

The role of multi-modality imaging for the assessment of left atrium and left atrial appendage: a clinical consensus statement of the European Association of Cardiovascular Imaging (EACVI), European Heart Rhythm Association (EHRA) of the European Society of Cardiology (ESC)

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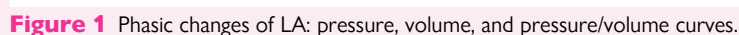
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The assessment of structural, architectural (tissue characteristics), functional, or electrophysiological changes associated with LA cardiopathy relies on the rational use of imaging modalities by appreciating their capabilities and limitations to derive clinically relevant information (*Table 1*).

Transthoracic echocardiography (TTE) is the primary imaging modality to evaluate LA size and function. Since LA enlargement is asymmetrical, anteroposterior LA diameter measured from the M-mode or 2D parasternal long-axis image can significantly underestimate LA size.^{8,9} Yet, for some specific clinical conditions, the cut-offs for LA dilatation remain based upon anteroposterior linear measurements, such as in risk stratification in hypertrophic cardiomyopathy (HCM).^{10,11} LA volume quantification by 2D echocardiography (2DE) and, preferentially, by 3D echocardiography (3DE) is the accurate way to evaluate LA size in clinical practice.¹² The measurement of LA volume by 2DE requires the acquisition of LA-focused apical views to maximize both the LA width and length in four- and two-chamber (should be similar in both views) to avoid foreshortening,¹⁰ because the LA and the LV are not co-axial.¹³ The bi-plane area-length method systematically yields larger LA volumes than the Simpson's bi-plane disc summation method as it assumes an ellipsoid shape.¹⁴ The Simpson's method is preferred over the area-length method for clinical use because of fewer geometric assumptions than the area-length method.⁵ LA volume calculated by 2DE correlates well with measurements obtained using 3DE, cardiac

Maximal LA volume has been the most clinically used parameter; yet, there is an important body of evidence supporting the role of phasic LA volumes (particularly LA minimal volume) and function. LA function may demonstrate alterations prior to volume changes.¹² The same dedicated 2DE views used for maximal LA volume calculation can be employed to obtain LA phasic function parameters. From the LA volumes, the total emptying volume (EV) is calculated as (LAVmax – LAVmin), the passive EV as (LAVmax – LAVpreA), and the active EV as (LAVpreA – LAVmin). Total emptying fraction (LAEF); (total EV/LAVmax), passive LAEF;

