides a more intuitive way of understanding the spatial distribution of cells. In
3D, the pair correlation $g_\gamma(r)$ for a particular distribution of particles is defined as
the probability of finding a particle at distance of $r$ away from a given reference
particle, relative to that for a distribution of completely uncorrelated particles.
Therefore, $g_\gamma(r)=1$ for a distribution of uncorrelated particles. If the particle
positions are correlated, the local number density at a distance $r$ from the
reference particle is determined by $\overline{n}g_r(r)$, where $\overline{n}$ is the average number density of the entire system. The concept is the same for 2D. For an isotropic distribution of spheres of similar size, the pair correlation function $g_A(r)$ estimated from a 2D section is a rough approximation for the 3D pair correlation function $g_V(r)$ [59].

Estimating the structure function and pair correlation function from histological slides will eventually improve the understanding of tumor scattering. Chapter 4 points out that the BSC curves can be different for cell pellets and tumors of the same cell line (e.g., LMTK), although both the cell pellets and tumors are composed of primarily the same type of cells. It is hypothesized that the difference in BSC might have come from the difference in the spatial distribution of tumor cells. The 2D histological section study will provide an opportunity to test the hypothesis by exploring exactly how the distribution is different. Furthermore, the cell spatial distributions for different tumor structures can be compared, which will provide guidelines for tumor structure modeling.

5.2 Methods

An H&E stained tissue section was viewed under a light microscope (Olympus BX–51, Optical Analysis Corporation, Nashua, NH, USA). For each tissue slide, a 40 X TIF format picture was taken using the digital camera that was in sync with the microscope (see Figure 4.13 for a typical picture). Each image contained a 50 µm scale bar on the lower right of the image. The dimension of the image was $2560 \times 1920$ pixels, with a resolution of 5.72 pixels per micron. Only an area
of 1920 × 1920 pixels was analyzed for pair correlation and structure function estimation. This area covers the majority of the original image except a 640 × 1920 rectangle on the right side that contains the scale bar. The analyzed image area corresponds to a physical dimension of 336 × 336 µm². For each scanned biological sample type, two to three pictures from separate tissue sections were taken and analyzed.

A custom MATLAB routine was developed to allow manual determination of the nuclear center for each cell on the image (the nucleus center was assumed to overlap the cell center). The manual determination process was completed by clicking on the nucleus center on the image opened in MATLAB (Figure 5.1). The nuclear center coordinates were automatically recorded for further processing. As an example, 700 nuclear centers were found from the image shown in Figure 5.1, and plotted separately in Figure 5.2.

5.2.1 Structure function estimation

A matrix of the dimension 1920 × 1920 was created. The matrix dimension was the same as the pixel dimension of the image (Figure 5.1). The matrix elements that were corresponding to nuclear centers were assigned a value of one, with all remaining matrix elements being assigned a value of zero. The nuclear centers were essentially gridded. The gridding will not cause a problem because the gridding resolution (5.7 µm⁻¹) is sufficiently high compared to the cell/nuclear size. A fast Fourier transform was performed on the matrix. The squared modulus of the 2D Fourier transform (Figure 5.3), normalized to unity for large wave numbers, is the 2D structure function for all directions. A radial symmetry of the
structure function is observed, suggesting the isotropy of the system. Therefore, a radial averaging was performed, which resulted in a structure function $S_A(k)$ that is only dependent on the modulus of the wave number. For backscattering, we are interested in $S_A(2k)$. Converting $k$ to frequency in MHz using a speed of sound 1540 m/s will generate the frequency-dependent 2D structure function (e.g., Figure 5.4).

5.2.2 Pair correlation function estimation

The pair correlation function was estimated using the algorithm described in [60]. The algorithm is briefly summarized as follows. To calculate $g_A(r)$, a value of distance resolution $dr$ was picked first. The distance resolution cannot be too small because of limited number of points on the 2D image (Figure 5.2). It cannot be too large either, to avoid blurring any important structures in the pair correlation curve. A distance resolution of 0.87 µm (equivalent to five pixels of the original image) was chosen. For each distance $r$ at which the pair correlation function was to be calculated, each nuclear center was chosen as a reference point in turn. The number of nuclear centers that were at a distance between $r$ and $r + dr$ away from the reference point was counted, and averaged for all the reference points. This number was then normalized by $2\pi rdr$ (the area of the ring), and divided by the average number of nuclear centers per unit area. The edge effect was also properly accounted for. Specifically, when calculating the pair correlation function based on reference nuclear centers near the edge of the image, the circle of some radius $r$ may extend outside the image. The edge effect was correctly accounted for by determining how much angular extent of the circle
lies within the image. An example of estimated pair correlation function is shown in Figure 5.5.

5.3 Results and Discussions

5.3.1 Simulations

For comparison purposes, the estimated pair correlation functions and structure functions from three simulated point distributions are presented herein, before presenting the results from the tissue histological images. The three simulated 2D point distributions are: (a) the points are randomly distributed in the image (Figure 5.6a); (b) the points are clustered, with weaker clustering (Figure 5.6b); and (c) the points are clustered, with stronger clustering (Figure 5.6c). The same number density is used for the three simulations (900 points on a 336 × 336 µm² area).

The pair correlation function extracted from Figure 5.6a is approximately unity over the entire range of analyzed distance (Figure 5.7), as expected. The two clustering point distributions yield pair correlation functions that have high values at closer distances, and the pair correlation function gradually reduces to unity as the distance increases (Figure 5.7). The primary effect of clustering is the increased pair correlation function at closer distances. The stronger the clustering is, the higher the pair correlation function is at closer distances.

