

Acoustic Shadow Detection: Study and Statistics of B-Mode and Radiofrequency Data

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Abstract

An acoustic shadow is an ultrasound artifact occurring at boundaries between significantly different tissue impedances, resulting in signal loss and a dark appearance. Shadow detection is important as shadows can identify anatomical features or obscure regions of interest. A study was performed to scan human subjects (N=37) specifically to explore the statistical characteristics of various shadows from different anatomy and with different transducers. Differences in shadow statistics were observed and used for shadow detection algorithms with a fitted Nakagami distribution on radiofrequency speckle (RF) or cumulative entropy on brightness-mode (B-mode) data. The fitted Nakagami parameter and Entropy values in shadows were consistent across different transducers and anatomy. Both algorithms utilized adaptive thresholding, needing only the transducer pulse length as an input parameter for easy utilization by different operators or equipment. Mean Dice coefficients

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(\pm standard deviation) of 0.90 ± 0.07 and 0.87 ± 0.08 were obtained for the RF and B-mode algorithms, which is within the range of manual annotators. The high accuracy in different imaging scenarios indicate that the shadows can be detected with high versatility and without expert configuration. The understanding of shadow statistics can be used for more specialized techniques to be developed for specific applications in the future, including pre-processing for machine learning and automatic interpretation.

Keywords: Acoustic Shadow, Ultrasound, Speckle, Radiofrequency, Segmentation

1 Introduction

2 Ultrasound devices have become increasingly affordable and portable, en-
3 couraging applications such as point-of-care ultrasound (Bouhemad et al.,
4 2011), novice usage (Sippel et al., 2011), and analysis by machine learning
5 (Ghose et al., 2013). However, ultrasound is susceptible to unique artifacts
6 that increase the difficulty of interpretation and processing of images. One
7 artifact is an acoustic shadow, which occurs when an ultrasound wave crosses
8 a boundary of two materials with high impedance differences (Kremkau and
9 Taylor, 1986). The wave is almost completely reflected and depicted beyond
10 the boundary is a continuous dark region and a loss of anatomical features.
11 Shadows occur in air-tissue, tissue-bone, and tissue-lesion interfaces. Shad-
12 ows can aid interpretation, such as identifying gall stones (Good et al., 1979)
13 or spinal levels (Galiano et al., 2005). However, shadows, such as from poor
14 transducer contact, can lead to misinterpretation of anatomy, particularly by
15 novice users and automated processing algorithms. Thus, the identification
16 of shadows is an important preprocessing step in many applications.

17 Several methods have been used in literature to detect shadows and il-
18 lustrative examples are discussed. Geometric techniques model the path of
19 an ultrasound signal for an expected image along the scanline using a ran-
20 dom walk (Karamalis et al., 2012). Pixels are then flagged as a shadow
21 if it is below a heuristic confidence threshold of 0.25. However, geometric
22 techniques require knowledge of ultrasound transducer properties to param-
23 eterize random walk weights, such as the focal length, radius of curvature,
24 and thickness. The technique is therefore challenging to implement across
25 different ultrasound equipment. This also reduces applicability for machine

26 learning applications as accurate transducer parameter labels are required
27 for each image.

28 Pixel gray level methods ignore the transducer properties and analyze
29 only the graphical properties of an image (Hellier et al., 2010). Shadows have
30 been detected on brain images by analyzing the entropy along a scanline to
31 flag pixels of sudden low entropy as a potential shadow. These techniques
32 achieved a comparable Dice similarity coefficient as geometric methods but
33 require specific thresholding, window sizing, filtering, and image mask pa-
34 rameterization for different anatomy and transducers. The drawback is again
35 the need for parameterization and tuning, which requires image processing
36 expertise and prior knowledge of specific applications.

37 Machine learning methods have gained significant interest in medical
38 imaging analysis. To our knowledge, no machine learning method has demon-
39 strated the capability of general shadow detection from multiple types of
40 anatomy. Deep learning methods have identified features in a specific im-
41 age sets that contain shadows, such as neuroanatomical regions in cranial
42 scan (Milletari et al., 2017) or spinal levels in a posterior scan (Hethering-
43 ton et al., 2017). Although machine learning has the potential of providing
44 automated feature recognition in multiple applications, a large data set is re-
45 quired for an algorithm to recognize certain features. Ultrasound imaging is
46 highly variable due to unique artifacts, operator techniques, and equipment.
47 In addition, shadows are a common feature that occur in various imaging
48 scenarios. Previous techniques focused on a single anatomical region and
49 training data was from a consistent imaging scenario. However, it is difficult
50 to construct a training data set with the generality required to recognize

51 shadows in different scenarios usable for a variety of ultrasound applications.