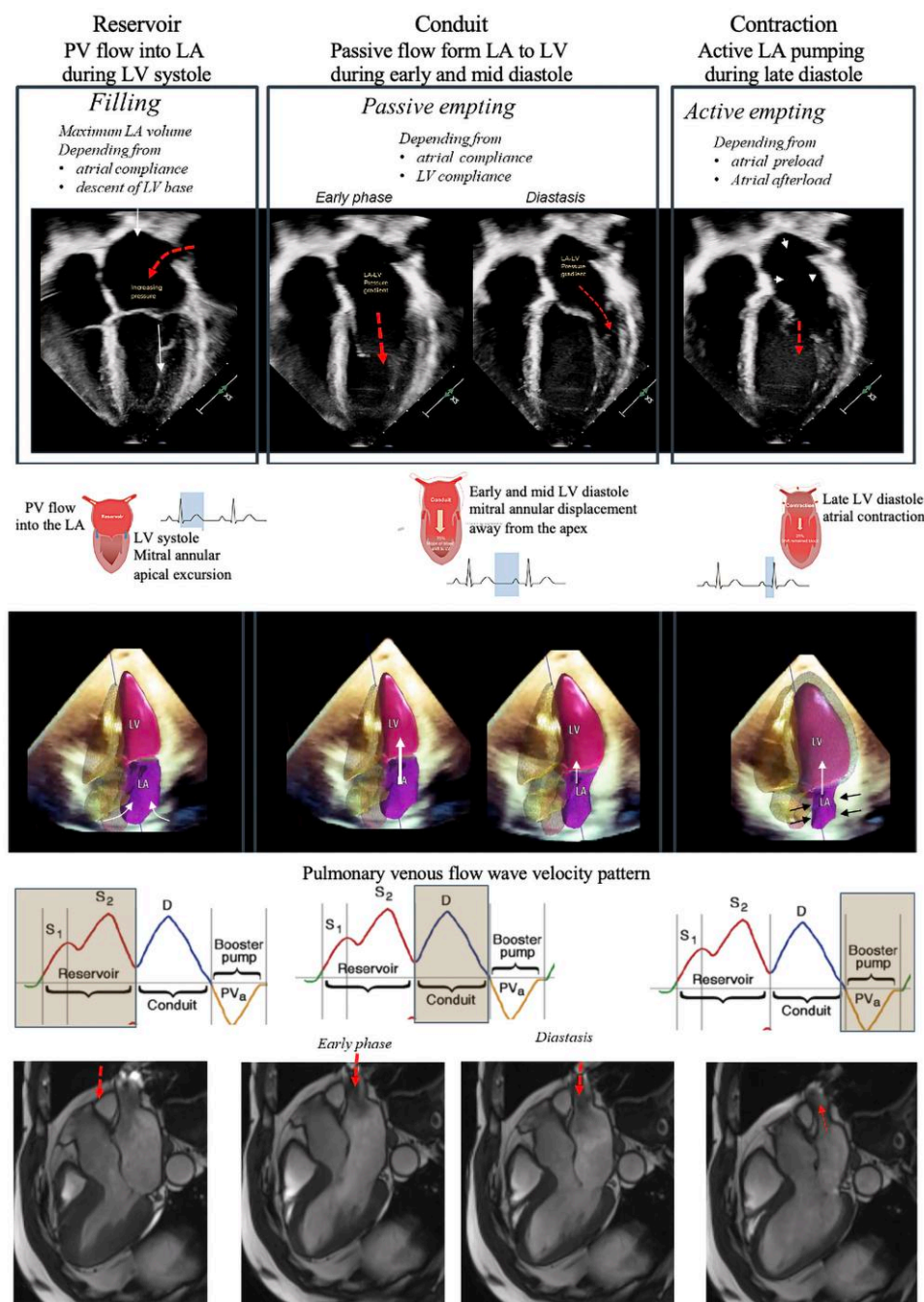


Figure 2 Translation of LA phasic function into imaging. Note the interaction between mitral annular descent by LV contraction, LA reservoir, atrial contraction, and PV flow.

(passive EV / LAVmax), and active LAEF; (active EV / LAVpreA) are calculated as indices of reservoir, conduit, and contractile function, respectively (Figure 3).¹³ Because indexing LA size to body surface area accounts for gender difference, only the indexed value should be reported.²²

Because of manual endocardial border tracing at three different time points of the cardiac cycle, assessment of LA phasic function by 2DE is time-consuming and prone to errors. LA function can be measured faster and automatically, without geometric assumptions by using dedicated software packages for LA volume quantitation from 3DE datasets (Figure 4).¹³ The limitations of 3DE at present include low spatial resolution and the need

for dedicated equipment and training. There is paucity of data regarding its incremental prognostic value compared to 2DE.

Mitral inflow velocity during atrial contraction (A wave) and late diastolic mitral annular velocity (a') are Doppler metrics of LA function. Blunted or absent A and a' waves during sinus rhythm are typical findings of atrial stunning. Of note, these parameters are load- and angle-dependent and present during sinus rhythm only. Speckle-tracking strain has been a highly sought-after technique for LA function quantification. It is semi-automated, less angle-dependent, and less affected by artefacts than tissue Doppler-based measurements. LA strain

Table 1 Goal-directed multi-modality imaging for LA and LAA

	TTE	TOE	ICE	CCT	CMR	PET
LA volume	+++ ^a	+	+	+++	+++	-
LA morphology	+++ ^a	-	+	+++	+++	-
LA phasic function	+++ ^a	-	-	++	+++	-
LA strain	+++	-	-	+	++	-
LAA morphology	-	++	++	+++	++	-
LAA function	-	+++	+	-	+	-
Pulmonary vein flow	++	+++	-	-	-	-
Pulmonary vein pattern	-	++	++	+++	+++	-
LAA thrombus	-	+++	+++	+++	++	-
LA fibrosis	-	-	-	-	++	++ ^b
Peri-atrial epicardial fat	+ ^c	-	-	+++	+++	-
Real-time image integration with EAM	-	-	-	+++	+++	-
Inflammation	-	-	-	-	-	++ ^d

Abbreviations: CCT, cardiac computerized tomography; CMR, cardiac magnetic resonance; EAM, electroanatomic mapping; ICE, intracardiac echocardiography; PET, positron emission tomography; TTE, transthoracic echocardiography; TOE, transthoracic echocardiography.

^aIdeally by 3DE.

^bPET with fibroblast activating protein inhibitors.

