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# On the relationship between dynamic contrast enhanced ultrasound parameters and the underlying vascular architecture extracted from acoustic angiography

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# Abstract

Dynamic contrast-enhanced ultrasound (DCE-US) has been proposed as a powerful tool for cancer diagnosis by estimation of perfusion and dispersion parameters reflecting angiogenic vascular changes. This work aims at identifying which vascular features are mainly reflected by the estimated perfusion and dispersion parameters through comparison with Acoustic Angiography (AA). AA is a high resolution technique that allows quantification of vascular morphology. 3D AA and 2D DCE-US bolus acquisitions monitored growth of fibrosarcoma tumors in 9 rats. AA-derived vascular properties were analyzed

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along with DCE-US perfusion and dispersion in order to investigate the differences between tumor and control, and their evolution in time. AA-derived microvascular density and DCE-US perfusion showed good agreement, confirmed by their spatial distributions. No vascular feature was correlated with dispersion. Yet, dispersion provided better cancer classification than perfusion. We therefore hypothesize that dispersion characterizes vessels that are smaller than those visible with AA.

*Keywords:* Acoustic angiography, Dynamic contrast-enhanced ultrasound, Cancer, Dispersion, Perfusion, Ultrasound contrast agents

## 1 Introduction

Malignant tissue shows a set of alterations from being tissue that can be 2 used as markers to detect it (Koumoutsakos et al., 2013). Of particular in-3 terest for cancer imaging are the altered vascular architecture and the conse-4 quent changes in blood supply. Angiogenic vessels grow to nourish the tumor 5 and support its proliferation. These vessels have been found to be tortuous, 6 to grow chaotically, without the typical vessel hierarchy, and with a high occurrence of arteriovenus shunts. Many of these properties can be recognized 8 with contrast-enhanced ultrasound techniques, which have shown promising 9 results for distinguishing malignant tissue from benign (Brock et al., 2013; 10 Gessner et al., 2013; Kuenen et al., 2013b, 2011; Mischi et al., 2012; Quaia, 11 2011; Shelton et al., 2015). 12

Dynamic contrast-enhanced ultrasound (DCE-US) captures the contrast-13 agent passage through the vascular bed after its injection in the patient's 14 bloodstream. Specifically, it registers the local evolution of gray-level inten-15 sity at each pixel, referred to as the time intensity curve (TIC), which reflects 16 the varying ultrasound contrast agent (UCA) concentration. The recorded 17 intensities are then converted into UCA concentration with a linearization 18 function specific to the employed ultrasound scanner (Rognin et al., 2008), 19 yielding an indicator dilution curve (IDC) for every pixel in the video. Vari-20 ous characteristics of IDCs have been proven to be useful for distinguishing 21 malignant from benign tissue (Mischi et al., 2012). 22

Several approaches have been adopted to extract information from IDCs
derived from DCE-US bolus acquisitions. Some heuristic features of the
IDCs, such as the wash-in time and the peak intensity, are related to cancer

(Mischi et al., 2012; Zhao et al., 2010). Multiple other techniques employ 26 IDC fitting by analytical models, such as the lognormal, gamma, and local 27 density random walk (LDRW) model (Strouthos et al., 2010). Functional 28 parameters of the curves (e.g. area-under-the-curve) are extracted and dis-29 played in colormaps, aiming to obtain a clearly distinguishable malignant 30 region. All these approaches mainly attempt to quantify perfusion, which 31 is motivated by the presence of ample arteries feeding the tumor, increased 32 microvacular density (MVD), and presence of arteriovenous shunts. Despite 33 this, clinical evidence has shown that cancerous lesions in the prostate can 34 also be iso- or hypo-perfused (Brock et al., 2013). Indeed, it is known that 35 tumor tissue has higher resistance to blood flow (Narang and Varia, 2011). 36 This induces a couterbalancing factor that complicates predictions about 37 the level of blood supply within the tumor, as compared to surrounding tis-38 sue (Cosgrove and Lassau, 2010). Furthermore, the MVD inside the tumor 30 can be strongly heterogeneous, creating highly perfused regions as well as 40 hypoxic, avascular regions. Therefore, assessment of perfusion alone is in-41 sufficient for reliable cancer diagnostics. These findings have motivated the 42 development of contrast ultrasound dispersion imaging (CUDI), a method 43 which enables assessment of UCA dispersion, in addition to quantification of 44 perfusion (Kuenen et al., 2011; Mischi et al., 2012). 45

