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Assessment of Left Atrial Deformation and Function by 2-Dimensional Speckle Tracking Echocardiography in Healthy Dogs and Dogs With Myxomatous Mitral Valve Disease

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Background: The assessment of left atrial (LA) function by 2-dimensional speckle tracking echocardiography (STE) holds important clinical implications in human medicine. Few similar data are available in dogs.

Objectives: To assess LA function by STE in dogs with and without myxomatous mitral valve disease (MMVD), analyzing LA areas, systolic function, and strain.

Animals: One hundred and fifty dogs were divided according to the American College of Veterinary Internal Medicine classification of heart failure: 23 dogs in class A, 52 in class B1, 36 in class B2, and 39 in class C + D.

Methods: Prospective observational study. Conventional morphologic and Doppler variables, LA areas, and STE-based LA strain analysis were performed in all dogs and results were compared among groups. Correlation analysis was carried out between LA STE variables and other echocardiographic variables.

Results: Variability study showed good reproducibility for all the tested variables (coefficient of variation <16%). Left atrial areas, fractional area change, peak atrial longitudinal strain (PALS), peak atrial contraction strain, and contraction strain index (CSI) differed significantly between groups B2 and C + D and all the other groups (overall $P < .001$), whereas only PALS differed between groups B1 and A ($P = .01$). Left atrial areas increased with progression of the disease, whereas LA functional parameters decreased. Only CSI increased nonsignificantly from group A to group B1 and then progressively decreased. Thirty-one significant correlations ($P < .001$, $r > .3$) were found between conventional left heart echocardiographic variables and LA areas and strain variables.

Conclusions and Clinical Importance: Left atrial STE analysis provides useful information on atrial function in the dog, highlighting a progressive decline in atrial function with worsening of MMVD.

Key words: Canine; Cardiology; Left atrium; Strain.

The left atrium (LA) plays an essential role in cardiac performance by means of 3 main functions, namely reservoir, conduit, and booster pump function.¹ Reservoir phase represents the collection of pulmonary venous flow during ventricular systole, conduit phase is the passage of blood from the pulmonary veins to the left ventricle (LV) in early diastole, whereas booster pump function represents the active contraction in late diastole. Assessment of LA function is essential for clinical evaluation and prognostic purposes in dogs affected by myxomatous mitral valve disease (MMVD).² Several

Abbreviations:

2D	2-dimensional
Ao	aortic root
BW	body weight
CHF	congestive heart failure
CSI	contraction strain index
CV	coefficient of variation
FAC	fractional area change
LA _{Amax}	left atrial maximal area
LA _{Amin}	left atrial minimal area
LA	left atrium
LV	left ventricle
MMVD	myxomatous mitral valve disease
MR	mitral regurgitation
MV	mitral valve
PACS	peak atrial contraction strain
PALS	peak atrial longitudinal strain
STE	speckle tracking echocardiography
TDI	tissue Doppler imaging
TTE	transthoracic echocardiography

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methods have been used to assess LA function in humans, including 2-dimensional (2D) and 3-dimensional echocardiography, computed tomography, and magnetic resonance.^{3,4} In dogs, the majority of studies have been focused on conventional echocardiographic variables, such as calculation of LA phasic sizes (areas and volumes),^{5–8} and pulsed-wave Doppler evaluation of transmittal and pulmonary venous flow.⁹ Only 1 study evaluated the prognostic relevance of LA function in dogs with MMVD, showing a worse prognosis in those animals with a poorer LA systolic function.² However, most of the studied echocardiographic

parameters of LA function are highly preload dependent and they are influenced by changes in hemodynamic load associated with mitral regurgitation (MR), thus raising the need for more sophisticated echocardiographic techniques.¹⁰ Assessment of atrial deformation profiles obtained by tissue Doppler imaging (TDI) and its derived variables, strain and strain rate, has been recently proposed as an alternative method of exploring LA mechanics in both humans^{3,4,10} and dogs.¹¹ Nevertheless, a number of potential drawbacks of this approach should be considered, including suboptimal reproducibility, angle dependence, and the confounding effect of noise artifacts.^{3,4,10,11} Many of these limitations might be overcome by 2D speckle tracking echocardiography (STE), the most recent and promising ultrasound technology for the direct evaluation of LA function from standard grayscale echocardiographic images.^{3,4,10} Specifically, regional myocardial deformation can be calculated from continuous frame-by-frame tracking of the tiny echo-dense speckles created by the interference between the ultrasound beam and myocardium. The movement of the speckle during the cardiac cycle follows myocardial movement, and a change between speckles is assumed to represent myocardial deformation. In this manner, STE allows for the assessment of myocardial function independent of cardiac translation and angle of insonation.^{3,4,10} In people, reference range values, feasibility, and reproducibility of STE for the study of LA mechanics have been extensively documented.^{12–15} Furthermore, the clinical and prognostic importance of STE variables has been demonstrated in numerous pathophysiologic conditions typically associated with abnormal LA function, including MMVD, atrial fibrillation, systemic hypertension, and dilated, as well as hypertrophic, cardiomyopathy.^{3,4,10,16,17} The majority of STE studies in dogs to date focused on the evaluation of mechanical ventricular properties.^{18–27} Few data are available regarding the application of this technique for LA assessment^{28–30}, and no studies have focused on the evaluation of LA STE-derived strain in dogs with cardiac disease.