^cOnly thickness.

^dPET with fluorodeoxyglucose.

Table 2 Reference echocardiographic values of LA volumes¹³

	3D			2D
	Women	Men	Overall	Overall
LAVI _{max} (mL/m ²)	31 (27–45)	31 (19–52)	32 (28–36)	24 (21–28)
LAVI _{min} (mL/m ²)	10 (5–18)	11 (4–21)	10 (8–12)	8 (6–10)
LAVI _{preA} (mL/m ²)	18 (10–30)	18 (9–32)	18 (14–21)	14 (12–18)
Total LAEF (%)	68 (53–79)	66 (51–80)	67 (63–71)	67 (62–74)
Active LAEF (%)	40 (18–61)	41 (20–60)	41 (35–48)	46 (39–53)
Passive LAEF (%)	45 (22–60)	43 (23–61)	44 (38–49)	41 (32–48)

Values are given as median (Interquartile range). Abbreviations: LAVI, left atrial volume index; max, maximum; min, minimum.

assessment requires dedicated, focused image acquisitions to optimize lateral and temporal resolution. The use of dedicated software with an automated LA wall detection algorithm improves measurement reproducibility. The region of interest (ROI) should fit in the thin wall, accurately track the motion of the mitral annulus, and avoid the strong signals of the adjacent and stationary pericardial tissue, which—if included—may underestimate strain values.²³ LA function parameters [reservoir (LASr), conduit, and contraction strain] are computed as LA time-strain curves over the cardiac cycle (Figure 5). Analysis, reporting, and interpretation should follow the EACVI/ASE/Industry Taskforce consensus statement on LA strain analysis.²⁴ The reference timing of zero strain should be set at end-diastole (R–R gating). The alternative method is when zero strain is set at the onset of atrial contraction (P–P gating). Importantly, however, the R–R gating is less prone to errors

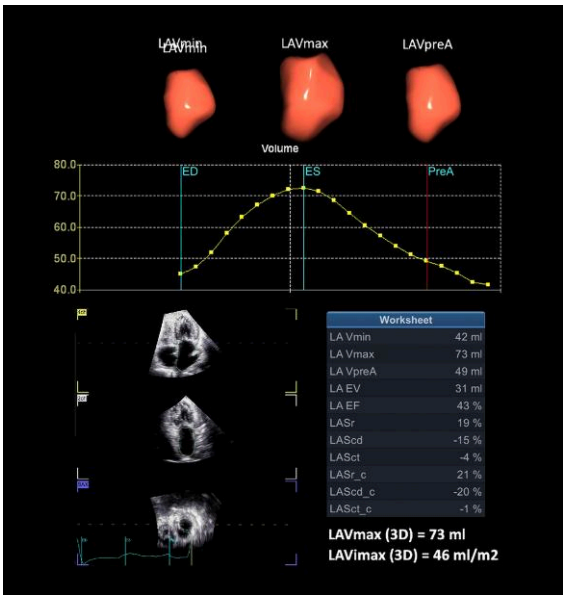


Figure 3 Phasic volumes by 3DE.

than the P–P gating and can be used during AF. Yet, high reproducibility (intraclass correlation 0.93 and 0.90, respectively) of both methods was shown in the MASCOT-HIT study.²⁵ Strain rate during ventricular systole, early diastole, and late diastole correspond to reservoir, conduit, and contractile function, irrespective of the R–R or P–P gating. Of note, LA load impacts volume and strain measurements regardless of the method used. Normal LA phasic strain values were reported in the healthy population from a meta-analysis including 40 studies in 2017 and from the NORRE study in 2018^{19,26} (Table 3). Additionally, reference ranges of LA phasic function by strain were published from 1329 healthy participants in the HUNT study (analyses performed by General Electric HealthCare EchoPAC system). Accordingly, bi-plane mean $\pm$ 2SD reference values are as follows, in females and males, respectively: %LASr, 33.2 (17.2–49.2), 32.7 (16.0–49.3); %LA conduit strain, –17.0 (–30.8 to –3.3), –15.7 (–29.1 to –2.3); %LA contraction strain, –16.2 (–25.4 to –7.0), –17.0 (–26.9 to –7.0).²⁷ Combining early diastolic mitral inflow (E) and annulus (e') velocities with LASr enables obtaining an index for the estimation of LA stiffness by the formula: $E/e'/LASr$.²⁸

Advice table 2 Use of TTE

Evaluation of the LA by TTE requires the acquisition of atrial-focused views.	
It is advised to assess LA size by volume (bi-plane 2D or 3D) quantification and to use the same tool for intra- and inter-individual comparisons.	
LA phasic volumes and strain by TTE are the mainstay of assessing LA function.	
Volumetric cut-offs obtained from different echo techniques cannot be used interchangeably.	
R–R gating is advised for the assessment of LA phasic function by 2D strain.	
LA size estimation should not be limited to M-mode or 2D linear measurements.	