CUDI aims at quantifying the UCA dispersion due to the architecture of the vascular tree and complex multipath trajectories available for UCA transport. The main hypothesis that lies in the foundation of the method states that dispersion reflects structural vascular changes induced by angiogenesis. The first CUDI approach involved modelling of the IDCs in time

domain with a LDRW model and extraction of a dispersion-related parame-51 ter from the fitted model (Kuenen et al., 2011). An important complication 52 associated with this approach was poor signal to noise ratio, hindering the 53 fitting procedure and decreasing its reliability. This problem has been mit-54 igated by spatiotemporal similarity analysis (Kuenen et al., 2013a; Mischi 55 et al., 2012). In a promising implementation, this approach involves calcu-56 lation of an average correlation coefficient measuring the similarity of a TIC 57 at a pixel and its surrounding pixels (Kuenen et al., 2013b). A theoretical 58 description of the problem within the framework of the LDRW model has 59 shown that the correlation coefficient between IDCs is monotonically related 60 to the dispersion coefficient (Kuenen et al., 2013a). Moreover, this approach 61 has demonstrated its superior performance compared to perfusion-related 62 parameters at localizing prostate cancer in a clinical setting (Kuenen et al., 63 2013b). This method has been validated against cell differentiation reflected 64 with the Gleason score for prostate cancer (Schalk, 2017). Another study 65 identified that regions of low dispersion correlated with those of high MVD, 66 quantified by immunohistology (Saidov et al., 2016). However, in this study 67 detailed chatacterization of the vascular architecture (e.g. tortuosity and 68 vessel size) was not available. 69

Acoustic angiography (AA) can provide accurate characterization of the vascular architecture: it is a high-resolution technique, capable of imaging individual microvessels (Gessner et al., 2013; Shelton et al., 2015). AA permits imaging vessels at a high resolution of 100-200 µm at 2 cm depth with minimal signal from tissue. While transmitting ultrasonic waves at frequencies in the order of a few MHz, close to the UCA bubble resonance frequency,

it records the nonlinear response of the contrast agents in a high frequency 76 range centered at 30 MHz. This technique grants the possibility to quantify 77 vessel density and morphology measures such as the sum of angles metric 78 (SOAM) and distance metric (DM) (Rao et al., 2016; Shelton et al., 2015). 79 These parameters have been reported to be significantly different for malig-80 nant and benign tissue (Gessner et al., 2013; Shelton et al., 2015). Thereby, 81 AA gives the opportunity to validate whether these features are reflected in 82 DCE-US due to the different character of UCA perfusion and dispersion in 83 these vessels. 84

The aim of this work is to determine whether DCE-US is able to char-85 acterize the underlying vascular architecture. It involves DCE-US and AA 86 imaging of fibrosarcoma tumors and control regions in a longitudinal study of 87 9 rats. AA and DCE-US acquisitions were performed every 3 days, at 4 time 88 points, starting with the day when the tumors could be palpated. An overall 80 comparison of the tumor's and control's vascular properties was performed. 90 Additionally, a longitudinal study of these properties was conducted, aiming 91 to find similar trends in features extracted from the two different techniques 92 of DCE-US and AA. 93

# 94 Materials and Methods

# 95 Rat Models

Fibrosarcoma tumor implantation was performed in rats according to a previously applied protocol (Streeter et al., 2011). The tumor models were established from propagated tumor tissue provided by the Dewhirst Lab at Duke University. Before surgery the (Fischer 344) rats were anesthetized with isoflurane; their left flank was then shaved and disinfected. An incision ( $\sim 2 \text{ mm}$ ) was made above the quadriceps muscle, and a sample of tumor tissue ( $\sim 1 \text{ mm}^3$ ) was positioned under the skin. The incision was closed with 1-2 staples. This procedure was performed at 3 different time points with 9 rats in total. Rats belonging to the same series were operated on the same day.

On day 8 after implantation, the first ultrasound acquisition was per-106 formed if the tumors were palpable. Otherwise, we waited for 2-3 days for 107 subsequent assessment. When the tumors were palpable, UCA was injected 108 in the rats' tail vein through a 24 gauge catheter while the animals were anes-109 thetized with vaporized isoflurane in oxygen. DCE-US was performed on the 110 tumor-bearing flank for assessment of perfusion and dispersion. The AA ac-111 quisition protocol immediately followed the DCE-US acquisition to minimize 112 the amount of time each animal spent under anesthesia. The beginning of 113 the DCE-US and AA acquisitions were different between the series, start-114 ing with day 8, day 11, and day 13, respectively. For all but one animal, 115 subsequent imaging acquisitions were performed with an interval of 3 days, 116 amounting to 4 time points in total. One rat was an exception since we 117 were not able to inject the contrast (for both modalities) in its tail vein, and 118 managed to image only at the first and third time points. All experiments 119 were performed at the University of North Carolina at Chapel Hill, approved 120 by the Institutional Animal Care and Use Committee at the University of 121 North Carolina at Chapel Hill. 122

#### 123 Image acquisition

# 124 DCE-US bolus injection protocol

A UCA bolus of  $2 \times 10^8$  microbubbles was injected in the rats' tail vein. The contrast agent used in this study was made in-house; it has a lipid shell and perfluorcarbon core, similar to Definity<sup>®</sup> (Latheus Medical Imaging/U.S.A, N. Billerica). A 15L8-S probe was utilized with a Siemens Sequoia scanner in Cadence Pulse Sequencing mode at an insonifying central frequency of 7 MHz. The acquired DCE-US recordings were stored in DI-COM format.