Therefore, the aims of this study were (1) to document the feasibility and reproducibility of STE in the assessment of LA function in dogs; (2) to compare selected STE indices of LA function between healthy dogs and dogs with MMVD at different stages; and (3) to evaluate the association between these indices and traditional echocardiographic variables.

Materials and Methods

Animals

Dogs were presented to the cardiology unit of the Veterinary Teaching Hospital of the University of Bologna for cardiac evaluation on the basis of cardiac murmur assessment, clinical signs referable to heart disease (ie, dyspnea, cough, and syncope), for breed screening, or for pre-anesthetic assessment. Dogs were prospectively recruited from January 2011 to December 2014 and included in the study if they had clinical and echocardiographic signs typical for MMVD and a body weight (BW) <21 kg. Dogs with congenital cardiac malformations, dilated cardiomyopathy, or

other acquired cardiac diseases were excluded. Dogs were permitted to receive cardiac medications for congestive heart failure (CHF) if considered necessary by the clinician. Healthy dogs were recruited from students, university staff, and owners as the control group. These animals were considered clinically healthy based on cardiovascular examination, noninvasive systemic arterial blood pressure measurements, and complete transthoracic echocardiographic examination.

Procedures

All dogs underwent a complete physical examination, cardiac auscultation, and noninvasive systemic arterial blood pressure measurement. Complete transthoracic echocardiography (TTE) was performed according to standard techniques³¹ by 2 investigators (MBT and MC) using ultrasound units^{a,b} equipped with phased array transducers of various frequencies and continuous ECG tracing. The diagnosis of MMVD was made based on clinical examination and echocardiographic criteria including thickening, prolapse, or both thickening and prolapse of the mitral valve (MV) leaflets on 2D echocardiography and MR on color Doppler examination. Wall thickness and chamber diameter of the LV were obtained from 2D guided M-mode images acquired from a right parasternal short-axis view at the level of the papillary muscles. In particular, LV end-diastolic and end-systolic dimensions were normalized to the aortic root (Ao) diameter and compared with previously published reference intervals, in order to identify the presence of LV dilatation.³² Left atrial diameter, Ao diameter, and Ao area were measured on 2D images obtained from a right parasternal short-axis view at the basilar level,⁵ and a LA/Ao ratio <1.5 was considered normal.³³ Left ventricular diastolic inflow was recorded from the left apical 4-chamber view placing the sample volume at the level of the MV on the ventricular side. Three consecutive measurements were averaged for each variable. Thoracic radiography in 2 orthogonal projections was performed to exclude or confirm the presence of CHF (pulmonary edema, pleural effusion, or both) in all dogs with left atrial dilatation or according to clinical signs of cough, dyspnea, or both. Radiographic evidence of interstitial opacity in the caudo-dorsal fields in the presence of cardiomegaly was considered indicative of CHF. If CHF was present at the time of presentation, dogs were first stabilized and a complete TTE was then performed for the first time or repeated. Only dogs in sinus rhythm during echocardiography were included in the study.

According to the guidelines for the diagnosis and treatment of MMVD,³⁴ dogs were divided into different groups as follows: control, healthy dogs with normal LV and LA dimensions (group A); dogs having MMVD with normal LV and LA dimensions (group B1); dogs having MMVD with LV, LA, or both dimensions above the reference limits^{32,33} but without confirmed radiographic evidence of CHF (group B2); dogs with symptomatic MMVD and radiographic evidence of pulmonary edema/pleural effusion; or dogs with at least 1 confirmed episode of CHF in the past (group C + D).