Transoesophageal echocardiography

Transoesophageal echocardiography (TOE) is well suited to image the LA and the LAA, given their proximity to the oesophagus. However, LA sizing is performed by TTE rather than TOE, because the ultrasound scan sector cannot entirely encompass LA on TOE. Exclusion of thrombi, search of cardioembolic sources, characterization of the PVs before AF ablation, procedural planning, and guidance for LAA occlusion are the most common indications for imaging LA and LAA by TOE.²⁹ Although TOE is semi-invasive, it is safe in renal insufficiency. It provides adequate image quality in AF and offers information about associated cardiac pathologies and haemodynamics.

Technical considerations

Multiple TOE planes are used to explore the LA and LAA from the mid-oesophageal view. With 2D probes, LAA can be easily visualized between 45 and 90° with counterclockwise rotation and gentle anteflexion of the transducer. Once the LAA is visualized, it is centralized on the screen, and a thorough evaluation is performed by sweeping from 0 to 135°. However, 3D probes allow visualization of bi- or tri-plane simultaneously which facilitates the sweep process (Figure 6). A 3D zoom acquisition with a wide sector (since high temporal resolution is not specifically required for atrial structures) and excluding the atrial roof in the nearfield (avoiding obstruction of the view) is generally performed for a live assessment and further cropping. From the 3D zoom acquisition, a live ‘en face’ view of the LAA orifice can be obtained. Further perpendicular cropping and orientation of the 3D dataset enable the characterization of the LAA morphology. Photo-realistic rendering with light adjustments or CCT-like rendering with improved transparency (Glass) has also been recently introduced to improve the qualitative assessment by highlighting the tissue–blood interface (Figure 7).³⁰

The TOE also provides anatomic and functional information about PVs. Notably, TOE underestimates PV dimensions, especially for the inferior veins.³¹ The left upper PV is visualized in its long axis adjacent to LAA with further counterclockwise rotation between 45 and 110°. Visualization of the left lower PV requires slight vertical manipulation or further angulation to 120°. The right upper PV can be visualized after clockwise rotation of the probe with anteflexion from either 0, 45, or 120–135° (on the left and right side of the screen, respectively) adjacent to the IAS and superior vena cava. The right lower PV is the most difficult to align with the Doppler beam and is best seen from the extreme clockwise rotation of the probe at 0° without anteflexion immediately inferior to the right upper PV. PV Doppler interrogation shows systolic (S), diastolic (D), and atrial contraction (A) waves during sinus rhythm (Figure 8A–D). PV flow velocities are relevant for the assessment of LA pressure, mitral regurgitation (MR) severity, and PV stenosis after PV isolation. The left and right PV orifices are widely separated, and the pyramidal 3D data set cannot include them all in a single image. From the best 2D views obtained, the left and right PV orifices can be seen 2 by 2 by 3D zoom acquisition from an optimal orthogonal display, and rotation of the dataset as shown in figure (Figure 8E and F). The left PVs are adjacent to LAA, and the right PVs run adjacent to the IAS.

LAA function

Flow in the LAA is visualized using colour Doppler with a low Nyquist limit, and flow velocities are measured by pulsed-wave Doppler. Four phases have been described: emptying (contraction) velocity, filling velocity, a biphasic systolic reflection wave, and an early diastolic emptying wave²⁹ (Figure 9). Emptying velocity ranges from 63 ± 29 to 83 ± 25 cm/s and filling velocities range from 54 ± 17 to 61 ± 18 cm/s. Velocities can decrease because of high LA