#### <sup>132</sup> AA continuous infusion protocol

A continuous infusion of microbubbles was administered using a syringe pump (PHD 2000, Harvard Apparatus) at a rate of  $1.5 \times 10^8$  microbubbles per minute. AA imaging was performed with a dual-frequency single-element transducer transmitting at 4 MHz, and receiving around 30 MHz. The 3D AA images were acquired plane by plane, with a step size of 100 µm.

# <sup>138</sup> DCE-US bolus data processing

#### 139 Preprocessing

All the bolus recordings were filtered with a Gaussian filter, as previously performed in (Mischi et al., 2012), using a kernel of 0.13 mm equal to 1.6 pixels. This value improved the signal-to-noise-ratio at the cost of additional spatial correlation between TICs at neighbouring pixels. The TIC power of every pixel was evaluated as the root mean square of the TIC after the baseline was removed. Regions with a level of TIC power below -22 dBs of the maximum TIC power over all images were excluded from further analysis

(shown in black in Fig. 1 a.). This limited the effect of random noise on the 147 parameters of interest (Kuenen et al., 2014). Characteristic of DCE-US is 148 multiplicative noise: noise proportional in its power to the signal amplitude. 149 By eliminating regions with low TIC power, we avoided erroneous parameter 150 estimation from regions with low signal power where random noise dominates. 151 After this, the intensity values of the remaining regions were linearized by 152 inverting the logarithmic compression function implemented in the adopted 153 scanner, yielding the IDCs. 154

#### 155 Assessment of dispersion

An average correlation coefficient was calculated for every pixel between 156 its own IDC and those at its surrounding pixels within a ring-shaped kernel 157 (Mischi et al., 2012) with an inner radius of 0.6 mm and an outer radius 158 of 2 mm. The inner radius was chosen equal to the lateral resolution of 159 the preprocessed bolus recordings at  $\sim 2$  cm depth as identified with local 160 autocorrelation analysis. Details about the latter procedure can be found in 161 (Mischi et al., 2012). The lateral resolution was taken as a reference since 162 it was worse than the axial resolution. The outer radius of the kernel was 163 set equal to the size of 2 mm, which a tumor can usually reach without 164 neovascularization (Folkman, 1971). The time window over which the IDCs 165 were correlated to each other was selected to maximize the area under the 166 receiver operating characteristic curve for tumor classification, resulting in a 167 value of 17 seconds as proposed in previous work (Panfilova et al., 2016). This 168 is the only informative segment of the IDC (Fig. 2) due to early recirculation, 169 as often observed in small animals (Stapleton et al., 2009). In this work, the 170 beginning of the analyzed time window was set with 3 seconds before the 171

<sup>172</sup> appearance time, ensuring the wash-in phase to be entirely captured.

# 173 Assessment of perfusion

Wash-in-rate was adopted to assess perfusion and computed as the slope of a line fitted to the IDC in the 2-second interval after appearance time, as illustrated in Fig. 2. The value of 2 seconds was chosen to reflect the rise of UCA concentration in the initial part of the IDCs in all acquired clips.

178 AA data processing

The AA volumes were interpolated to reduce the inter-plane distance to 179 50 microns and make the pixels isotropic. Visible vessels were manually seg-180 mented and characterized in terms of vessel dimensions: vessel length (VL) 181 and mean radius (MR). VL was computed as the length of the vessel segment 182 identified between successive branching points, and MR was computed as the 183 mean radius of this vessel segment along its length. Vessel tortuosity was as-184 sessed with the distance metric and the sum of angles metric (Bullitt et al., 185 2003). The DM was computed as the ratio of vessel length to the Eucledian 186 distance between its beginning and end. The SOAM was calculated as the 187 sum of angles between successive points on the vessel centerline divided by 188 VL, using the same formula as in (Bullitt et al., 2003), but excluding the 180 torsional angle. Besides these individual vessel properties, MVD was calcu-190 lated as a global characteristic of the tumor at a given timepoint, defined as 191 the number of visible vessel segments divided by tumor volume. The volume 192 vascular density (VVD) was computed with a moving 3D isotropic kernel in 193 the central slice of the tumor ( $\sim 1 \text{ mm}$  in thickness). Otsu's method (Vala 194 and Baxi, 2013) was used to select a threshold to separate noise from vessel 195

<sup>196</sup> signal within the central slices; the percentage of pixels with vessel signal
<sup>197</sup> from the overall number of pixels in the 3D kernel was calculated.

#### 198 Statistical analysis

The DCE-US parameters were spatially downsampled by a factor 7 in both directions, equal to the resolution of the preprocessed images. This was performed to exclude spatial correlation and prepare the data for the statistical tests that require sample independence.