Similarly, the structure function extracted from Figure 5.6a is approximately unity over the entire frequency range being analyzed. Clustering increases the
lower-frequency component of the structure function. The strong clustering is, the higher the low-frequency structure function is (Figure 5.8).

The simulation results are presented for two reasons. The simulation results allow for comparison between simulations and the actual results from tumors. More importantly, the results demonstrate the feasibility of extracting the pair correlation function and structure function from a 2D region of similar dimensions and numbers of points.

5.3.2 Cell pellets

The estimated 2D structure function curves are presented in Figure 5.9 for high-concentration (74% volume fraction; same as Concentration II in Chapter 3) cell pellets of four cell lines: CHO, MAT, 4T1, and LMTK. The general trend of the 2D structure function curves appears to be similar to the structure function calculated from the ultrasonic scattering data (Figure 3.6), and to the polydisperse structure function curves in Chapter 3.

A one-to-one comparison between the 2D structure function and the structure function from ultrasonic scattering is shown in Figure 5.10, Figure 5.11, and Figure 5.12 for CHO, MAT, and 4T1, respectively. The peak position agree very well (within 5 MHz) between the two structure function curves for each of the three cell lines. The shape agreement is also observed for CHO and MAT. The magnitude agreement is not as good; only MAT shows good agreement in structure function curves (Figure 5.11). For CHO, the acoustic structure function is in general lower than the 2D structure function across the frequency range (Figure 5.10). The peak magnitude barely exceeds unity for the acoustic structure
function for CHO. The acoustic structure function curve could have been underestimated due to number density uncertainty, etc.

In general, the 2D structure function curves at least show convincing qualitative agreement with the curves and theories presented in Chapter 3. Also, the peak position of the 2D structure function curve shows good correlation with cell size: 4T1 has the largest cell size (cell radius: 8.9 µm) out of the four cell lines, whereas the other three cell lines have similar but smaller cell sizes (cell radius: 6.8, 7.3, and 7.3 µm for CHO, MAT, and LMTK, respectively). Therefore, the structure function peak for 4T1 is at the lowest frequency compared to other cell lines, and the structure function peak of the other cell lines are located at similar frequencies.

The estimated pair correlation curves (Figure 5.13) for the four cell lines give insight into what aspects of cell arrangement have cause the structure function behavior. All four cell lines have a zero pair correlation for approximately \( r < 5 \mu m \). This is because the cells are considered non-overlapping spheres. The width of the zero-correlation region is determined by the cell diameter. The 4T1 curve has the widest zero-correlation region because of the large cell radius. Note that the 2D pair correlation function is only a very rough approximation of the 3D case, because cells at different depths could appear to be slightly overlapping in 2D, reducing the width of the zero-correlation regions. Although very significant, the non-overlapping condition alone is not the reason for the observed structure function behavior. What has caused the observed structure function behavior is a combination of the non-overlapping condition and
the high concentration. Because of the high concentration, the pair correlation curve immediately sees a sharp peak above unity after the zero-correlation region. If the cell concentration had been much lower, then the sharp peak could not have been observed, and the resulting structure function curve would have been closer to unity.

5.3.3 Tumors versus cell pellets

The structure function and pair correlation function were estimated for 4T1, MAT, LMTK, and EHS tumors. Necrotic regions were not analyzed for MAT tumors, because it is hard to find the nuclear center in the necrotic region. Therefore, the MAT results may not fully reflect the entire MAT tumor structure.

The tumor structure functions are compared to the cell pellet structure functions for 4T1, MAT, and LMTK (Figure 5.14). The tumors appear to have less structure function effect than the cell pellets do. The tumor has a lower structure function than the cell pellet of the same cell type at frequencies lower than 50 MHz, meaning that the tumor cell arrangement has a less destructive effect in the destructive frequency range. In the constructive frequency range (around the scattering peak), the tumor shows a less constructive effect as well. The observed 2D structure functions in Figure 5.14 are well correlated to the BSC curves in Figures 4.1 – 4.3 also. Overall the tumor structure function curves are smoother than the cell pellet curves. This could suggest that the cell arrangement is more random in tumors than in cell pellets.

The pair correlation function comparison between cell pellets and tumors (Figure 5.15) shows three important features: (a) the zero-correlation region is
wider for the cell pellets than for the tumor; (b) the correlation peak occurs at a
closer distance for the tumor than for the cell pellet; (c) the peak values are higher
for the cell pellet than for the tumor. The three features reveal the structural
difference between cell pellets and tumors. Feature (a) might be explained by the
fact that some cells may be pushed in tumors so that the centers between certain
cell pairs are shortened. The same reason could be attributed to feature (b).
Feature (c) suggests that the cell arrangement in tumors is more random than in
cell pellets, consistent with structure function observation.

5.3.4 EHS and clustering

The EHS tumor is different from 4T1, MAT, and LMTK in the cell
arrangement: the cells are grouped in EHS, with typically less than 20 cells per
group, whereas other tumors do not show such a clustering phenomenon. To show
how cell clustering affects the structure function in tumors, the 2D structure
functions are compared for all the four tumor types (Figure 5.16). The tumor
structure functions seem to be similar, except that the EHS show higher structure
function values around 25 MHz compared to other tumor types. This phenomenon
can explain the experimental BSC behavior of EHS compared to other tumor
types (e.g., Figure 4.11): EHS shows more scattering at around 25 MHz than
other tumor types. Therefore, the results suggest that the additional scattering at
25 MHz for EHS is related to the EHS tumor cell arrangement.