52 There are two objectives to this paper. First, to address the need for
53 understanding general characteristics of shadows, a study was conducted to
54 scan multiple anatomy and transducers specifically to analyze the statistics of
55 different types of shadows. Second, to address existing needs for versatile de-
56 tection with minimal parameterization, previous methods were then extended
57 utilizing statistical thresholding of radiofrequency (RF) or brightness-mode
58 (B-mode) data to detect shadows from various imaging scenarios. The two
59 methods are illustrated in a flowchart in Fig. 1.

60 **Materials and Methods**

61 *Data Collection*

62 Ultrasound RF and B-mode data were acquired by scanning 37 adult
63 participants with informed written consent, approved by the University of
64 British Columbia Research Ethics Board (Study ID: H18-01199). The scans
65 included a forearm scan near the distal end of the pronator quadratus, an
66 elbow scan near the cubital fossa, and a rib scan on the anterior surface of
67 right ribs 11-12. Each scan was taken with both a curvilinear (Model C5-
68 2/60, Ultrasonix Medical Corporation, Richmond, BC, Canada) and linear
69 (Model L14-5/38, Ultrasonix Medical Corporation, Richmond, BC, Canada)
70 transducer. Different transducer settings were used for each anatomical re-
71 gion and transducer, summarized in Table 1. Shadows were expected to
72 occur due to superficial and deep bones and from an air gap created by the
73 lateral edges of the transducer not being in flush contact with the skin. The
74 experiment was designed to generate a dataset from various imaging scenar-

ios to explore general shadow characteristics and to validate the versatility
of the two simple shadow detection methods. The pulse lengths measured
for the different transducers are reported in Table 1.

Radiofrequency Speckle Analysis

To analyze shadows, windows of speckle were analyzed on the RF signal. Speckle occurs from interference of randomly distributed microscopic scatterers, resulting in a granular appearance on the image. To produce B-mode images, manufacturers often employ image enhancement algorithms, such as logarithmic compression, nonlinearly alter speckle patterns. B-mode image formation can also be manipulated by an operator to visually enhance an image, such as adjusting time-gain compensation or dynamic range. Thus, the underlying speckle analysis in RF signals can provide shadow detection usable across different machines and operators. However, the original speckle pattern contains information of the acoustic interactions in tissue. (Burckhardt, 1978). By analyzing the RF signal distribution, we can statistically characterize the distributions in tissue compared to shadow regions. We expect tissue to resemble speckle modeled by known distributions and expect shadow to resemble different distributions, which may be a mixture of lessened speckle due to the signal loss and background electronic noise. Previous studies have attempted despeckling methods on images containing shadows (Aysal and Barner, 2007) by using filters based on a Rayleigh-like distribution. As such, even if shadow regions do not exactly resemble known speckle distributions, they may still be characterized to a sufficient extent with known distributions for a maximum likelihood fit. The fitted parameters can then be used to differentiate between shadow and non-shadow regions.

100 One of the first models for speckle is the one parameter Rayleigh distribu-
 101 tion to model the probability density of a random walk (Burckhardt, 1978).
 102 The Rayleigh distribution is capable of modeling fully developed speckle,
 103 which does not occur in limited scattering (Tuthill et al., 1988). More gen-
 104 eralized models have been applied such as the Rician, Homodyned-K, and
 105 Nakagami distributions to characterize speckle (Destripes and Cloutier,
 106 2010). The utility of speckle has been demonstrated in the literature to
 107 classify tumorigenicity of breast lesions (Byra et al., 2016) or levels of liver
 108 fibrosis (Ho et al., 2012) by categorizing image regions based on the speckle
 109 pattern. Shadow characterization presents a simpler problem as shadow and
 110 non-shadow regions contain significantly different speckle patterns. Thus,
 111 the Nakagami distribution expressed in Eq. 1 was chosen to model speckle.
 112 The Nakagami distribution provides greater generality than the Rayleigh
 113 distribution while being more computationally efficient than the Rician or
 114 Homodyned K distributions (Destripes and Cloutier, 2010):

$$\Phi(x, m, \omega) = 2\left(\frac{m}{\omega}\right)^m \frac{1}{\Gamma(m)} x^{(2m-1)} e^{-\frac{m}{\omega}x^2} \quad (1)$$

115 where x is RF intensity, m is the shape parameter or Nakagami m parameter,
 116 ω is a scale parameter and $\Gamma(m)$ is the gamma distribution.