Left Atrial Areas and Speckle Tracking Echocardiography

The same operator (MBT) that was blinded to the dogs' classification performed this part of the analysis. For the analysis of LA areas and STE, a left apical 4-chamber view was optimized in order to visualize the LA walls. Then, 3 consecutive cardiac beats cine loops were acquired and stored in DICOM format for subsequent offline analysis. Dedicated software^c was used to review the cine loops, the beat with the best image quality was chosen, and the endocardial border of the LA was automatically drawn by the software and LA areas provided during the entire cardiac cycle. If

necessary, manual editing of the tracing was done in order to correct software errors in tracking the endocardial border. Left atrial maximal area (LAAmax) and minimal area (LAAmin) expressed in cm^2 were automatically calculated by the software and were then indexed to Ao area. Left atrial fractional area change (FAC) was calculated with the formula: $\text{FAC} = [(LAA_{\text{max}} - LAA_{\text{min}}) / LAA_{\text{max}}] \times 100$, and expressed as a percentage.²

The same cine loops and beats used to measure LA areas were successively employed for the STE analysis. The operator selected 3 points on the LA (2 on the MV annulus and 1 on the LA roof), and the software automatically drew a region of interest on the entire LA wall dividing it into 7 regions (from the basilar segments to the roof of the LA) (Fig 1). When necessary, the operator manually edited the tracking of each segment, and those segments that showed artifactual values (due to lung interposition or echo drop-out at the level of the region of the *fossa ovalis* or the pulmonary veins inlet) were excluded from the analysis. The number of segments considered for each dog was recorded. Time/intensity curves were then generated for each segment displacing strain values (as percentages) on the y-axis and time (in seconds) on the x-axis over an entire cardiac cycle. The software also generated a mean curve of all the segments that was used by the operator to measure the peak strain value during LV contraction (peak atrial longitudinal strain [PALS]), and just before LA active contraction, on the peak of the P wave of the ECG trace (peak atrial contraction strain [PACS]). Finally, the contraction strain index (CSI) was calculated by the formula: $\text{CSI} = [(PALS/PACS) \times 100]$ (Fig 1). This last parameter represents the contribution of LA contraction on LV filling.¹⁷

Variability Study

To assess measurement variability, 6 dogs enrolled into the study were randomly selected. For each dog 1 single echocardiographic study was then chosen, thus a total of 6 different echocardiographic studies were analyzed. For interobserver variability, 2 operators with different levels of experience in echocardiography (MBT, GR) analyzed and measured the LA areas (LAAmax and LAAmin) and FAC, and LA STE variables (PALS, PACS, CSI). To assess interday intra-observer variability, the same operator (MBT) repeated the measurements for each of the 6 dogs on different days (at least 7 days apart). Attention was made to analyze the same cardiac cycle for each loop. Finally, to assess interday intra-observer acquisition variability, 5 echocardiographic studies from the same number of dogs from those enrolled in the control group, were repeated 1 week apart. Then, the same operator (MBT) measured LA areas and LA STE variables. The variability was then quantified as the coefficient of variation (CV) by use of the equation $\text{CV} = (\text{mean difference between measurement/mean of measurements})/100$ and expressed as a percentage. The degree of variability was arbitrarily defined as follows: $\text{CV} < 5\%$, very low variability; 5–15%, low variability; 16–25%, moderate variability; or $> 25\%$, high variability.³⁵

Statistical Analysis

Descriptive statistics were used for age, sex, and BW. A 1-way ANOVA and a chi-square test were performed to evaluate the group effect (A, B1, B2, C + D) on BW and age, and sex,