The pair correlation function comparison between EHS and other tumor types
(Figure 5.17) provides more detailed information about cell spatial arrangement.
EHS has a slightly higher peak around 10 µm than other tumor types. More
interestingly, there is a dip around 15 µm in the EHS curve but not in other tumor curves (Figure 5.17). This dip might be a result of the clustering structure. There is empty space between different clusters, resulting in decreased probability of finding a nuclear center in that space. If this hypothesis is true, then the dip position would be related to the cluster size.

Comparing the EHS pair correlation function with the simulation results for clustered points (Figure 5.7), it is hard to find any similarity between the two. In fact, the EHS pair correlation function appears to be more similar to that of non-clustering tumors (although noticeable difference exists). Note that non-overlapping condition was not applied to the simulations. Therefore, the comparison between EHS and simulation results suggests that the effect of clustering on structure function is significantly reduced due to the non-overlapping condition. Also, the close distance between the EHS tumor clusters may have further reduced the effect of clustering on structure function.

5.3.5 Significance and limitations

The 2D structure function and pair correlation function estimated from histological sections provide direct evidence to show that the cell spatial arrangement affects acoustic scattering in real tumors. The result also qualitatively confirms the polydisperse structure function model developed in Chapter 3. It provides guidelines as to how to model the structure function properly. Overall the results significantly improved the understanding of how tumors scatter ultrasound.
However, there are several limitations in the 2D histological section study. First, the 2D structure function and pair correlation function are not exactly the same as those in 3D. Therefore, we would not be able to directly divide the BSC by the 2D structure function to estimate the incoherent BSC. Second, the number of histological sections analyzed for each sample type is not large enough. Only three images on average have been analyzed for each sample type. Third, manual determination of the nuclear center is time-intensive, and subject to human error. Automatic procedures are currently not available.

5.4 Chapter Summary

This chapter has introduced histology analysis methods that can be used to relate histological features to scattering and the structure function. The method of extracting the 2D structure function and pair correlation function from histological sections is shown to be feasible. The 2D structure function and pair correlation function have been successfully estimated from 2D histological sections for various sample types. The comparisons among different sample types improve our understanding of how tumor microstructure (particularly cell arrangement) affects scattering. Specifically, the tumor cells are shown to be more randomly distributed than in cell pellets. Tumor cell clustering is shown to moderately affect the structure function, and should be taken into consideration when scattering models are being developed.
5.5 Figures

Figure 5.1 An optical microscope image (40x magnification; 1920 × 1920 pixels) superimposed with nuclear centers in yellow dots that were manually determined. Displayed are 700 nuclear center points.
Figure 5.2 Nuclear centers extracted from Figure 5.1 plotted on the XY plane. The coordinates were gridded (sampled), with a spatial sampling rate of 5.7 µm⁻¹.
Figure 5.3 Two-dimensional plot of the squared modulus of the fast Fourier transform of the points plotted on Figure 5.2.
Figure 5.4 An estimated structure function versus frequency curve, obtained by radial averaging of the modulus of the 2D Fourier transform shown in Figure 5.3.
Figure 5.5 An estimated pair correlation function versus distance curve, obtained from points presented in Figure 5.2.
Figure 5.6 Simulated 2D distributions of random points for, (a) a completely random distribution, (b) a clustering distribution with weaker clustering, and (c) a clustering distribution with stronger clustering.
Figure 5.7 Estimated pair correlation functions from the point distributions shown in Figure 5.6.
Figure 5.8 Estimated structure functions from the point distributions shown in Figure 5.6.
Figure 5.9 Comparison among the average estimated structure functions from the histological sections for high-concentration CHO, 4T1, MAT, and LMTK cell pellets. Each curve is an average of three independent measures on different histological images.
Figure 5.10 Comparison between the 2D structure function estimated from histological slides and the structure function obtained from ultrasonic backscattering coefficients (see Figure 3.6a) for high-concentration CHO cell pellets. The blue curve is an average of three independent measures on different histological images.
Figure 5.11 Comparison between the 2D structure function estimated from histological slides and the structure function obtained from ultrasonic backscattering coefficients (see Figure 3.6b) for high-concentration MAT cell pellets. The blue curve is an average of three independent measures on different histological images.
Figure 5.12 Comparison between the 2D structure function estimated from histological slides and the structure function obtained from ultrasonic backscattering coefficients (see Figure 3.6c) for high-concentration 4T1 cell pellets. The blue curve is an average of three independent measures on different histological images.
Figure 5.13 Comparison among the average estimated pair correlation functions from the histological sections for high-concentration CHO, 4T1, MAT, and LMTK cell pellets. Each curve is an average of three independent measures on different histological images.
Figure 5.14 Comparison between the average 2D structure function curves for (a) 4T1 cell pellets and tumors, (b) MAT cell pellets and tumors, and (c) LMTK cell pellets and tumors.
Figure 5.15 Comparison between the average pair correlation curves for (a) 4T1 cell pellets and tumors, (b) MAT cell pellets and tumors, and (c) LMTK cell pellets and tumors.
Figure 5.16 Comparison among the average 2D structure function curves for 4T1, MAT, LMTK, and EHS tumors.
Figure 5.17 Comparison among the average 2D pair correlation function curves for 4T1, MAT, LMTK, and EHS tumors.
This chapter discusses modeling the scattering of different tumor structures. Two cases are considered herein, the tumor cells are homogenously distributed (e.g., 4T1, LMTK), and the tumor cells are clustered (e.g., EHS). The effect of the structure function on scatterer parameter estimation is discussed. A new scattering model that is capable of detecting tumor cell clustering is developed.