#### <sup>203</sup> Comparison between tumor and control

Dispersion and perfusion values were divided into two groups. The tumor 204 group was composed of the manually selected tumor regions (inside the red 205 contour, Fig. 1 a.) from all rats at all time points binned together. The 206 control group was taken from pixels outside the tumor contour, dilated by 207  $\sim 1 \text{ mm}$  (in blue, Fig. 1 a.). The region between the red and blue contour 208 was excluded from analysis to avoid erroneous pixel assignment to tumor 209 or control, since DCE-US information was not considered sufficiently com-210 prehensive for such accurate tumor delineation, as required by e.g. ablation 211 therapy and surgery. The AA parameters were extracted in a similar fashion: 212 vessels were taken from within the tumor region and outside it in the same 213 flank (Fig. 3). Vessel segments on the border of the selected contour, whose 214 belonging to a tumor or control group was debatable were disregarded from 215 analysis. 216

An Anderson-Darling goodness of fit hypothesis test was performed on all the parameter distributions to check for data normality. Since all the distributions were identified as non-Gaussian, a Mann Whitney non-parametric test was performed to establish the significance (p-value) of the difference between tumor and control. No additional subsampling or upsampling was performed to make the control and tumor data sets balanced, since the Mann Whitney test can be applied to data sets with distributions of different size (Mann and Whitney, 1947).

The Cohen's d was used as a measure of the 'effect size' (Sullivan and Feinn, 2012) that the tumor has on the underlying vasculature, calculated as the difference between the means of two distributions divided by the standard deviation of the control. The values of the Cohen's d term allow to classify the difference between two distributions according to 4 categories: small, medium, large, and very large for values of 0.2, 0.5, 0.8, 1.3, respectively.

# <sup>231</sup> Longitudinal study of tumor and control

A longitudinal study of the tumor evolution was performed with the 232 Kruskall-Wallis post hoc test, evaluating the differences among the distribu-233 tions of dispersion and perfusion, and vascular features of tumor and control 234 at 4 time points. The Kruskall-Wallis test (Kruskall and Wallis, 1952) does 235 not require equal sample sizes, which is an advantage considering that our 236 data set is unbalanced and incomplete: data is missing for one tumor at two 237 time points as well as control at several time points for the large tumors. 238 Moreover, the number of visible vessels is different for every image acquisi-239 tion. For all rats, all parameter values were binned together according to the 240 time point of the acquisition. 241

The statistic test calculation is influenced by the number of observations and can result in different outcomes for different sample sizes (Kruskall and Wallis, 1952). Since the number of pixels provided more samples for dispersion and perfusion compared to the number of vessels extracted with AA, these pixels were randomly subsampled to yield the same number of samples as vessels per each representative dataset of tumor and control at each time point. The only parameter that remained different in terms of group size is the MVD; being a global parameter that characterizes the entire tumor and control at a specific time point.

After the post-hoc tests were performed, the Pearson correlation coefficient was computed between the medians of the parameters showing similar longitudinal trends.

#### <sup>254</sup> Mapping of vascular properties on the bolus acquisition plane

During the DCE-US bolus acquisistions the operator always tried to im-255 age the largest cross-section of the tumor, and keep the same orientation of 256 the probe as used for AA. However, it was noticed that these precautions 257 were not sufficient to reliably identify the DCE-US plane within AA: even a 258 movement of the order of  $\sim 1 \text{ mm}$  alters the imaged vascular pattern of a tu-259 mor. It was noticed that the perfusion maps highlight larger vessels, clearly 260 visible in the AA (Fig. 2 b. and d.). These vessels were used as markers 261 to locate the bolus recording plane in the AA volume. For this, a dedicated 262 tool was developed, allowing to freely scroll through the AA volume planes 263 and change their orientation. 264

The selection of the plane was performed by visual inspection, choosing an image containing as many as possible vessel markers present in the perfusion maps. A slice in the AA volume of  $\sim 1$  mm thickness was selected and an extention of the skeletonization algorithm described in (Meiburger et al., 2006) was applied to extract MVD (Fig. 2 e.), MR (Fig. 2 f.), VL, and SOAM. This slice thickness was chosen to be of the order of the elevational resolution in the bolus recordings and sufficiently large to register vessel segments. This allowed a qualitative comparison of the spatial distribution of the vascular features with those of dispersion and perfusion in the same plane.

All the image processing and statistical analysis was performed with Matlab software (the MathWorks, Natick, MA).

# 277 **Results**

278 Statistical analysis

#### 279 Comparison between tumor and control

For all the extracted parameters, tumor and control have significantly different distributions (p < 0.001). However, the magnitude of the differences, expressed in Cohen's d, spans a wide range (Fig. 4), showing a marginal effect size for the DM (Fig. 4 c.), and small to very large differences for the rest of the parameters.

#### 285 Longitudinal study of tumor and control

Since the DM showed almost no difference between tumor and control, it was excluded from the longitudinal analysis. Boxplots with all parameter values binned according to the time points are shown in Fig. 5, while Fig. 6 illustrates the results of the post hoc Kruskall-Wallis test, color-coded according to the significance level of the intra-distribution differences.

The dispersion median is relatively constant in time for both tumor and control, showing a significant difference for control and tumor distributions (Fig. 5 a., Fig. 6 a.). Tumor perfusion is significantly different from the control at all time points (Fig. 5 b., Fig. 6 b.), peaking for the tumors at the second time point. Interestingly, the longitudinal trend of the control's perfusion seems to mimic the tumor's trend in time, however, at a smaller magnitude, not identified as significant with the post hoc test.