117 To characterize shadows, the raw RF data was first processed by com-
 118 puting the echo envelope of each scanline with a Hilbert transform. This was
 119 performed on an averaged RF signal from three image frames. This creates
 120 a pre-scan converted image, visually similar to B-mode but without filtering
 121 to alter speckle. Next, the RF image was divided into overlapped windows
 122 with a width of a single RF scanline and a length of three times the pulse

length. We expect the width of a single RF scanline to be on the order of magnitude of a resolution cell, which is on the same order of magnitude as the correlation length (Wagner and Insana, 1988). The window length was demonstrated in literature to be sufficiently large to capture multiple wavelengths and scattering events while being small enough to be useful in differentiating different regions on the millimeter scale (Byra et al., 2016). Next, each window was fit to a Nakagami distribution using a maximum likelihood estimate to compute a map of Nakagami parameters m and ω , as shown in Fig. 2.

Then, for each ultrasound image, Otsus method was applied to its Nakagami ω map to automatically compute a ω threshold for each individual image as we expect separate distributions for shadow and non-shadow regions. This was sufficient as the ω parameter is significantly different for shadow regions with abundant speckle and non-shadow regions with minimal speckle. Then, for each scanline, the axially deepest data point that is above the threshold is labeled as the shadow boundary and all data points below are labeled as a shadow.

The Nakagami shape parameter, m , was also investigated, though there was not sufficient delineation between parameter values in shadow and non-shadow regions for this parameter to be effective in thresholding. The distributions of the two parameters are displayed for shadow and non-shadow regions in Fig. 5.

B-mode Scanline Analysis

Many ultrasound machines do not provide access to RF data for speckle analysis. Thus, a previous pixel gray level shadow detection method on B-

mode images was modified and extended. Scanline entropy was investigated on B-mode images to characterize different types of shadows, but with the addition of adaptive thresholding of entropy to address the need for usability with minimum configuration. B-mode analysis was performed on an averaged image from three image frames, similar to RF analysis. First, the cumulative scanline entropy is computed for each pixel, similar to the “Rupture Criterion” (Hellier et al., 2010), with approximate window size fixed as three times the pulse length, η , as defined in Eq. 2. This is the same window size as the RF analysis.

$$S_{i,j} = \sum_{k=1}^{3\eta} I(i-k, j) \log_2 \frac{I(i-k, j)}{I(i+k, j)} + I(i+k, j) \log_2 \frac{I(i+k, j)}{I(i-k, j)} \quad (2)$$

where $S_{i,j}$ is the cumulative entropy at pixel i on scanline j , η is the pulse length, and $I(i)$ is the gray level (0-255) of pixel (i, j) . For the case of curvilinear images, radial scanlines were linearly interpolated between the two symmetric lateral edges of the image.

Next, Otsu’s method is applied onto the entropy map of each image to automatically compute a threshold entropy value, similar to RF analysis. The intuition of the threshold is different than in RF analysis. In RF analysis, the threshold separates patches of intense and minimal speckle. In B-mode analysis, the threshold separates pixels of a shadow boundary, which has high entropy, and pixels away from shadow boundary, which include shadow and non-shadow regions. Thus, shadows can be identified by finding the last pixel on a scanline with an entropy higher than the threshold, representing a bright shadow boundary.

170 *Validation*

171 A trained annotator (RH) manually outlined the boundary of the shadow
172 regions on B-mode images. The manual regions were used as a gold standard,
173 as manual identification is common in clinical practice and has been used in
174 previous literature for comparison (Hellier et al., 2010). A Dice coefficient
175 was computed to compare similarity of manual and automated shadow de-
176 tection. The manual outline was used to define four regions for classification
177 of statistical parameters: a non-shadow region above the boundary, a shadow
178 region below the boundary, a “transition region”, which is a window defined
179 as three pulse lengths long axially below the boundary, and a “deep shadow
180 region”, which is the data below the transition region. The validation was
181 repeated with the RF and entropy window increased and decreased by 50%.
182 The Ljung-Box Q-test was used to measure residual autocorrelation of the
183 Dice coefficients. A Wilcoxon rank sum test has been performed between
184 Nakagami parameter values in shadow and non-shadow regions and between
185 entropy values in shadow and non-shadow regions.