6.1 Background and Introduction

Commonly used scattering models for tumors are based on the assumption that the scatterer positions are uncorrelated. The fluid-filled sphere model and the Gaussian form factor model are two such examples. For these two models, the BSC is determined by two parameters, the effective scatterer size and the effective acoustic concentration (EAC). The EAC is the product of the number of scatterers per unit volume and the square of the fractional change in the impedance between the scatterer and the surrounding medium. The effective scatterer size affects the slope and the magnitude of the BSC versus frequency curve. The EAC only affects the magnitude of the BSC curve.

Previous chapters have shown that the spatial correlation among scatterer positions in tumors plays a significant role in scattering, and modifies the BSC curve substantially. Therefore, models such as the fluid-filled sphere model and
the Gaussian form factor model may not give accurate parameter estimates. It is important to evaluate how the structure function affects model parameter estimates, by comparing the parameters estimated using models that do not take into account the structure function versus parameters estimated using models that take into account the structure function. Also, a comparison between different structure function models would be interesting.

For the above reasons, three models are compared in this chapter to evaluate the effect of structure function on parameter estimates: the fluid-filled sphere model (denoted FF), the fluid-filled sphere model incorporating the structure function estimated from 2D histological slides (denoted SF0), and the fluid-filled sphere model incorporating a polydisperse structure function (denoted SF1; the polydisperse structure function is Polydisperse Model I described in Chapter 3). SF0 is included in comparison because the 2D structure function estimated from histology is the only available structure function estimated directly from tumors, although 2D and 3D can be different. SF1 is included in comparison because it is a theoretical model that works reasonably well for high-concentration cell pellets at high frequencies where there is a peak in the BSC curve. The theoretical structure function used in SF1 is not the same curve as the estimated 2D structure function used in SF0. Therefore, SF0 and SF1 represent two different structure function models.

The structure function for tumors with cell clustering is different than that for tumors that have a homogenous cell distribution (Figure 5.16). SF1 is developed for a homogenous cell distribution, but not for a clustering structure. A model that
is capable of detecting the clustering structure is desirable. For the EHS tumors, the clustering of cells not only changes the structure function, but may also alter the scattering sites as well – the cell clump may become a scattering site as well because of possible impedance contrast between the clump and the extracellular matrix. Therefore, a model that assumes two primary scatter sizes is developed for the EHS tumor and is applied for detecting clustering.

6.2 Effect of Structure Function on Model Parameter Estimation

6.2.1 Methods

Three models (FF, SF0, SF1) are applied to the experimental BSC data from MAT, 4T1, LMTK, and EHS tumor data. The scatterers are assumed to follow a gamma distribution as described in Equation (3.2). To reduce the number of unknowns for the inverse problem, the Schulz width factor \( z \) is assumed to be 50 for all three models, based on measurements made in Chapter 3.

The first model, FF, is calculated using

\[
BSC_{FF}(2k) = \frac{1}{2} \pi \bar{n} \gamma_0^2 k^{-4} \left[ 1 - \chi - k \psi' + \frac{1}{4} k^2 (\zeta'' + \chi'') \right].
\]  

(6.1)

where \( \bar{n} \) is the scatterer number density, \( \gamma_0^2 \) is the mean-square variation in acoustic impedance per scatterer as in Equation (2.5), the product \( \bar{n} \gamma_0^2 \) is the EAC, and the quantities \( \chi, \psi', \zeta'', \) and \( \chi'' \) are defined in Equation (A.5). Equation (6.1) is an analytical expression of the BSC for fluid-filled spheres with gamma-distributed radii. Equation (6.1) is a slightly modified form of Equation (19) in [53].
The second model, SF0, is calculated by

\[ BSC_{SF0}(2k) = BSC_{FF}(2k) \cdot S_o(k), \]  

(6.2)

where \( BSC_{FF}(2k) \) is calculated using Equation (6.1), and the structure function \( S_o(k) \) is assumed to be known, calculated for each tumor type by performing a seventh-order polynomial fit to the corresponding structure function curve presented in Figure 5.16.

The third model, SF1, is calculated by

\[ BSC_{SF1}(2k) = BSC_{FF}(2k) \cdot S_i(k), \]  

(6.3)

where the structure function \( S_i(k) \) is calculated using Equation (A.1).

The unknowns for model FF are the average radius \( a \) and the effective acoustic concentration EAC (dB/mL). The unknowns for model SF0 are the same as for model FF. The unknowns for model SF1 are the average radius \( a \), the EAC, and the volume fraction. A least squares minimization is performed using an exhaustive search approach, to find parameter values that minimize the mean square error between the log-scaled experimental and theoretical BSCs.

Considering that the parameter estimation is often dependent on frequency, three frequency ranges are analyzed: 11 – 31 MHz (data from the 20-MHz transducer), 11 – 52 MHz (data from the 20- and 40-MHz transducers), and 11 – 105 MHz (data from the 20-, 40- and 80-MHz transducers). Frequency ranges that contain only the data from 40- or 80-MHz transducers are not used, because the high-frequency BSC curve for tumors tends to be horizontal.
6.2.2 Results

The estimated scatterer radii and effective acoustic concentrations for MAT, 4T1, LMTK, and EHS tumors using the three models are shown in Figure 6.1 (11-31 MHz), Figure 6.2 (11-52 MHz), and Figure 6.3 (11-105 MHz). The estimated scatterer volume fraction using model SF1 is shown in Figure 6.4.