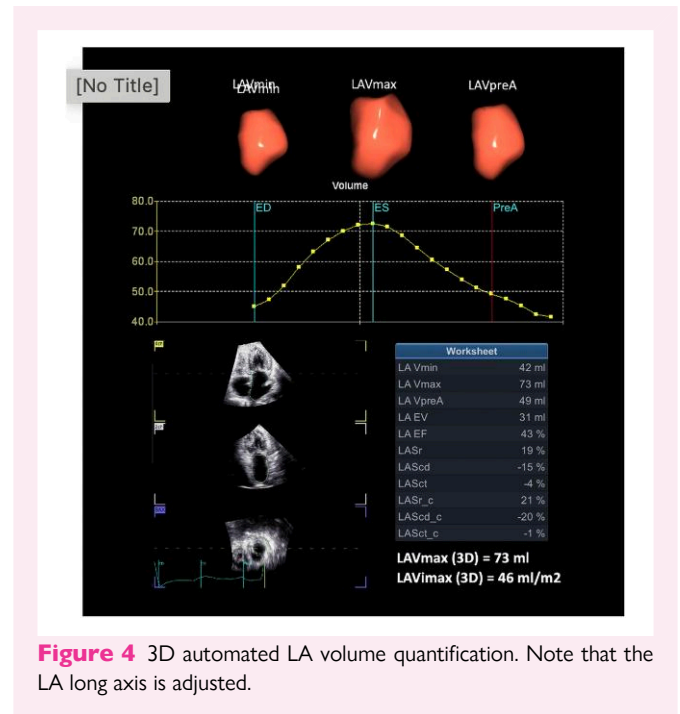


Figure 4 3D automated LA volume quantification. Note that the LA long axis is adjusted.

pressure. A sawtooth pattern with low velocities is observed during AF. Velocities <40 cm/s in sinus rhythm are associated with an increased risk of stroke.³²

Advice table 3 For the use of TOE

- Multipanar imaging with or without 3D is indispensable for complete visualization of LAA morphology.
- TOE is the preferred technique for the assessment of LAA, function, and thrombus.
- TOE is the preferred technique for the assessment of PV flow.
- The use of TOE is inappropriate for the assessment of LA size or volume.

CMR

CMR allows a comprehensive 3D evaluation of LA anatomy, structure, and function. Its potential to reveal the extent of atrial fibrosis by late gadolinium hyperenhancement (LGE) imaging is an advantage.³³

Assessment of morphology

CMR steady-state free-precession (SSFP) cine images provide excellent blood-endocardium and epicardium-fat contrast, allowing good delineation of the atrial endocardial borders and LA size. This allows precise measurement of LA volumes, either using a bi-plane area–length method from two- and four-chamber cine images (Figure 10A) or using the Simpson’s disc summation method on a stack of adjacent short-axis images from the atrioventricular ring to the roof of the LA. Since LV and LA are not co-axial, the LA volumes derived from the bi-plane area–length method may underestimate the true LA size because the long-axis cine images are normally aligned to the LV axis during