186 As an initial experiment, a gelatin phantom was created with slits of
187 wood embedded at 0.75cm and at 2.50cm to create a region of shallow
188 and deeper shadows on both edges of the phantom. The gain was varied
189 and both RF and B-mode methods were employed to test the feasibility of
190 the methods on a clearly visible shadow, shown in Fig. 3. When comparing
191 to manual segmentation, all detected shadows resulted in a Dice coefficient
192 of above 0.95, with the lowest score being the entropy method applied on a
193 high-gain image. This provides support that extreme operator adjustments
194 on the B-mode image may affect pixel gray level detection methods more

195 than RF methods.

196 Results

197 Examples of detected shadows from both methods are highlighted in gray
198 in Fig. 4 in different shadow detection scenarios. The Dice coefficients for
199 both methods for different anatomy and transducers are shown in Table
200 2. The mean Dice coefficients (\pm standard deviation) were 0.90 ± 0.07 and
201 0.87 ± 0.08 for RF and B-mode methods. Manual annotation was repeated
202 five times with a mean Dice coefficient of 0.92 ± 0.02 for all images and trans-
203 ducers. The Dice coefficient did not change by more than 0.03 when the
204 window size was varied by 50%.

205 With the benefit of a varied dataset, general statistics of shadows can
206 be analyzed, as summarized in Table 3 and Table 4. The distributions of
207 Nakagami parameters and entropy for the different regions are visualized
208 in Fig. 5. For shadow detection, the parameters differentiating a shadow
209 and non-shadow are of particular interest. Shadows were observed to have a
210 mean Nakagami ω parameter of 4.14 ± 0.40 and a mean entropy of $1.03 \pm$
211 0.29 whereas non-shadows were observed to have a mean ω of 6.24 ± 0.92
212 and 2.20 ± 0.81 . Wilcoxon rank sum p values were less than 0.002 between
213 Nakagami ω parameter distributions in shadow and non-shadow regions and
214 less than 0.001 between entropy distributions in shadow and non-shadow
215 regions, indicating that shadow and non-shadow regions have statistically
216 different distributions for ω and entropy. The values of entropy and Nakagami
217 ω are consistent across different transducers and anatomical regions. The
218 variance of entropy and Nakagami ω in one imaging region and transducer

219 setting is less than the variance across different regions and transducers for
220 shadows and non-shadows.

221 Discussion

222 The RF and B-mode shadow detection developed achieved a comparable
223 Dice similarity coefficient to manual detection for all anatomy and transducer
224 types ($p < 0.025$). The previous studies using B-mode entropy reported a
225 mean Dice coefficient of 0.91 ± 0.07 between manual annotators (Hellier et al.,
226 2010). An important feature of shadow detection is being able to differentiate
227 between a shadow and simply high attenuation of the signal. Both scenarios
228 result in an eventual loss of signal. Shadow detection, however, has a char-
229 acteristic high gray level shadow boundary before a significant loss in signal,
230 compared to gradual signal losses in attenuation. The high Dice similarity
231 coefficient indicates that both methods were capable of this distinction. This
232 is also visualized in Fig. 4, where regions of low gray level without a bright
233 shadow boundary were correctly labeled as non-shadow. The high accuracy
234 supports the versatility of the detection method as both methods are able
235 to identify shadows across different anatomy and transducers with minimum
236 configuration.

237 For a general observation for shadows, the computed Nakagami ω param-
238 eters of all manually outlined shadows indicate that there is a statistically
239 significant difference between shadow and non-shadow regions, regardless of
240 anatomy and transducer and even with the error in the transition regions
241 considered. The speckle and its statistics from shadows is thus distinct from
242 the speckle created by tissue, muscle, or fat. This observation can be utilized

243 in the future for further analysis of shadows.

244 In RF detection, both false positive and false negative errors most fre-
245 quently occurred immediately below a shadow boundary as opposed to B-
246 mode detection where errors were in various regions. To study the frequent
247 areas of error further, the “transition region” immediately below a man-
248 ually annotated shadow boundary and a “deep shadow region” below the
249 transition region was investigated. The Nakagami ω parameter of transition
250 regions of all anatomy and transducers were within a standard deviation of
251 both shadow and non-shadow regions. The deeper shadow regions were ob-
252 served to have a lower Nakagami ω parameter than shadow regions and with
253 a lower standard deviation as summarized in Table 3. The spread of the
254 speckle also significantly decreases after the transition region. This indicates
255 that the transition region cannot be fully distinguished from either a shadow
256 or non-shadow and presents as it is statistically similar to the two. This
257 is likely the cause of the errors, as the speckle distribution is much more
258 consistent in the deep shadow regions compared to any other region. Phys-
259 ically, speckle interactions appear to gradually lessen after a brightest point
260 on a scanline, possibly due to incomplete total reflection at a boundary. The
261 boundary is thus is not an instantaneous division between non-shadow and
262 shadow, rather, there is a transition region with statistics between a shadow
263 and non-shadow before the speckle fully resembles a shadow.