Comparing FF and SF0, FF underestimates the scatterer radius or the effective acoustic concentration in most cases compared to SF0. Comparing SF0 and SF1, SF0 always underestimates the scatterer radius and overestimates the effective acoustic concentration, regardless of which frequency range is being used. Comparing the frequency ranges, all the models tend to have lower radius estimates and higher EAC estimates. However, the volume fraction estimated from model SF1 does not show a clear dependence on which frequency range is used for MAT, 4T1, and LMTK tumors (Figure 6.4). The volume fraction estimates for those three tumor types are in a reasonable range. The EHS shows a higher volume fraction estimate than the other tumor types. This observation allows for distinction between EHS and other tumor types. However, a higher volume fraction estimate for EHS tumors is not consistent with the fact that EHS tumors have a greater amount of the extracellular matrix than the other three tumor types. This inconsistency may be caused by cell clustering in EHS.

6.2.3 Discussions

The comparison among three models shows the model parameter estimates are sensitive to the structure function. Different estimated values are obtained when different structure functions are used. The structure function can alter both the
magnitude and the frequency dependence of the BSC curve. Therefore, both the scatterer size and the effective acoustic concentration estimations may be affected.

For the lower frequency range (11 – 31 MHz, Figure 6.1) typically used in most QUS studies, the structure function has values that are much lower than unity, which effectively reduces the BSC magnitude. The widely used FF model that does not take into account this effect underestimates the effective acoustic concentration (Figure 6.1b). The SF1 model that works reasonably well for the high-concentration cell pellets overestimates the effective acoustic concentration compared to SF0 as a reference, because the structure function in SF1 has values that are much lower than the measured 2D structure function used in SF0. Therefore, the polydisperse structure function models developed in Chapter 3 appear to overestimate the regularity of the spatial arrangement of tumor cells. The polydisperse structure functions work better for cell pellets because the cell pellets have a more regular cell arrangement than tumors, as is evidence in Figure 5.14 where the cell pellet structure functions are lower than the tumor structure functions at lower frequencies.

The phenomenon that the estimated scatterer size and effective acoustic concentration depends on the analysis frequency has been observed frequently in QUS studies. This dependency is undesirable, because it obscures the biological meaning of the parameters. Parameters estimates that are independent of analysis frequency may be more robust for tissue characterization purposes. Model SF1 has an advantage in this aspect because it is capable of generating scatterer
volume fraction estimate that is independent of analysis frequency, despite the fact that the model underestimates the randomness in tumor cell arrangement. Note that model SF1 is developed for a homogenous distribution of cells, hence the scatterer volume fraction estimate for EHS tumors is not accurate and can only be used for classification purpose.

6.3 A Model for the EHS Tumor

This subsection introduces a model that describes the clustering feature of the EHS tumor cells. The model assumes that the cell clumps are scattering sites in addition to the individual spheres. The BSC is expressed as:

\[
\frac{1}{3} \sum_{i} \left( \frac{0}{CI_i} \right) \left( \frac{EAC_i}{EAC} \right) fBSC_i + \frac{1}{3} \sum_{i} \left( \frac{EAC_i}{EAC} \right) fBSC_i,
\]

(6.1)

where \( fBSC \) is the fluid-filled sphere BSC, \( N \) is the number of spheres per cluster, \( EAC \) is the acoustic concentration of the spheres, and \( CI_i \) is the clustering index which means the ratio of cluster \( EAC \) to sphere \( EAC \) in the model. The size distribution of the spheres may be taken into account in the model also. For the subsequent analysis, the spherical radius is assumed to follow a Schulz distribution. A Schulz width factor \( z = 50 \) is used to take into account the polydispersity of the spherical radius.

The reasons why the clustering index \( CI_i \) may be used to characterize the “clustering strength” are the follows: (1) For a given \( N \), if most of the spheres are not clustered and only a small portion of the spheres are clustered, then the number density, consequently the \( EAC \), of the clusters would be lower compared to the case when more spheres are clustered. (2) If the impedance contrast
between the clusters and the extracellular matrix is higher, the scattering from the cluster would be higher as a result.

The proposed model is applied to MAT, 4T1, LMTK, and EHS tumors to estimate the clustering index $C_\text{I}_0$ (in dB) of each tumor type (Figure 6.5). The EHS shows the highest clustering index for all the frequency ranges, suggesting that the model is able to detect the clustering structure. Also, the MAT shows a higher clustering index than the 4T1 and LMTK, which might be attributed to the necrosis in the MAT tumors. The necrosis in the MAT tumors effectively creates a clustering phenomenon because of missed cells due to necrosis.