The VVD is stably higher for tumor, while the MVD seems to follow a similar trend to that of perfusion, peaking for tumors at the second time point. However, the result of the MVD post hoc test is difficult to compare to others since the number of samples is different: only one value of MVD per time point is available, while the other parameters were subsampled according to the number of segmented vessels in the AA volume at a given time point.

The post hoc results, illustrated by colormaps in Fig. 6, are comparable for dispersion, the VVD, the VL, and the SOAM. However, no significant correlation between the medians of the dispersion levels and the mentioned AA parameters has been identified. As for perfusion, the mean perfusion in tumors and their MVD showed a significant correlation coefficient of 0.572 (p < 0.001) and inclusion of both control and tumor values resulted in a Pearson correlation coefficient of 0.67 (p < 0.001).

## <sup>312</sup> Mapping of vascular properties on the bolus acquisition plane

The spatial parametric maps of the AA skeleton confirmed our observation that there is a correlation between regions of high perfusion and elevated MVD (Fig. 1 b. and f.). No spatial correspondance was found between dispersion and the other AA - derived parameters.

# 317 Discussion

Dispersion shows a large difference (Cohen's d = 1.68) between tumor 318 and control, exhibiting stable performance at tumor detection as it develops. 319 Perfusion shows a lower discrimination power than dispersion, that is high for 320 younger tumors, peaking at time point 2, and decreases with tumor growth. 321 Interestingly, the perfusion level in the control around the tumor is also 322 elevated (Fig. 5 b.), showing a similar trend as in the tumor itself. This 323 may reflect that the overall perfusion of tissue around the tumor is increased 324 and influenced by the tumor. This effect has been shown for the SOAM, 325 which exhibits intermediate values between that of tumor and control in 326 tissue adjacent to the tumor (Rao et al., 2016). Moreover, it has been shown 327 for the fibrosarcoma model that the vascular source is often located in the 328 periphery of the tumor (Ponce et al., 2007; Tozer et al., 1990; Viglianti et al., 329 2004). 330

Dispersion of the control stays stable over time, indicating that dispersionrelated changes mainly occur within the tumor itself, and not in the surroundings. The spatial perfusion and dispersion maps are complementary, showing different patterns of highlighted regions (Fig. 1 b. and c.). Perfusion highlights large vessels, as well as regions with high MVD.

The SOAM indicates that the tumor has more tortuous vessels, exhibiting a similar trend to that of dispersion (Fig. 5 a., g.) and comparable results for the post-hoc test (Fig. 6 a., g.). Nevertheless, the effect size difference, as indicated by Cohen's d, is much lower for the SOAM than for dispersion. In general, the control regions in this experiment show a higher tortuosity than we previously observed for these rats, expressed by the DM in (Shelton et al.,

2015). Direct comparison of the SOAM in this work and in (Shelton et al., 342 2015) is not available since the calculation of the SOAM has been adjusted 343 since then. The unusually high tortuosity for control may be caused by the 344 presence of the bowel region in some of the AA images, which was excluded 345 from analysis in earlier studies, and may have elevated tortuosity. Previous 346 data also shows that the SOAM exhibited an intermediate level of tortuosity 347 in tissue up to 1 cm away from a tumor, with a mean tortuosity between 348 that of tumors and non tumor-bearing animals (Rao et al., 2016). The dis-349 crimination power of the SOAM in our data set increases for smaller vessels 350 (Cohen's d = 0.14 for vessels with a radius > 0.11 mm, 0.28 with an interme-351 diate radius, and 0.43 with a radius < 0.09 mm). Therefore, its relation with 352 the extracted DCE-US features can not be fully appreciated due to the finite 353 resolution of AA. Similarly, a previous study has shown that the difference 354 in MVD between tumor and control increases for smaller vessels (Sedelaar 355 et al., 2001). Therefore, it may be that the SOAM, MVD, and other metrics 356 extracted in this study are related to dispersion; however, mainly smaller 357 vessels' properties have a significant influence on it. Supporting this hypoth-358 esis is the former observation that regions with increased MVD correspond 359 to those with low dispersion (Saidov et al., 2016), as derived from immuno-360 histology. The immunohistology derived MVD was based on evaluation of 361 tomato lectin binding to the endothelial cells and therefore characterized the 362 presence of vessels of all sizes. 363

Spots of increased vascular density or large vessels were detected with perfusion colormaps. The correlation between median perfusion level and MVD is the only significant inter-parameter agreement found in this work.