264 In the transition region of B-mode images, the entropy values were similar
265 but consistently higher than non-shadow values. This is expected as entropy
266 is the highest when there is the greatest change in pixel gray level, which oc-
267 curs at a shadow boundary, even with the a non-instantaneous non-shadow

268 to shadow transition. However, the averaged entropy of all non-shadow re-
269 gions have a greater spread than the Nakagami parameters, likely due to
270 the differing operator settings used. Thus, B-mode detection may not be as
271 consistent as RF detection.

272 Limitations

273 In our study, although a range of frequencies and equipment were used,
274 the parameters were still limited and not all combinations were explored.
275 To further validate the detection method, future work would include a more
276 extensive investigation of these parameters, such as with a random parameter
277 grid search, to provide more support for widespread clinical use.

278 As both RF and B-mode images search for a threshold for the shadow
279 boundary, it is possible to misinterpret a reverberation artifact as a begin-
280 ning of a shadow. Reverberation at a shadow boundary would cause a similar
281 bright region followed by a dark region, which visually appears like a shadow
282 boundary despite being an artifact in a shadow region. This is a limitation in
283 our method and future work includes integration of reverberation identifica-
284 tion, such as identifying echo time duration to know what pulses correspond
285 to anatomical interaction (Win et al., 2010), would be required to reduce
286 reverberation errors.

287 There is a limitation with analysis using the Nakagami distribution in
288 that the fitted Nakagami distribution to model scatterers change depending
289 on transducer frequency. Previous literature observed that in the 36-58MHz
290 frequency range, the Nakagami m parameter decreased near the theoretical
291 lower limit compared to a higher Nakagami m parameter value at 10MHz

292 signal (Cloutier et al., 2004). This was reported to be due to the spatial
293 organization of the cells being "on the order of a fraction of the wavelength"
294 and a Nakagami distribution cannot model the scatterers of red blood cells
295 at this frequency. Due to this and from limitations of the equipment used in
296 our study, we cannot conclude that shadow detection with Nakagami anal-
297 ysis will be accurate in higher frequencies beyond the values tested. Future
298 studies are required to analysis the performace of shadow detection in higher
299 frequencies. Diagnostic ultrasound commonly uses a frequency range of 2-
300 15MHz (Jensen, 2007) and higher frequencies are limited to subspecialized
301 cases such as optical ultrasound (Pavlin et al., 1992). Shadow detection is ex-
302 pected be applicable in most use cases without issues from the high frequency
303 behaviour of the Nakagami distribution.

304 There is a limitation for diagnostic usage of the proposed shadow method
305 in cases where acoustic shadowing does not exhibit the characteristic bright
306 boundary followed by a dark region. In cases where there is partial or incom-
307 plete shadowing, such as small calcifications in the placenta (Abramowicz
308 and Sheiner, 2008). In these cases, there is a resemblance of a shadow, where
309 the calcification is brighter and the region below is noticeably darker, but
310 not with a brightness difference as extreme as shadowing from the ulna and
311 the regions below retain speckle similar to tissue. Although calcifications
312 are pathologically important to recognize, the proposed shadow detection
313 method would likely be unable to detect the partial shadowing from these
314 calcifications. The proposed method would be applicable only in cases of
315 more complete shadowing, which would still be practical for significant gall
316 and kidney stones, for instance.

317 In previous literature, shadows were defined qualitatively (Kremkau and
318 Taylor, 1986) as a sudden loss of signal and brightness. The observed transi-
319 tion region in this study suggests that the qualitative definition of a shadow
320 may be insufficient for accurate detection. One algorithm may detect the
321 shadow starting immediately after the brightest location, or another may
322 use a convention such as a full width at half maximum to define where the
323 signal has sufficiently low gray level to resemble the start of a shadow. There
324 is a decision point required for a clear definition for where a shadow begins
325 to improve shadow detection accuracy, both from a signaling perspective for
326 image processing and a visual perspective for manual inspection.