6.4 Chapter Summary

The effects of structure function on scatterer parameter estimation have been discussed. The scatterer radius and effective acoustic concentration of the widely used fluid-filled sphere model have shown to be sensitive to the choice of structure function. The polydisperse structure functions developed in Chapter 3 underestimated the randomness of the cell arrangement in tumors. More accurate structure functions for tumors are required to model tumor scattering more accurately. As an example of modeling unique tumor structures, the new model developed for EHS has been shown to be able to detect clustering and model the clustered tumor cell structure.
6.5 Figures

Figure 6.1 (a) Estimated scatterer radii, and (b) estimated effective acoustic concentrations for MAT, 4T1, LMTK, and EHS tumors using three models (FF, SF0, and SF1) in the frequency range of 11 – 31 MHz. The estimated parameter value for a tumor type represents the average from all the samples of that tumor type, and error bars represent the standard deviation.
Figure 6.2 (a) Estimated scatterer radii, and (b) estimated effective acoustic concentrations for MAT, 4T1, LMTK, and EHS tumors using three models (FF, SF0, and SF1) in the frequency range of 11 – 52 MHz. The estimated parameter value for a tumor type represents the average from all the samples of that tumor type, and error bars represent the standard deviation.
Figure 6.3 (a) Estimated scatterer radii, and (b) estimated effective acoustic concentrations for MAT, 4T1, LMTK, and EHS tumors using three models (FF, SF0, and SF1) in the frequency range of 11 – 105 MHz. The estimated parameter value for a tumor type represents the average from all the samples of that tumor type, and error bars represent the standard deviation.
Figure 6.4 Estimated volume fractions for MAT, 4T1, LMTK, and EHS tumors using model SF1 in three different frequency ranges. The estimated parameter value for a tumor type represents the average from all the samples of that tumor type, and error bars represent the standard deviation.
Figure 6.5 Estimated clustering indexes for MAT, 4T1, LMTK, and EHS tumors in three different frequency ranges. The estimated parameter value for a tumor type represents the average from all the samples of that tumor type, and error bars represent the standard deviation.
CHAPTER 7

CONCLUSIONS AND FUTURE WORK

This dissertation investigates the scattering from tumors by studying the scattering from biological media of different degrees of complexity. Our understanding of the acoustic scattering processes in tumors has been improved through this investigation. The comparison between lower and higher concentrations of cell pellets demonstrates that the concentration affects the spatial arrangement of cells, and thus the structure function and BSC. The structure function of high-concentration cell pellets is modeled reasonably well by the proposed polydisperse structure function models. The comparison between cell pellets and tumors of the same cell types, and between different tumor types, reveals that unique tumor structures could have special scattering patterns. The estimated 2D structure function and pair correlation function from histological slides confirm the significance of structure function in dense media, support the proposed structure function models for high-concentration cell pellets, and reveal the cell arrangement difference between cell pellets and tumors. The unique clustering tumor structure can be detected using models that take into account the clustering structure. The modeling of the scattering from tumors with homogenously distributed tumor cells requires further study. This chapter concludes the dissertation and suggests possible directions for future studies.
7.1 Structure Function Estimation and Modeling for Cell Pellets

The structure function has been isolated from the BSC by comparing two concentrations of cell pellets. The experiment could be improved by including more concentrations and extending to lower concentrations. For model validation purposes, physical phantoms of similar scatterer sizes and concentrations to the cell pellets could be produced and scanned.

Another future direction could be to study the effect of the finite transducer beamwidth on the estimation of the structure function. The theories and models discussed in this dissertation are based on the assumption that the particles are scanned by a plane wave extending infinitely in space. The sound field produced by a weakly-focused single-element transducer has a finite boundary. The scattering volume is thus limited. The structure function of a limited scattering volume can be different from that of a much larger volume. Theoretical and experimental studies should address issues such as determining the minimal required scattering volume for obtaining a satisfactory estimate of the structure function.

7.2 The Comparison between Cell Pellets and Tumors

The comparison between cell pellets and tumors of the same cell types is the key to understanding tumor scattering. The observed difference in BSC between cell pellets and tumors of the same cell types has been attributed to the unique tumor structures such as regions of necrosis and the presence of blood vessel in tumors, and the structure function differences between cell pellets and tumors.
However, the above factors still do not fully explain why the homogenous LMTK tumor has an entirely different BSC behavior than the LMTK cell pellet does. Investigating additional factors that may cause the difference will significantly improve tumor scattering modeling. Further studies could be focused on histology analysis. Automatic or semi-automatic image processing techniques that are capable of detecting objects on a histological image can be developed and applied to analyze the histology systematically. The idea of object detection should not be limited to nucleus or cytoplasm. Rather, any object that is of the scale of interest could be detected and analyzed. Further, different staining techniques could be explored. Structure differences that cannot be distinguished by the currently used H&E staining might be seen if another staining method is applied. Additionally, the two-dimensional impedance map (2DZM) techniques that are currently being developed by other researchers will have the potential to be used for identifying the additional factors that caused the difference in scattering between LMTK cell pellets and tumors.

7.3 Structure Function Estimation from Histological Sections

Directly measuring the structure function from histology is a useful approach to understanding the structure function. The current study on estimating the structure function from 2D histological sections could be improved by a number of future steps. First, more images should be analyzed to yield a more accurate estimation of the 2D structure function. Different magnifications, different image sizes can be explored as well. Second, an automatic or semi-manual cell (or nucleus) center detection algorithms could be developed to allow for the analysis of more images.
Third, theoretical stereological studies can be pursued to develop computational techniques for estimating the 3D structure function from the 2D structure function.

Future studies may also extend to measuring the 3D structure function directly by analyzing series adjacent thin slices, similar to the 3DZM approach.