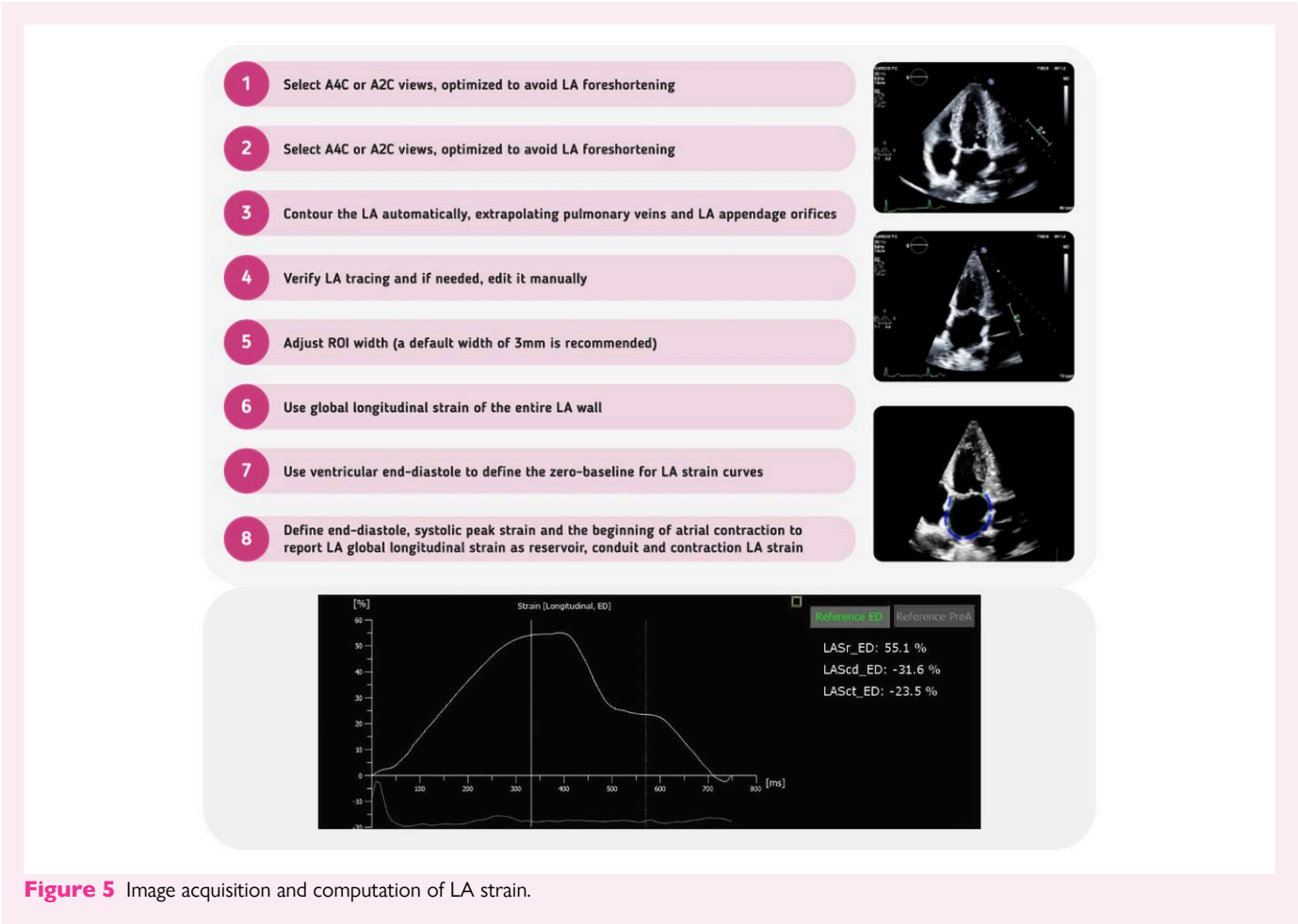


Table 3 Reference values of LA phasic strain (NORRE study)¹⁹

	All ages	Age 20–40 years	Age 40–60 years	Age >60 years
LAS reservoir	42.5 (36.1–48.0)	46.8 (42.3–52.4)	40.9 (35.4–46.1)	35.5 (30.9–41.9)
LAS contraction	–16.3 (–12.9 to –19.5)	–15.6 (–11.9 to –19.0)	–16.3 (–13.2 to –19.6)	–16.8 (–13.6 to –21.4)
LAS conduit	–25.7 (–20.4 to –31.8)	–30.6 (–26.8 to –36.5)	–24.1 (–19.7 to –29.3)	–18.6 (–14.7 to –22.6)

Values are given as median (interquartile range).

acquisition. The Simpson's method is more precise and overcomes the geometric assumption and limitations of the area–length method; however, it requires additional scanning time to acquire the LA short-axis stack. The reference values obtained from the two- and four-chamber cine images are given in Table 4.³⁴

More precise evaluation of atrial morphology, especially of the LAA and PV, can be achieved by using 3D angiographic (MRA) images (Figure 10C) which can be acquired either using free breathing of navigator gated MRA after gadolinium injection or using non-contrast balanced SSFP MRA.³⁵ Like CCT, these images can be merged with 3D electro-anatomical maps using landmarks or surface registration for AF ablation³⁶ (Figure 10E). Evaluation of the anatomy of the PV pre-ablation and PV stenosis post-ablation is also feasible with CMR; however, less data is available as compared to CCT.³⁷

CMR allows understanding of tissue characteristics and detection of atrial fibrosis by high-resolution 3D LGE imaging³⁸ (Figure 10E). LGE on the LA wall has been detected in several conditions such as AF,^{39,40} mitral valve diseases,⁴¹ and cardiac amyloidosis (CA).⁴² Yet, assessment of atrial LGE is performed visually at best by experienced centres without a consensus for quantification and is not possible if the image quality is suboptimal. Variability among observers has been a significant disadvantage.³⁸ Further work is required to standardize atrial LGE imaging to enable its widespread clinical use.