327 The findings in this study result in several implications. First, the statis-
328 tics of acoustic shadows have been investigated on a dataset with shadows
329 occurring from multiple scenarios as opposed to specific cases where shadows
330 are observed. This provided a more generalizable observation that shad-
331 ows can be characterized by distinctive speckle distributions in different of
332 anatomy and equipment and that there exists a transition region before the
333 loss of speckle in a shadow. Second, the shadow detection methods demon-
334 strated high accuracy, indicating that the same shadow detection method
335 can be used with different transducer or imaging location. In future stud-
336 ies, the speckle statistics observed can be used to develop further models for
337 anatomical features containing shadows. In machine learning algorithms, an
338 initial network could be used with the shadow detection methods presented.
339 Future studies would also have to take into consideration the most frequent
340 source of error of shadow detection as the shadow boundary.

341 **Conclusions**

342 Acoustic shadows from different imaging scenarios were investigated. RF
343 and B-mode methods were developed for acoustic shadow detection requiring
344 only the transducer pulse length as the input parameter. When comparing to
345 manual detection, the methods achieved a Dice similarity coefficient within
346 range of manual observers. The work focused on applying shadow detection
347 and statistical analysis to a varied dataset of three different anatomical loca-
348 tions and two different transducer to provide a representative understanding
349 of general acoustic shadows. The statistics of acoustic shadow indicate that
350 shadows contain a distinct speckle distribution compared to non-shadows and
351 the speckle characteristics transition at the shadow boundary. The statistical
352 findings of shadows can aid interpretation of ultrasound images in the future
353 using speckle analysis. The versatility of the shadow detection method has
354 the potential to improve the interpretation of ultrasound images with shadow
355 artifacts or to serve as a pre-processing step for machine learning methods.

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361 **References**

- 362 Abramowicz JS, Sheiner E. Ultrasound of the Placenta: A Systematic Ap-
363 proach. Part I: Imaging. *Placenta*, 2008;29:225–240.
- 364 Aysal TC, Barner KE. Rayleigh-maximum-likelihood filtering for speckle re-
365 duction of ultrasound images. *IEEE Trans. Med. Imaging*, 2007;26:712–
366 727.
- 367 Bouhemad B, Brisson H, Le-Guen M, Arbelot C, Lu Q, Rouby JJ. Bedside
368 ultrasound assessment of positive end-expiratory pressure-induced lung re-
369 cruitment. *American Journal of Respiratory and Critical Care Medicine*,
370 2011;183:341–347.
- 371 Burckhardt CB. Speckle in ultrasound B-mode scans, 1978.
- 372 Byra M, Nowicki A, Wróblewska-Piotrkowska H, Dobruch-Sobczak K. Clas-
373 sification of breast lesions using segmented quantitative ultrasound maps
374 of homodyned K distribution parameters. *Med. Phys.*, 2016;43:5561–5569.
- 375 Cloutier G, Savery MDD, Garcia D, Durand LG, Foster SF. Non-Gaussian
376 statistics and temporal variations of the ultrasound signal backscattered
377 by blood at frequencies between 10 and 58 MHz. *Acoust. Soc. Am.*,
378 2004;116:566–577.
- 379 Destrepes F, Cloutier G. A critical review and uniformized representation of
380 statistical distributions modeling the ultrasound echo envelope. *Ultrasound*
381 *Med. Biol.*, 2010;36:1037–1051.

382 Galiano K, Obwegeser AA, Bodner G, Freund M, Maurer H, Kamelger FS,
383 Schatzer R, Ploner F. Ultrasound guidance for facet joint injections in
384 the lumbar spine: A computed tomography-controlled feasibility study.
385 *Anesthesia and Analgesia*, 2005;101:579–583.

386 Ghose S, Oliver A, Mitra J, Martí R, Lladó X, Freixenet J, Sidibé D, Vilanova
387 JC, Comet J, Meriaudeau F. A supervised learning framework of statis-
388 tical shape and probability priors for automatic prostate segmentation in
389 ultrasound images. *Medical Image Analysis*, 2013;17:587–600.

390 Good LI, Edell SL, Soloway RD, Trotman BW, Mulhern C, Arger Pa. Ultra-
391 sonic properties of gallstones. Effect of stone size and composition. *Gas-
392 troenterology*, 1979;77:258–263.

393 Hellier P, Coupé P, Morandi X, Collins DL. An automatic geometrical and
394 statistical method to detect acoustic shadows in intraoperative ultrasound
395 brain images. *Medical Image Analysis*, 2010;14:195–204.