7.4 Tumor Scattering Modeling

Tumor scattering modeling has the potential to be improved by taking into account the structure functions. Accurate analytical structure functions for tumors are not yet available. Such structure functions could be developed by modifying existing structure functions such as the polydisperse structure functions introduced in Chapter 3, according to the cell spatial arrangement in tumors. Alternatively, empirical structure function curves could be derived from the structure function curves for the tumors at various cell radii and volume fractions.

The new model introduced in Chapter 6 for EHS modeling has been shown to work well in detecting the clustering feature of EHS. However, it has not been tested on other tumors that have a similar clustering cell arrangement. The model will be more convincing if it is tested in future studies on another tissue with a clustering cell structure. Nevertheless, the new model is encouraging in the sense that it demonstrates how sensitive QUS is to unique tumor structures, and how the unique tumor structure may be detected by scattering models.

In a practical point of view, it could be a future step to develop scattering models that are sensitive to a particular structure, or scattering models that can
yield parameters that are not dependent on the analysis frequency. Such models will improve the applicability of QUS techniques. A long-term goal of this investigation is to provide scattering models that can be used clinically for tissue characterization. Future studies and analysis have to be extended to *in vivo* data.
APPENDIX A

ANALYTICAL EXPRESSION OF THE STRUCTURE FUNCTION FOR POLYDISPERSE MODEL I

The structure function expressed in Equation (3.1) is a double integral, where the probability density function $f(x)$ was given in Equation (3.2), and the partial structure functions $H_{ij}$ was given by Blum and Stell [51]. The analytical expression of Equation (3.1) has been derived in Equation (2) of [52]. For backscattering, Equation (2) of [52] may be modified as:

$$S(2k) = 1 - \frac{\bar{n}}{4k^3(\Xi^2 + \Psi^2)} \times$$

$$\left\{ \lambda \left[ \lambda (\Xi \delta_1 - \Xi \delta_4) + \lambda' (\Xi \delta_2 - \Xi \delta_4) + \mu (\Xi \delta_1 + \Psi \delta_4) + \mu' (\Xi \delta_2 + \Psi \delta_4) \right] 
+ \lambda' \left[ \lambda (\Xi \delta_1 - \Xi \delta_3) + \lambda' (\Xi \delta_2 - \Xi \delta_3) + \mu (\Xi \delta_1 + \Psi \delta_3) + \mu' (\Xi \delta_2 + \Psi \delta_3) \right] 
+ \mu \left[ \lambda (\Xi \delta_1 + \Psi \delta_4) + \lambda' (\Xi \delta_2 + \Psi \delta_4) + \mu (\Xi \delta_1 - \Psi \delta_4) + \mu' (\Xi \delta_2 - \Psi \delta_4) \right] 
+ \mu' \left[ \lambda (\Xi \delta_1 + \Psi \delta_3) + \lambda' (\Xi \delta_2 + \Psi \delta_3) + \mu (\Xi \delta_1 - \Psi \delta_3) + \mu' (\Xi \delta_2 - \Psi \delta_3) \right] \right\}, \quad (A.1)$$

where

$$\Xi = 1 - (2\pi / \Delta)(1 + \frac{1}{2} \pi \xi / \Delta) \bar{n}(2k)^3 (2k \zeta' - \psi')$$

$$- (2\pi / \Delta) \bar{n}(2k)^{-3} [(\zeta' - \zeta')' + (\frac{1}{4} \pi \xi / \Delta)(\zeta' - \zeta'')]$$

$$- (\pi / \Delta)^2 \left( \frac{\bar{n}}{4k^2} \right)^2 [(\zeta - 1) (\zeta' - \zeta') - (\zeta' - \zeta')^2]$$

$$- (2k \zeta' - \psi')(2k \zeta'^{''' - \psi'}') + (2k \zeta'^{''} - \psi')^2 \right), \quad (A.2)$$
\[\Upsilon = \left(2\pi / \Delta\right) \left(1 + \frac{1}{2} \pi \xi_3 / \Delta\right) \bar{n} (2k)^{-3} \left(\chi + 2k^2 \zeta'' - 1\right) - \left(2\pi / \Delta\right) \bar{n} (2k)^{-2} \left[2k\zeta'' - \psi' + \left(\frac{1}{4} \pi \xi_2 / \Delta\right) (2k\zeta''' - \psi')\right] \]

\[-\left(\pi / \Delta\right)^2 \left(\frac{\bar{n}}{4k^2}\right)^2 \left[(2k\zeta' - \psi')(\chi'' - \zeta'')\right] \quad \text{(A.3)}\]

\[+ 2\left(\pi / \Delta\right)^2 \left(\frac{\bar{n}}{4k^2}\right)^2 (2k\zeta'' - \psi')(\chi' - \zeta') \]

\[-\left(\pi / \Delta\right)^2 \left(\frac{\bar{n}}{4k^2}\right)^2 (2k\zeta''' - \psi'')(\chi - 1) \]

and

\[\delta_1 = \left(\pi / \Delta\right) \left[2 + (\pi / \Delta) \left(\xi_3 - \frac{\bar{n}}{2k} (2k\zeta''' - \psi'')\right)\right] \]

\[\delta_2 = \left(\pi / \Delta\right)^2 \frac{\bar{n}}{2k} (2k\zeta'' - \psi') \]

\[\delta_3 = -\left(\pi / \Delta\right)^2 \frac{\bar{n}}{2k} (2k\zeta' - \psi) \]

\[\delta_4 = (\pi / \Delta)[2k - (\pi / \Delta) \frac{\bar{n}}{2k} (\chi' - \zeta')] \]