Cine and 3D MRA allow also the assessment of LAA size and morphology and measurement of its ostial diameter and depth. For assessing LAA thrombus, CMR with inversion time myocardial delayed enhancement acquisition has the highest accuracy (99.4%) followed by contrast-enhanced (97.6%) and cine CMR 93.9%) tools.⁴³ Hence,

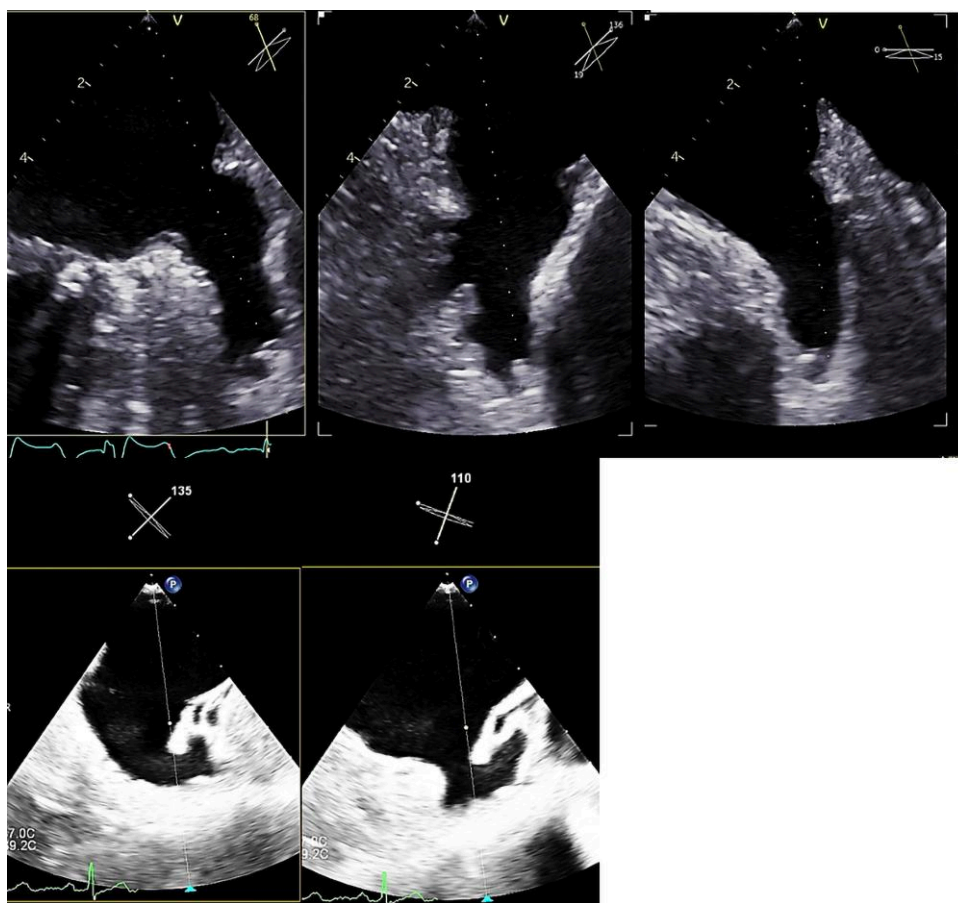


Figure 6 Multiplane images of the LAA showing different morphologies.

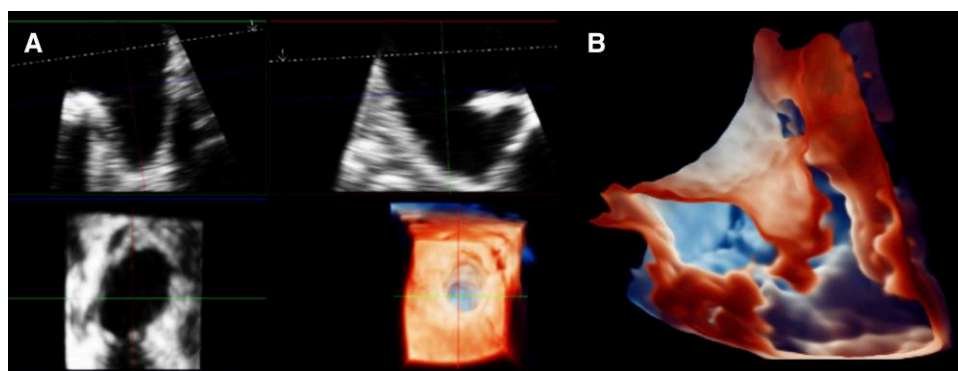


Figure 7 (A) Zoom-mode acquisition and multiplane display of LAA morphology by cropping and re-orienting the 3D data set with 3D photo-realistic rendering. (B) Glass view of the LAA.

CMR is an alternative to TOE or CCT in centres having adequate experience in LAA image acquisition and interpretation, depending on the resources.