The Kruskall-Wallis test is ideally constructed for a study design when 367 subjects are randomly assigned to different groups, so that each subject ap-368 pears in one group only (Kruskall and Wallis, 1952). Moreover, the subjects 369 within the group must be independent. We realize that these assumptions 370 are not strictly valid in this study, since we observe the tumor evolution in 371 the same rats over time and since the vessels selected from the same rat are, 372 strictly speaking, not independent. However, we do not expect these limita-373 tions to be crucial for deriving a meaningful conclusion about the significant 374 trends in time. 375

Imaging initialization was different among 3 series of experiments, start-376 ing with day 8, 11, and 13 after tumor implantation, as explained before. We 377 consider that combining all the rats together according to the number of the 378 acquisition is justified as the imaging was initialized according to the same 379 strategy: when the tumors became palpable. However, since we waited for 380 2-3 days for subsequent assessment if tumors were not pulpable on day 8, in 381 future work it may be benifitial to assess the tumors every day or evaluate 382 all tumors in a single cohort. This would ensure that the development of the 383 imaged tumors is more consistent. 384

It is often observed that the wash-out phase is masked by recirculation in small animals. (Stapleton et al., 2009) shows that for a range of administered UCA doses the wash-out phase is more prominent in mice. Different UCA doses should therefore be investigated in our future work, since a prominent wash-out phase, in our experience, enhances the performance of CUDI (Kuenen et al., 2013b). A clear wash-out would also allow evaluating the wash-out as a complementary perfusion parameter.

An important limitation of this study is the 2D character of the extracted 392 parameters of dispersion and perfusion. The results of the post hoc tests, 393 therefore, must still be taken with caution since it was performed for 3D 394 vascular features evaluated in the whole tumor volume and 2D dispersion 395 and perfusion that leave us blind to out of plane information and restrict us 396 to the central tumor slice, which is not always representative of the whole 397 tumor (Streeter et al., 2011). We mitigated this limitation by performing 398 an additional spatial comparison of the parameter maps in the same plane, 399 matched with the help of large vessels identified in the perfusion maps. The 400 agreement between perfusion and MVD, noticed in the longitudinal trends, 401 was also identified in the spatial distribution of these parameters in the same 402 plane, raising more confidence to the finding that perfusion and MVD are 403 correlated. 404

An improved study design should either include 3D DCE-US (Schalk 405 et al., 2015), giving more accurate overall tumor characteristics, or a regis-406 tration procedure, allowing to fix the orientation of the probes and identify 407 the location of the DCE-US plane within the AA volume. The finding that 408 perfusion highlights large vessels can be used to further improve registration. 409 The abscence of any parameters correlated with dispersion may pinpoint 410 to the limitation of AA as a validation method for CUDI: while enabling very 411 high resolution ultrasound imaging, it may not be sufficient to find out which 412 vascular properties substantially influence dispersion, since dispersion may 413 be mainly defined by properties of subresolution vessels. In this respect, it is 414 possible to direct our attention to superlocalization methods that overcome 415 the limit of diffraction: they are able to track singe bubbles and determine 416

their exact positions by finding the centers of their point spread functions 417 (Cox and Beard, 2015; Errico et al., 2015). Another possible reason for the 418 abscence of vascular parameters that correlate with dispersion is that the 419 adopted dispersion parameter, is in fact related to both dispersion and flow 420 velocity (Kuenen et al., 2013a). Different vascular parameters may contribute 421 to the separate terms of dispersion and flow velocity, while we assessed their 422 combination. In this regard, it would also be of interest to apply another 423 analysis to the DCE-US bolus recordings that allows to separate dispersion 424 and velocity contributions (van Sloun et al., 2017). 425

# 426 Conclusions

In this work, dispersion demonstrated its superior performance at tumor classification compared to perfusion, as previously found for prostate cancer (Kuenen et al., 2013a,b; Mischi et al., 2012). Perfusion colormaps highlight large vessels and regions of elevated MVD. The vascular factors that determine the dispersion level remain yet to be found, as well as the role of vessels with a diameter below 100-200 μ in defining perfusion levels.

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- 441 Abbreviations
- 442 AA Acoustic angiography
- 443 CUDI Contrast ultrasound dispersion imaging
- 444 DCE-US Dynamic contrast-enhanced ultrasound
- 445 DM Distance metric
- 446 IDC Indicator dilution curve
- 447 LDRW Local density random walk
- 448 MR Mean radius
- 449 MVD Microvascular density
- 450 SOAM Sum of angles metric
- 451 TIC Time intensity curve
- 452 UCA Ultrasound contrast agent
- 453 VL Vessel length
- 454 VVD Volume vascular density

- <sup>455</sup> Brock M, Eggert T, Rein JP, Roghmann F, Braun K, Bjrn L, Sommerer F,
- J. N, Bodman C. Multiparametric ultrasound of the prostate: Adding contrast enhanced ultrasound to real-time elastography to detect histopathologically confirmed cancer. J Urol, 2013;189:93–98.
- <sup>459</sup> Bullitt E, Gerig G, Pizer SM, Lin W, Aylward SR. Measuring tortuosity of
  the intracerebral vasculature from mra images. IEEE Trans Med Imaging,
  <sup>461</sup> 2003;22:1163–1171.
- <sup>462</sup> Cosgrove D, Lassau N. Imaging of perfusion using ultrasound. Eur J Nucl
  <sup>463</sup> Med Mol Imaging, 2010;37:65–85.
- <sup>464</sup> Cox B, Beard P. Super-resolution ultrasound. Nature, 2015;527:451–452.
- Errico C, Pierre J, Pezet S, Desailly Y, Lenkei Z, Couture O, M. T. Ultrafast ultrasound localization microscopy for deep super-resolution vascular
  imaging. Nature, 2015;527:499–502.
- Folkman J. Tumor angiogenesis: theraputic implications. N Engl J Med,
  1971;285:1182–1186.
- <sup>470</sup> Gessner R, Frederick C, Foster F, Dayton P. Acoustic angiography: A new
  <sup>471</sup> imaging modality for assessing microvascular architecture. Int J Biomed
  <sup>472</sup> Imaging, 2013;2013.
- <sup>473</sup> Koumoutsakos P, Pivkin I, Milde F. The fluid mechanic of cancer and its
  <sup>474</sup> therapy. Annu Rev Fluid Mech, 2013;45:325–355.
- 475 Kruskall WH, Wallis WA. Use of ranks in one-criterion variance analysis. J
  476 Am Stat Assoc, 1952;47:583–621.