\[\delta_5 = (\pi / \Delta)^2 \left(\frac{\bar{n}}{2k} (\chi - 1) + k\xi_2\right) \]

\[\delta_6 = (\pi / \Delta)^2 \frac{\bar{n}}{2k} (\chi'' - \zeta'') \quad \text{(A.4)}\]

and

\[\zeta' = bc \]
\[ \zeta^* = b^2 c(c + 1) , \]
\[ \zeta'' = b^3 c(c + 1)(c + 2) , \]
\[ \psi = v_1^{c/2} \sin[c \tan^{-1}(2bk)] , \]
\[ \psi' = bcv_1^{(c+1)/2} \sin[(c + 1) \tan^{-1}(2bk)] , \]
\[ \psi'' = b^2 c(c + 1)v_1^{(c+2)/2} \sin[(c + 2) \tan^{-1}(2bk)] , \]
\[ \mu = 2^c v_2^{c/2} \sin[c \tan^{-1}(bk)] , \]
\[ \mu' = 2^{c+1} bcv_2^{(c+1)/2} \sin[(c + 1) \tan^{-1}(bk)] , \]
\[ \chi = v_1^{c/2} \cos[c \tan^{-1}(2bk)] , \]
\[ \chi' = bcv_1^{(c+1)/2} \cos[(c + 1) \tan^{-1}(2bk)] , \]
\[ \chi'' = b^2 c(c + 1)v_1^{(c+2)/2} \cos[(c + 2) \tan^{-1}(2bk)] , \]
\[ \lambda = 2^c v_2^{c/2} \cos[c \tan^{-1}(bk)] , \]
\[ \lambda' = 2^{c+1} bcv_2^{(c+1)/2} \cos[(c + 1) \tan^{-1}(bk)] , \]
\[ v_1 = [1 + (2bk)^2]^{-1} , \]
\[ v_2 = [4 + (2bk)^2]^{-1} , \]

(A.5)
\[ \xi_2 = \overline{n} b^2 c (c + 1), \quad \xi_3 = \overline{n} b^3 c (c + 1)(c + 2), \quad \Delta = 1 - \pi \xi_3 / 6, \]

\[ b = 2a / (z + 1), \quad c = z + 1. \]

We would like to point out two typographic errors in the expression of \( \Xi \) in Equation (2) of [52]. The expression of \( \Xi \) is correctly printed in Equation (A.2) above. Also, we would like to point out that Equation (A.5) was modified from Table I of [53] for backscattering.
APPENDIX B

ANALYTICAL EXPRESSION OF THE STRUCTURE FUNCTION FOR POLYDISPERSE MODEL II

We start from Equation (3.4) to derive the analytical expression of the structure function for Polydisperse Model II derived from the fluid-filled sphere form factor. The probability density function \( f(x) \) and the partial structure functions \( H_{ij} \) in Equation (3.4) were the same as those in Equation (3.1). The scattering amplitude \( \Phi_i(2k) \) in Equation (3.4) takes the following form derived from the fluid-filled sphere form factor:

\[
\Phi_i(2k) = \frac{\gamma_z}{8k} \left[ \sin(2kx_i) - 2kx_i \cos(2kx_i) \right], \quad (B.1)
\]

where \( \gamma_z = \frac{2(Z_0 - Z)}{Z} \), which describes the acoustic impedance contrast between the sphere \( Z \) and the background \( Z_0 \). However, the acoustic impedance contrast is irrelevant to Polydisperse Model II because \( \frac{\gamma_z}{8k} \) is a common factor between the numerator and denominator of the fraction in Equation (3.4), and is cancelled out.

Calculating the double integral in Equation (3.4) gives the analytical expression of \( S(2k) \) for Polydisperse Model II as

\[
S(2k) = 1 + \frac{I_N(2k)}{I_D(2k)}, \quad (B.2)
\]
where

\[ I_N(2k) = (M - k\Lambda')/\bar{n}, \]

\[ I_D(2k) = -\frac{1}{8k^3(\Xi^2 + \bar{Y}^2)} \times \]

\[
\left\{ \begin{array}{l}
\Lambda [\lambda(\bar{Y}\delta_1 - \Xi\delta_6) + \Lambda'(Y\delta_2 - \Xi\delta_4) + M(\Xi\delta_1 + Y\delta_6) + M'(\Xi\delta_2 + Y\delta_4)] \\
+ \Lambda' [\lambda(\bar{Y}\delta_2 - \Xi\delta_4) + \Lambda'(Y\delta_3 - \Xi\delta_5) + M(\Xi\delta_2 + Y\delta_4) + M'(\Xi\delta_3 + Y\delta_5)] \\
+ M [\lambda(\Xi\delta_1 + Y\delta_6) + \Lambda'(\Xi\delta_2 + Y\delta_4) + M(\Xi\delta_1 - Y\delta_6) + M'(\Xi\delta_2 - Y\delta_4)] \\
+ M' [\lambda(\Xi\delta_2 + Y\delta_4) + \Lambda'(\Xi\delta_3 + Y\delta_5) + M(\Xi\delta_2 - Y\delta_4) + M'(\Xi\delta_3 - Y\delta_5)]
\end{array} \right\},
\]

and

\[ \Lambda = \psi - k(\zeta'' + \chi') \]
\[ \Lambda' = \psi' - k(\zeta'' + \chi'') \]
\[ M = 1 - \chi - k\psi' \]
\[ M' = \zeta' - \chi' - k\psi''. \]
REFERENCES