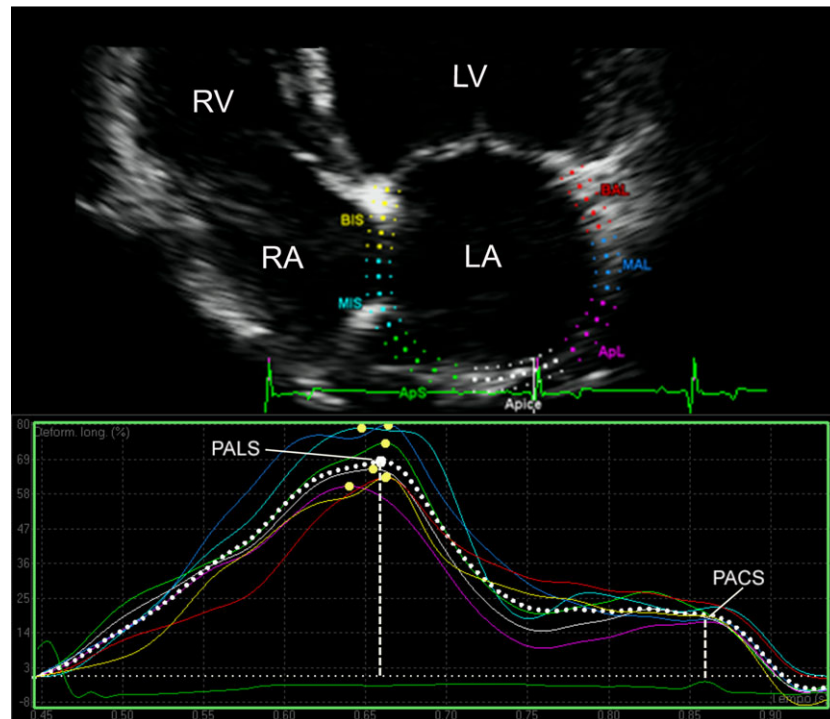


Fig 1. Representative image of the LA strain profile of a dog. On the top of the panel, the LA visualization is optimized in the left apical 4-chamber view. A region of interest is manually drawn to include the LA wall, which is automatically divided into 7 different segments with different colors. On the bottom of the panel, a trace screen of the speckle tracking echocardiography-derived strain (ϵ) is shown, which is expressed in percentage on the y-axis and time (in seconds) on the x-axis. Different colored lines are traced for each LA segment, and the mean ϵ value over time is presented as a dotted white line. The profile is characterized by 2 positive deflections, 1 during ventricular systole (PALS), and 1 just before LA contraction, on the P wave (PACS, peak contraction strain). The CSI is calculated from PALS and PACS as their difference expressed in percentage (see text for details). LA, left atrial; LV, left ventricle; RA, right atrium; RV, right ventricle; PACS, peak atrial contraction strain; PALS, peak atrial longitudinal strain; CSI, contraction strain index.

respectively. Normal distribution of the echocardiographic, Doppler, and STE-derived variables was assessed by a Shapiro–Wilks test. Variables not normally distributed were log-transformed. An ANCOVA model was used to test the fixed effects of the group of dogs, sex, as well as the continuous variables of heart rate, BW and age as covariates on echocardiographic, Doppler, and STE-derived variables. The hypothesis of the linear model was assessed by visual inspection of the residuals plots.

When a significant main effect was detected, a post hoc pairwise contrast were conducted by a Bonferroni correction. Data were reported as Least Squares means (LS-means) \pm Standard error (SE). The association between STE-derived variables and left heart measurements was tested with a Pearson correlation index (r) and considered significant for $P < .001$ and $r > .3$, then a linear regression model was applied.³⁶

All statistical analyses were performed by 2 commercial dedicated software programs.^{d,e} An overall value of $P < .01$ was considered significant.

Results

General Characteristics of the Study Population and Conventional Echocardiographic Variables

One hundred and fifty dogs met the inclusion criteria. Of these dogs, 23 (15%) were clinically healthy (group A), 52 (35%) had MMVD without cardiac enlargement (group B1), 36 (24%) had MMVD with cardiac enlargement (group B2), and 39 (26%) had symptomatic

MMVD (group C + D). Dogs in group A were younger compared with all the other groups, whereas dogs in group B1 were younger than those in group B2 and C + D ($P < .001$). Dogs of group B1 were significantly heavier than those in group C + D ($P = .028$). Male dogs were overrepresented in groups B1, B2, and C + D, whereas female dogs were overrepresented in group A; however, a statistically relevant difference was only observed between groups B2 and A ($P = .042$). Mixed-breed dogs were overrepresented ($n = 68$; 45%), followed by Cavalier King Charles Spaniels ($n = 10$; 7%), Yorkshire Terriers ($n = 9$; 6%), Miniature Poodles ($n = 7$; 5%), Maltese ($n = 6$; 4%), Shih Tzu ($n = 6$; 4%), and Cocker Spaniels ($n = 3$; 2%). All descriptive characteristics of the study population and all drugs used with potential impact on hemodynamic variables are summarized in Table 1.

All echocardiographic variables were normally distributed, except LAmax/AoArea and LAmin/AoArea. The heart rate was higher in dogs of group C + D compared with those of all other groups ($P < 0.001$). There was a progressive increase of MV E velocity, MV A velocity, and MV E:A ratio when comparing dogs of groups A and B1 to those of groups B2 and C + D ($P < .001$) (Table 2). None of the dogs had systemic arterial hypertension on noninvasive measurement.

Table 1. Demographic data from 150 dogs used in this study.

Variable	A	B1	B2	C + D	Overall P
No. of dogs	23	52	36	39	
Age (years)	7.0 \pm 0.6	9.3 \pm 0.4	11.1 \pm 0.5*	12.8 \pm 0.5***	<.001
Body weight (kg)	10.5 \pm 0.9	11.1 \pm 1.0	9.3 \pm 1.1	8.1 \pm 0.9	.028
Sex (male/female)	8/15	30/22	26/10	21/18	.042
Breed (no. of dogs)	Mixed breed (9) Cocker Spaniel (3)	Mixed breed (23) CKCS (8) Yorkshire Terrier (4)	Mixed breed (11) Yorkshire Terrier (5) Beagle, Bolognese, CKCS, Dachshund, Lagotto Romagnolo, Shih Tzu (2)	Mixed breed (25) Miniature Poodle (5) Shih Tzu (4) Maltese (2)	
Medications received (no. of dogs) [mean dose received]	Other breeds (7) None	Other breeds (9) None	Other breeds (8) None	Other breeds (3) Benazepril (39) [0.38 mg/kg q12–24h] Pimobendan (35) [0.31 mg/kg q24h] Amlodipine (5) [0.3 mg/kg q24h] Spironolactone (3) [2 mg/kg q24h] Furosemide (39) [2 mg/kg q12h]	

A, healthy control group; B1, dogs with MMVD without cardiac enlargement; B2, dogs with MMVD and cardiac enlargement; C + D, dogs with MMVD and clinical signs due to congestive heart failure; CKCS, Cavalier King Charles Spaniel; MMVD, myxomatous mitral valve disease.