396 Hetherington J, Lessoway V, Gunka V, Abolmaesumi P, Rohling R. SLIDE:
397 automatic spine level identification system using a deep convolutional neu-
398 ral network. *International Journal of Computer Assisted Radiology and
399 Surgery*, 2017;12:1189–1198.

400 Ho MC, Lin JJ, Shu YC, Chen CN, Chang KJ, Chang CC, Tsui PH. Using
401 ultrasound Nakagami imaging to assess liver fibrosis in rats. *Ultrasonics*,
402 2012;52:215–222.

403 Jensen JA. Medical ultrasound imaging. *Prog. Biophys. Mol. Biol.*,
404 2007;93:153–165.

405 Karamalis A, Wein W, Klein T, Navab N. Ultrasound confidence maps using
406 random walks. *Medical Image Analysis*, 2012;16:1101–1112.

407 Kremkau FW, Taylor KJ. Artifacts in ultrasound imaging. *Journal of Ultra-*
408 *sound in Medicine*, 1986;5:227–237.

409 Milletari F, Ahmadi SA, Kroll C, Plate A, Rozanski V, Maiostre J, Levin
410 J, Dietrich O, Ertl-Wagner B, Bötzel K, Navab N. Hough-CNN: Deep
411 learning for segmentation of deep brain regions in MRI and ultrasound.
412 *Computer Vision and Image Understanding*, 2017;164:92–102.

413 Pavlin CJ, Ritch R, Foster FS. Ultrasound biomicroscopy in plateau iris
414 syndrome. *American Journal of Ophthalmology*, 1992;113:390–395.

415 Sippel S, Muruganandan K, Levine A, Shah S. Review article: Use of ultra-
416 sound in the developing world. *Int. J. Emerg. Med.*, 2011;4:72.

417 Tuthill TA, Sperry RH, Parker KJ. Deviations from rayleigh statistics in
418 ultrasonic speckle. *Ultrasonic Imaging*, 1988;10:81–89.

419 Wagner RF, Insana MF. Fundamental Correlation Lengths of Coherent
420 Speckle in Medical Ultrasonic Images. *IEEE Trans Ultrason Ferroelectr*
421 *Freq Control*, 1988;35:34–44.

422 Win KK, Wang J, Zhang C, Yang R. Identification and removal of rever-
423 beration in ultrasound imaging. In: *Proceedings of the 2010 5th IEEE*
424 *Conference on Industrial Electronics and Applications, ICIEA 2010*, 2010.

425 **Figure Captions**

426 **Figure 1:** Processing steps for radiofrequency (RF) and B-mode shadow
427 detection. RF processing is used if RF data is available and involves
428 fitting the Nakagami distribution onto the echo envelope of each RF
429 scanline before adaptive thresholding with Otsus method. In many
430 cases, there may only be access to B-mode image data, for which an
431 entropy map is computed and similar adaptive thresholding is used to
432 detect shadows.

433 **Figure 2:** A visualization of the B-mode and RF parameter maps. The b)
434 Entropy Map was computed from processing of the a) original B-mode
435 image and the d) Nakagami ω map was computed from the c) echo
436 envelope. Note that the echo envelope contains noticeable speckle,
437 which has been used to fit a Nakagami distribution to characterize
438 shadow. The region at depth 2.50 cm and scanlines 32-40 is attenuation
439 and not a shadow. This is an important distinction in shadow detection
440 and both maps show the region as below a threshold to flag a shadow
441 boundary.

442 **Figure 3:** Images of both RF and B-mode shadow detection performed on a
443 gelatin phantom with two wooden slits embedded at a depth of 0.75cm
444 and 2.50cm. The phantom was made to simulate shallow, deep, and
445 non-shadow regions. The methods were capable of shadow detection
446 with a high accuracy (Dice coefficient ≥ 0.95), though noticeable errors
447 were present at high-gain images for the B-mode method. This is ex-
448 pected as B-mode methods rely on pixel gray level, which may vary

449 due to operator settings.

450 **Figure 4:** A comparison of the original B-mode images, the detected shad-
451 ows manual detection, RF detection, and B-mode detection. Both
452 detection methods perform similarly to manual detection. Both meth-
453 ods perform slightly less accurately on curvilinear images, likely due
454 to the reduced resolution from interpolating the scanlines. Most errors
455 of RF detection occur near the shadow boundary, likely due to the
456 transitioning speckle from non-shadow to shadow.