- Kuenen M, Saidov T, Wijkstra H, Mischi M. Contrast-ultrasound dispersion imaging for prostate cancer localization by improved spatiotemporal similarity analysis. Ultrasound Med Biol, 2013a;39:1631–1641.
- Kuenen M, Saidov T, Wijkstra H, Mischi M. Spatiotemporal correlation
  of ultrasound-contrast-agent dilution curves for angiogenesis localization
  by dispersion imaging. IEEE Trans Ultrason Ferroelectr Freq Control,
  2013b;60:2665–2669.
- Kuenen MPJ, Herold IHF, Korsten HHM, de la Rosette JJMCH, Wijkstra
  H. Maximum-likelihood estimation for indicator dilution analysis. IEEE
  Trans Biomed Eng, 2014;61:821–831.
- Kuenen MPJ, Mischi M, Wijkstra H. Contrast-ultrasound diffusion imaging for localization of prostate cancer. IEEE Trans Med Imaging,
  2011;30:1493–1502.
- Mann HB, Whitney D. On a test of whether one of two random variables is
  stochastically larger than the other. Ann Math Stat, 1947;18:50–60.
- Meiburger KM, Nam S, Chung E, Suggs LJ, Emelianov SY, Molinari F.
  Skeletonization algorithm-based blood vessel quantification using in vivo
  3d photoacoustic imaging. Phys Med Biol, 2016;61:7994–8009.
- <sup>495</sup> Mischi M, Kuenen MPJ, Wijkstra H. Angiogenesis imaging by spatiotempo<sup>496</sup> ral analysis of ultrasound-contrast-agent dispersion kinetics. IEEE Trans
  <sup>497</sup> Ultrason Ferroelectr Freq Control, 2012;59:621–629.
- <sup>498</sup> Narang AS, Varia S. Role of tumor vascular architecture in drug delivery.
  <sup>499</sup> Vol. 63, 2011.

- Panfilova A, Shelton S, Sloun R, Demi L, Wijkstra H, Dayton P, Mischi M.
   Does contrast ultrasound dispersion imaging reveal changes in tortuosity?
   a comparison with acoustic angiography, 2016.
- <sup>503</sup> Ponce A, Viglianti B, Yu D, Yarmolenko P, Michelich C, Woo J, Bally M, De<sup>504</sup> whirst M. Magnetic resonance imaging of temperature-sensitive liposome
  <sup>505</sup> release: Drug dose painting and antitumor effects. J. Natl Cancer Inst,
  <sup>506</sup> 2007;99:53-63.
- <sup>507</sup> Quaia E. Assessment of tissue perfusion by contrast-enhanced ultrasound.
   <sup>508</sup> Eur Radiol, 2011;21:604–615.
- Rao S, Shelton S, Dayton P. The 'fingerprint' of cancer extends beyond solid
  tumor boundaries: assessment with a novel ultrasound imaging approach.
  IEEE Trans Biomed Eng, 2016;63.
- <sup>512</sup> Rognin NG, Frinking P, Costa M, Arditi M. In-vivo perfusion quantifica<sup>513</sup> tion by contrast ultrasound: Validation of the use of linearized video data
  <sup>514</sup> vs. raw rf data. In: 2008 IEEE Int Ultrason Symp Proceedings. Piscat<sup>515</sup> away:IEEE, 2008. pp. 1690–1693.
- Saidov T, Heneweer C, Keunen M, Broich-Oppert J, Wijkstra H, Rosette J,
  Mischi M. Fractal dimension of tumor microvasculature by dce-us: preliminary study in mice. Ultrasound Med Biol, 2016;42:2852–2863.
- Schalk S. Towards 3d prostate cancer localization by contrast ultrasound
   dispersion imaging. Ph.D. thesis, 2017.
- Schalk SG, Demi L, Smeenge M, Millis DM, Wallace K, de la Rosette J,
  Wijkstra H, Mischi M. 4d spatiotemporal analysis of ultrasound contrast