Data are expressed as Least Squares-Means \pm SE and numerical data for sex prevalence. In bold values with statistical differences

* $P < .001$ compared with group A.

** $P < .001$ compared with group B1.

Table 2. Conventional echocardiographic and left atrial STE-derived variables from a population of 23 healthy dogs (A) and 127 dogs with MMVD at different stages.

Variable	A (n = 23)	B1 (n = 52)	B2 (n = 36)	C + D (n = 39)	Overall P Value
Conventional					
Heart rate (bpm)	137 ± 6	130 ± 4	135 ± 4.8	160 ± 4.6 ^{***§§§##}	<.001
MV E vel (cm/s)	71.3 ± 5.64	77.2 ± 3.71	112.7 ± 4.45 ^{***§§§}	139.1 ± 4.31 ^{***§§§,###}	<.001
MV A vel (cm/s)	63.2 ± 5.24	71.7 ± 3.49	83.7 ± 4.19 [*]	97.2 ± 4.03 ^{***§§§}	<.001
MV E/A	1.2 ± 0.09	1.1 ± 0.06	1.4 ± 0.07 [§]	1.6 ± 0.07 ^{***§§§}	<.001
STE					
LAAmax/AoArea	2.2 (1.9–2.4)	2.7 (2.4–3.0)	3.6 (3.3–4.1) ^{***§§}	5.2 (4.7–5.9) ^{***§§§,##}	<.001
LAamin/AoArea	0.9 (0.8–1.1)	1.3 (1.2–1.5)	2.0 (1.8–2.3) ^{***§§}	3.5 (3.0–4.0) ^{***§§§,###}	<.001
FAC (%)	55.1 ± 2.09	49.2 ± 1.39	44.7 ± 1.67 ^{***}	36.2 ± 1.60 ^{***§§§,##}	<.001
PALS (%)	60.4 ± 2.36	49.8 ± 1.57 ^{**}	40.6 ± 1.89 ^{***§§}	28.9 ± 1.81 ^{***§§§,###}	<.001
PACS (%)	26.3 ± 1.77	24.3 ± 1.18	16.5 ± 1.42 ^{***§§§}	10.6 ± 1.32 ^{***§§§,##}	<.001
CSI (%)	42.8 ± 3.24	49 ± 2.16	40.0 ± 2.59 ^{**}	33.9 ± 2.49 ^{§§§}	<.001

C + D, dogs with MMVD and clinical signs due to congestive heart failure; MV E vel, mitral valve E wave peak velocity; MV A vel, mitral valve A wave peak velocity; E/A, mitral valve peak E; and A wave velocity ratio; LAAmax/AoArea, left atrial maximal area-to-aortic area ratio; LAamin/AoArea, left atrial minimal area-to-aortic area ratio; FAC, left atrial fractional area change; PALS, peak atrial longitudinal strain; PACS, peak atrial contraction strain; CSI, contraction strain index; MMVD, myxomatous mitral valve disease; STE, speckle tracking echocardiography.

Data are expressed as LSmeans ± SE. LAAmax/AoArea and LAamin/AoArea mean values are reported after anti-log transformation and CI are reported into parentheses. In bold values with statistical differences.

**P* < .05 compared with group A (healthy dogs).

***P* < .01 compared with group A.

****P* ≤ .001 compared with group A.

§*P* < .05 compared with group B1 (dogs with MMVD without heart enlargement).

§§*P* < .01 compared with group B1.

§§§*P* < .001 compared with group B1.

#*P* < .05 compared with group B2 (dogs with MMVD and heart enlargement).

##*P* < .01 compared with group B2.

###*P* ≤ .001 compared with group B2.

Left Atrial Areas, LA STE Variables, and Correlation Analysis

For each dog, it was possible to obtain a good tracking profile of the atria and meaningful values for LA areas and strain. Similar to the linear measurements, LA areas also increased with progression of MMVD (Table 2). Specifically, LAAmax/AoArea and LAamin/AoArea were higher in dogs of group B2 compared with those of groups A and B1, whereas dogs of group C + D had higher values compared with those of all the other groups (*P* < .001). Fractional area change was lower in dogs of group B2 compared with that of group A, whereas dogs of group C + D had lower values compared with those of all the other groups (*P* < .001).