457 **Figure 5:** Histograms of Nakagami parameters and entropy values in shadow
458 and non-shadowing regions. The Nakagami ω and Entropy distri-
459 butions have a more noticeable delineation between shadowing and
460 non-shadowing distributions compared to the Nakagami m parameter,
461 which was not used to threshold shadow boundaries. Entropy is very
462 minimal in continuous dark shadow regions, which is expected due to
463 the minimal variations in pixel gray level.

464 **Tables**

465 **Table 1:** Transducer properties for different imaging scenarios.

466

	Anatomy	Frequency	Depth	Gain	Pulse Length
Linear Transducer (L14-5/38)	Forearm	11.0MHz	5.0cm	50%	0.6mm
	Elbow	11.0MHz	5.0cm	40%	0.6mm
	Ribcage	5.0MHz	10.0cm	30%	1.7mm
Curvilinear	Transducer(C5-2/60)	4.0MHz	5.0cm	50%	2.6mm
	Elbow	4.0MHz	5.0cm	40%	2.6mm
	Ribcage	3.3MHz	10.0cm	30%	5.5mm

467 **Table 2:** Mean Dice coefficients for different imaging scenarios \pm standard
468 deviation.

		RF	B-Mode
Linear (L14-5/38)	Forearm	0.91 \pm 0.05	0.89 \pm 0.06
	Elbow	0.94 \pm 0.06	0.90 \pm 0.07
	Ribcage	0.87 \pm 0.09	0.84 \pm 0.06
Curvilinear (C5-2/60)	Forearm	0.89 \pm 0.05	0.86 \pm 0.08
	Elbow	0.93 \pm 0.04	0.90 \pm 0.09
	Ribcage	0.83 \pm 0.08	0.83 \pm 0.10
Mean	All Anatomy	0.90\pm0.07	0.87\pm0.08

469 **Table 3 :** The mean Nakagami ω and Entropy values of different anatomy,

transducer, and shadowing region \pm standard deviation. Values are consistent among different transducers and anatomical regions. The variance of entropy and Nakagami ω in one imaging region and transducer setting is less than the variance across different regions and transducers for shadows and non-shadows.

	Linear (L14-5/38)			Curvilinear (C5-2/60)		
	Forearm	Elbow	Ribcage	Forearm	Elbow	Ribcage
Nakagami ω (Log Scale)						
Shadow	4.15 \pm 0.45	4.18 \pm 0.45	4.04 \pm 0.42	4.22 \pm 0.32	4.19 \pm 0.40	4.08 \pm 0.37
Non-Shadow	6.19 \pm 0.96	6.49 \pm 0.97	6.29 \pm 0.95	6.54 \pm 0.88	6.29 \pm 1.04	5.64 \pm 0.71
Transition	4.94 \pm 0.62	5.36 \pm 0.62	4.96 \pm 0.38	5.26 \pm 1.02	5.37 \pm 0.99	4.59 \pm 0.92
Deep Shadow	4.13 \pm 0.43	4.16 \pm 0.43	4.03 \pm 0.41	3.93 \pm 0.20	4.09 \pm 0.30	4.03 \pm 0.26
Entropy (Log Scale)						
Shadow	0.92 \pm 0.22	1.10 \pm 0.36	1.04 \pm 0.27	1.06 \pm 0.28	0.96 \pm 0.21	1.10 \pm 0.37
Non-Shadow	2.34 \pm 0.96	2.34 \pm 0.80	2.14 \pm 0.82	1.67 \pm 0.82	1.75 \pm 1.14	1.88 \pm 0.42
Transition	2.45 \pm 0.62	2.56 \pm 0.53	2.15 \pm 0.51	2.18 \pm 1.21	1.93 \pm 1.10	1.99 \pm 1.10
Deep Shadow	0.71 \pm 0.43	0.89 \pm 0.26	0.92 \pm 0.40	0.98 \pm 0.21	0.82 \pm 0.19	1.04 \pm 0.26

Table 4 : The mean Nakagami ω and Entropy values of all anatomy and transducers for different shadowing regions \pm standard deviation.

	Mean Nakagami ω (Log Scale)	Mean Entropy (Log Scale)
Shadow	4.14 ± 0.40	1.03 ± 0.29
Non-Shadow	6.24 ± 0.92	2.02 ± 0.81
Transition	5.08 ± 0.77	2.21 ± 0.84
Deep Shadow	4.06 ± 0.34	0.89 ± 0.27