- agent dispersion for prostate cancer localization: a feasibility study. IEEE
  Trans. Ultrason. Ferroelectr. Freq. Control, 2015;62:839–851.
- Sedelaar JM, Leenders G, Hulsbergen-van de Kaa C, Poel H, Laak J, Debruyne F, Wijkstra H, Rosette J. Microvessel density: Correlation between contrast ultrasonography and histology of prostate cancer. Eur Urol,
  2001;40:285–293.
- Shelton S, Lee Y, Lee M, Cherin E, Foster F, Aylward S, Dayton P. Quantifi cation of microvascular tortuosity during tumor evolution utilizing acoustic
   angiography. Ultrasound Med Biol, 2015;41:1869–1904.
- Stapleton S, Goodman H, Yu-Qing Z, Cherin E, Henkelman R, Burns P,
  Foster F. Acoustic and kinetic behavior of definity in mice exposed to high
  frequency ultrasound. Ultrasound Med Biol, 2009;35:296–307.
- Streeter J, R.C. G, Tsuruta J, Feingold S, Dayton P. Assessment of molecular imaging of agiogenesis with three-dimentional ultrasonography. Mol
  Imaging, 2011;10:460–468.
- Strouthos C, Lampaskis M, Sboros V, McNeilly A, Averkiou M. Indicator
  dilution models for the quantification of microvascular blood flow with
  bolus administration of ultrasound contrast agents. IEEE Trans Ultrason
  Ferroelectr Freq Control, 2010;57:1296–1310.
- Sullivan MG, Feinn R. Using effect size or why the p-value is not enough.
  J Grad Med Educ, 2012;4:279–282.

- Tozer GM, Lewis S, Michalowski A, Aber V. The relationship between regional variations in blood flow and histology in a transplanted rat fibrosarcoma. Br. J. Cancer, 1990;61:250–257.
- Vala HJ, Baxi A. A review on otsu image segmentation algorithm. Int J Adv
  Res Comput Sci Softw Eng, 2013;2:387–389.
- van Sloun RJ, Demi L, Postema AW, de la Rosette JJ, Wijkstra H, Mischi
  M. Ultrasound-contrast-agent dispersion and velocity imaging for prostate
  cancer localization. Med Image Anal, 2017;35:610–619.
- <sup>552</sup> Viglianti B, Abraham S, Michelich C, Yarmolenko P, MacFall J, Bally
  <sup>553</sup> M, Dewhirst M. In vivo monitoring of tissue pharmacokinetics of lipo<sup>554</sup> some/drug using mri: Illustration of targeted delivery. Magnet Reson Med,
  <sup>555</sup> 2004;51:1153–1162.
- Zhao H, Xu R, Ouyang Q, Chen L, Dong B, Huihua Y. Contrast-enhanced
  ultrasound is helpful in the differentiation of malignant and benign breast
  lesions. Eur J Radiol, 2010;73:288–293.

# 559 Figure Captions

Figure 1: DCE-US and AA images of the same plane, and maps of the 560 extracted features. a: maximum intensity projection of the DCE-US 561 video. The tumor is encircled by a red contour, while the region out-562 side the blue contour belongs to the control, separated by a margin 563 which was not included in the analysis. Regions with power below the 564 threshold of -22 dBs of the maximum intensity are displayed in black. 565 b-c: perfusion and dispersion colormaps, respectively. Regions with 566 power below -22 dBs of the maximum intensity are displayed in white. 567 d: Selected AA slice. e-f: vascular skeleton, colorcoded according to 568 the values of the microvascular desity, and mean radius, respectively 569 (yellow indicates low values, while red inicates high values). The num-570 bers in b and d illustate the vessels identified in the perfusion maps, 571 used as markers to locate the right plane in AA volumes. 572

Figure 2: A typical preprocessed indicator dilution curve. T1 shows the
appearance time, T0 is taken 3 seconds before appearance time. The
interval from T0 to T2 shows the interval of the IDC used for dispersion analysis. The tangens of the angle alpha of the line fitted to the
indicator dilution curve in the 2 seconds after appearance time is the
wash-in-rate.

- Figure 3: AA maximum intensity projection. The tumor region is indicated
  by the red contour, surrounded by the control region.
- Figure 4: Boxplots of tumor and control parameters, binned together from
  all time points. a: dispersion, b: perfusion, c: distance metric, d: sum

of angles metric, e: vessel length, f: vessel radius, g: microvascular density, h: volume vascular density. Cohen's d measure is indicated above the plots.

- Figure 5: Boxplots of tumor (T1, T2, T3, T4) and control (C1, C2, C3, C4)
  parameters, binned together at different time points. a: dispersion,
  b: perfusion, c: volume vascular density, d: microvascular density, e:
  vessel radius, f: vessel length, g: sum of angles metric.
- Figure 6: Results of the post hoc Kruskall-Wallis test performed on tumor
  and control parameters at four time points (indicated by T1, T2, T3,
  T4 and by C1, C2, C3, C4, respectively). The colors of the rows indicate whether the distribution is significantly different from the others,
  green and yellow representing different significance levels. a: dispersion, b: perfusion, c: volume vascular density, d: microvascular density,
  e: vessel radius, f: vessel length, g: sum of angles metric.