Of 1,050 LA segments (7 for each dog), only 61 segments (5.8%) were censored due to artifacts whereas the remaining 989 segments (94.2%) were analyzed. Regarding the strain values, dogs of group C + D had lower values for PALS compared with those of all the other groups, and lower PACS values compared only to groups A and B1; dogs of group B2 had lower values compared with dogs of groups B1 and A, whereas dogs of group B1 had lower PALS values compared with those of group A (*P* < .001). The CSI had significantly lower values in dogs of group C + D compared with those of group

B1 and in dogs of group B2 compared with those of group A (*P* < .001) (Table 2).

Results of the ANCOVA model did not show a significant effect of heart rate or age on each of the tested variables. Body weight had a significant negative effect on FAC, PALS and PACS, a positive effect on LAAmax/AoArea and LAamin/AoArea, and no effect on CSI.

The correlation analysis demonstrated several positive and negative correlations between the LA STE-derived variables and morphologic and Doppler variables of the LA and LV. In particular, 31 significant correlations (*P* < .001) with values of *r* > .3 were found between LA areas and functional variables (FAC, PALS, PACS, and CSI) and conventional left heart echocardiographic variables (Table 3; Fig 2).

Variability Study

The interobserver measurement variability was low for the 6 tested STE variables, with CV values always below 16%. Intra-observer interday measurement variability was always very low or low, with CV values always below 13% for PALS, PACS and CSI and below 7% for LAAmax, LAamin and FAC. Interday intra-observer acquisition variability was low (between 5% and 15%) for all variables except FAC, where the variability was 16%.

Table 3. Correlation analysis between speckle tracking echocardiography-derived variables and some selected conventional echocardiographic parameters in the 150 dogs used in this study.

Variable	LAAmax/ AoArea		LAAmin/ AoArea		FAC		PALS		PACS		CSI	
	<i>r</i>	<i>P</i>	<i>r</i>	<i>P</i>	<i>r</i>	<i>P</i>	<i>r</i>	<i>P</i>	<i>r</i>	<i>P</i>	<i>r</i>	<i>P</i>
Heart rate (bpm)	.268	<.001	.300	<.001	-.129	.116	-.183	.025	-.113	.170	-.025	.757
LA/Ao	.750	<.001	.756	<.001	-.538	<.001	-.609	<.001	-.539	<.001	-.303	.002
LVIDd/Ao	.697	<.001	.693	<.001	-.473	<.001	-.540	<.001	-.521	<.001	-.332	<.001
LVIDs/Ao	.452	<.001	.458	<.001	-.356	<.001	-.315	<.001	-.309	.001	-.187	.022
MV E vel (cm/s)	.644	<.001	.669	<.001	-.515	<.001	-.593	<.001	-.586	<.001	-.385	<.001
MV A vel (cm/s)	.405	<.001	.384	<.001	-.200	.014	-.279	.001	-.206	.012	-.028	.731
E/A	.312	<.001	.362	<.001	-.400	<.001	-.398	<.001	-.460	<.001	-.385	<.001

LAAmax/AoArea, left atrial maximal area-to-aortic area ratio; LAAmin/AoArea, left atrial minimal area-to-aortic area ratio; FAC, left atrial fractional area change; PALS, peak atrial longitudinal strain; PACS, peak atrial contraction strain; CSI, contraction strain index; LA/Ao, left atrium-to-aorta ratio; LVIDd/Ao, left ventricular internal diameter in diastole-to-aorta ratio; LVIDs/Ao, left ventricular internal diameter in systole-to-aorta ratio; MV E vel, mitral valve E wave peak velocity; MV A vel, mitral valve A wave peak velocity; E/A, mitral valve peak E and A wave velocity ratio.

In bold values with significant correlation ($P < .001$, $r > .3$).

Discussion

In the present study we evaluated novel echocardiographic indices of LA function in healthy dogs and dogs with MMVD. The major findings were: (1) LA STE analysis is feasible and reproducible in the dog, (2) several statistical significant differences exist in LA STE variables between healthy dogs and dogs with MMVD at different stages of severity, indicating a progressive impairment of LA function with the disease progression, and (3) a strong association is present between LA STE-derived indices and some conventional echocardiographic variables, supporting a reduction of LA function as the heart dilates.

Two-dimensional STE was feasible and repeatable for evaluation of LA longitudinal strain in awake, unanesthetized dogs with and without MMVD, with a lower variability compared with TDI-derived LA strain imaging in healthy dogs.¹¹ Specifically, intra- and interobserver variability was considered clinically acceptable (CV <16%) for all STE measurements. In addition, the obtained images were of sufficient quality to allow STE analysis, both in dogs with a structurally normal LA and in dogs with an enlarged LA. Similar data have been reported in healthy people, where adequate tracking quality was achieved in >95% of segments analyzed and inter- and intra-observer variability coefficients of measurements ranged approximately between 3% and 6%.^{13,37} The CV relative to LAAmax, LAAmin and FAC were slightly lower than those obtained for the STE variables (PALS, PACS, and CSI). This finding could be explained by the fact that the more complex software analysis needed to calculate the STE variables and the need of analyzing high-quality images, might increase their variability, whereas the simple area measurements, which require less computation and not necessarily high-quality images, allow higher repeatability. Speckle tracking echocardiography analysis allows an excellent assessment of the LA deformation profile during an entire cardiac cycle, closely following LA

physiology. In particular, during the period of LV systole, the LA acts as a reservoir allowing collection of blood from the pulmonary veins whereas the MV is closed. Consequently, longitudinal elongation of the LA walls occurs, resulting in a LA strain increase characterized by a peak at the end of LA filling, just before MV opening. During the successive conduit phase, the MV opens and the LA empties quickly; the shortening of LA walls results in a strain decrease, up to a plateau corresponding to the phase of diastole. Finally, after the P wave of the ECG, the LA acts as an active contractile chamber in order to complete the LV filling, reaching a second negative peak at the end of LA contraction.^{11,13,16} Starting from the above-described strain curve, specific STE variables of LA function can be calculated, including PALS and PACS, which reflect the LA function during its reservoir and booster pump phase, respectively.¹⁷ In addition, CSI can be also obtained from these 2 variables, representing the contribution of LA active contraction to the LV filling phase.¹⁷ Our results showed a linear decrease of LA STE-derived measurements including FAC, another index of LA function, from healthy dogs to dogs with progressively more severe MMVD. In addition, a negative correlation was found between the above-mentioned STE-derived variables and conventional echocardiographic measurements of left cardiac chamber size. These data are in agreement with previous reports focused on conventional echocardiographic assessment of LA function in dogs with MMVD,^{2,7} as well as human studies in patients investigated by STE,^{38–44} and suggest progressive LA dysfunction with worsening of MMVD and LA dilatation. These findings might be explained by the progressive LA volume overload due to MR and remodeling and by an increase in LV filling pressures in the advanced phases of the disease. Specifically, LA remodeling includes adaptive and maladaptive processes, such as alterations in the composition of extracellular matrix with excessive fibroblast proliferation, and myocyte hypertrophy, necrosis, and

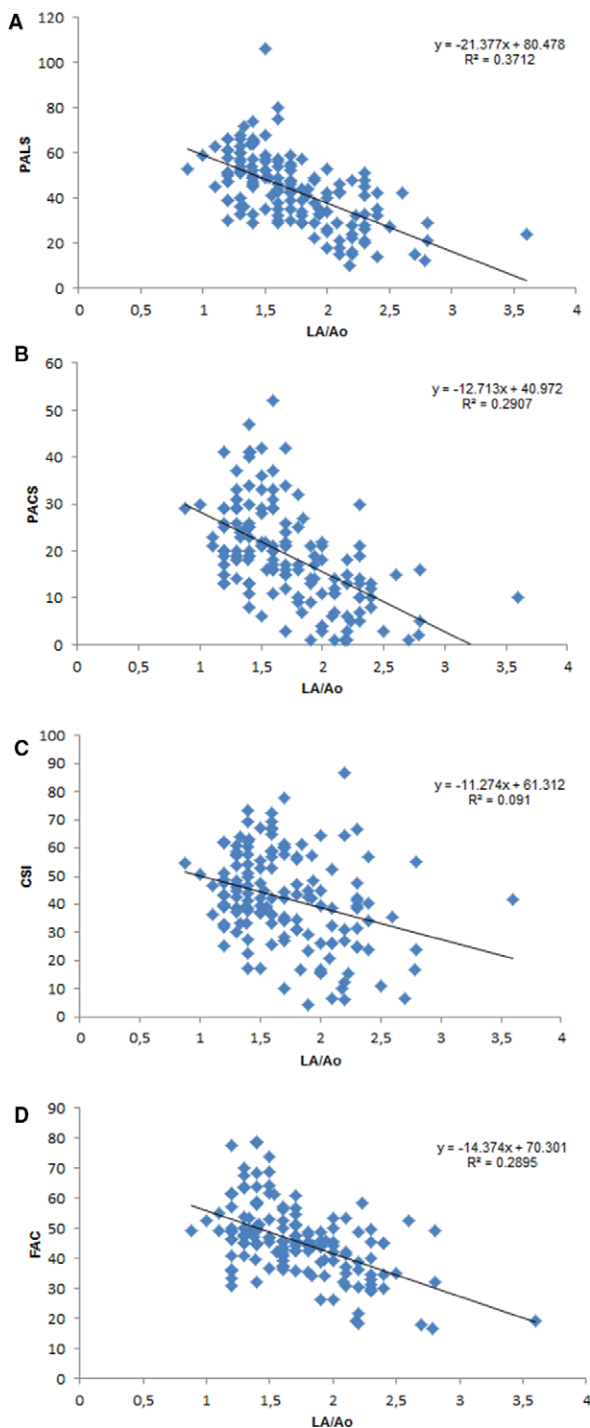


Fig 2. Linear regression analysis of 4 selected LA strain variables and FAC obtained from LA areas, compared with LA linear dimension normalized to aortic root diameter (LA/Ao). LA, left atrial diameter; AO, aortic diameter; PALS, peak atrial longitudinal strain; PACS, peak atrial contraction strain; CSI, contraction strain index; FAC, fraction area change.

apoptosis.^{45,46} All of these ultrastructural changes significantly affect LA myocardial wall properties, such as relaxation and compliance, ultimately leading to the observed reduction of LA phasic functions assessed by

STE.^{40,47,48} An interesting finding of the present study was the observed lower PALS in dogs with MMVD class B1 compared with that of healthy dogs, whereas LA areas and almost all STE-derived variables were not different between these groups of dogs. This finding suggests a possible early effect of MR on LA structure and function that was only evident by strain analysis in this study. Although canine MMVD progresses in the absence of significant ventricular myocardial fibrosis,⁴⁹ PALS has been proven to highly correlate with LA fibrosis in human patients with MR.³⁸ At present, studies investigating the role of LA fibrosis in dogs in the early stage of MMVD are lacking. On the other hand, because dogs of group A were significantly younger than those of the other groups, it cannot be excluded that an age-related effect might also have influenced this finding. Contraction strain index, a parameter indicative of LA active systolic function, showed a transient increase from dogs of group A to those of group B1 followed by a progressive reduction. This supernormal pattern has already been observed in humans with mild MR, and it is likely secondary to Starling's law usually applied to the ventricles.³⁷ In fact, the atria also enhance their contractile function in response to an increased preload during the early phase of MR, but this function subsequently declines as the LV filling pressure increases.³⁷

In the present study, we recorded relatively higher values of PALS, PACS, and CSI in our control group compared with those recently reported in healthy dogs.³⁰ This difference might be explained by the different equipment and software used for the analysis in the 2 studies, and also by the different demographic populations. The dogs enrolled in that previous study had a relatively broad range of breeds, age, and BW, whereas dogs of the present study were comparably older and smaller in size. Although no association was found between STE variables and BW and age in the reported study,³⁰ we cannot exclude a possible BW and age effect on STE variables.

Some limitations of the present study must be highlighted. First, technical aspects of STE-derived variables should be considered, because the reliability of measurements might be questioned in dogs with extremely remodeled LA. In these cases, excessive dilatation of the pulmonary vein inlet and displacement of the interatrial septum might complicate the tracking procedure. These structures were excluded from tracking analysis, therefore reducing the risk of producing artifactual data among the global LA strain. Moreover, we documented a low intra- and interobserver variability in all groups of dogs, including those of group C + D, where the described anatomical modifications are common. Second, dogs affected by MMVD were older than those of the control group. However, given the acquired and progressive nature of MMVD, it is difficult to find a perfectly age- and BW-matched control group of dogs without some degree of MR, and age did not have any significant effect on STE-derived variables in the ANCOVA model. Third, a possible influence of cardiac medications on LA function could not be evaluated in

the present study. However, only 39 of 127 (31%) dogs with MMVD received cardiac medications at the time of enrollment, and different drug combinations and dosages were employed. Finally, LA mechanical properties were not assessed for functional impairment by gold standard imaging modalities, such as cardiac computed tomography and magnetic resonance. However, given the need of general anesthesia and the related risks (especially in dogs with decompensated MMVD), these procedures were not carried out for ethical reasons.

In conclusion, we described the use of STE for the evaluation of LA strain and other functional indexes in healthy dogs and dogs with MMVD. These variables can be effectively obtained also in dogs in the advanced stages of MMVD and are useful to track the progressive decline in LA function as the disease progresses.

Footnotes

- ^a iU22 ultrasound system, Philips Healthcare, Monza, Italy
^b iE33 ultrasound system, Philips Healthcare, Monza, Italy
^c QLAB quantification software version 9.1, Philips Healthcare, Monza, Italy
^d SAS version 9.3, SAS Institute Inc., Cary, NC
^e MedCalc version 12.6.1.0, MedCalc Software, Ostend, Belgium

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Off-label Antimicrobial Declaration: Authors declare no off-label use of antimicrobials.

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