Finally, CMR (T1-weighted or cine SSFP images) can accurately quantify the volume and area of pericardial adipose tissue.⁴⁴ A recent

meta-analysis showed that LA epicardial adipose tissue (EAT) thickness was a strong parameter associated with the risk of AF recurrences after catheter ablation.⁴⁵

CMR may suffer from artefacts in case of AF with high heart rate and devices (particularly intracardiac defibrillator).

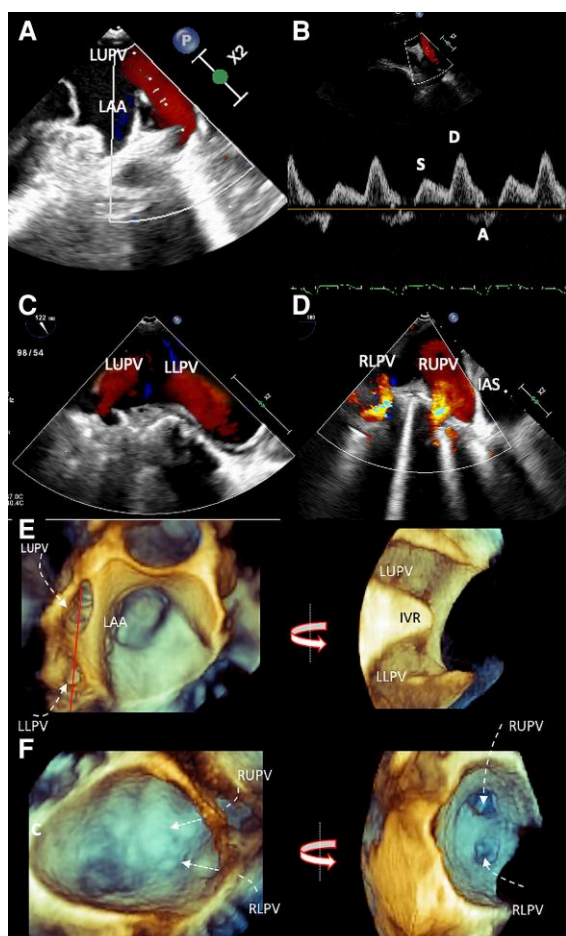


Figure 8 PVs by TOE. (A) Left upper PV, 60°, colour Doppler; (B) left upper PV, pulsed-wave Doppler; (C) left upper and lower PVs, 120°, colour Doppler; (D) right upper and lower PVs, 0° clockwise rotation; (E, F) the left and right PVs, 3D-rendered image.

Assessment of function

LA function can be assessed with CMR by computing LAEF from measurements of maximum and minimum volumes and phasic LA volume–time curves from the cine images (Figure 10B) (Table 4).^{34,46,47} Recent technology enabled LA strain quantification by automated feature tracking methods from two- and four-chamber cine images (Figure 10D). A recent meta-analysis showed that feature tracking vendor matters for the heterogeneity of measurements rather than the CMR vendor, sex, and age.⁴⁸ The pooled mean values of LA phasic strain are presented in Table 5. There are promising advances in measuring peak velocity and vorticity by 4D flow imaging paving the way for the assessment of atrial haemodynamics (Figure 10F).

CCT

Due to its excellent spatial resolution, CCT plays a pivotal role in defining the morphology of the LA, LAA, PVs, and function of the LA and in guiding electro-anatomical mapping. For evaluating LA, PVs, LAA morphology, and epicardial fat, a single arterial phase acquisition with ≥ 64 -slice CCT with ECG triggering is required. Prospective ECG triggering is used in patients with sinus rhythm and low heart rate whereas retrospective ECG triggering is appropriate in patients with high and non-stable heart rates.⁵⁰ In patients who are in AF during the scan, acquisition is more challenging, but the introduction of more recent technology allows adequate image quality even during AF.^{51,52} Figure 11 shows a typical 3D LA reconstruction with CCT. An additional delayed scan after contrast injection is mandatory for ruling out LAA thrombus.⁵³ While the time to delayed images varied in studies, most were acquired 30–180 s after the initial images.

Assessment of morphology

CCT has a higher spatial resolution than CMR and is a well-established technique to evaluate LA and LAA morphology and volume and PV patterns, to rule out LAA thrombus, and to detect peri-atrial adipose tissue.

CCT systematically detects higher LA volumes compared to 2DE and CMR because of several reasons.⁵⁴ First, due to higher temporal resolution, the proper image reconstruction windows for

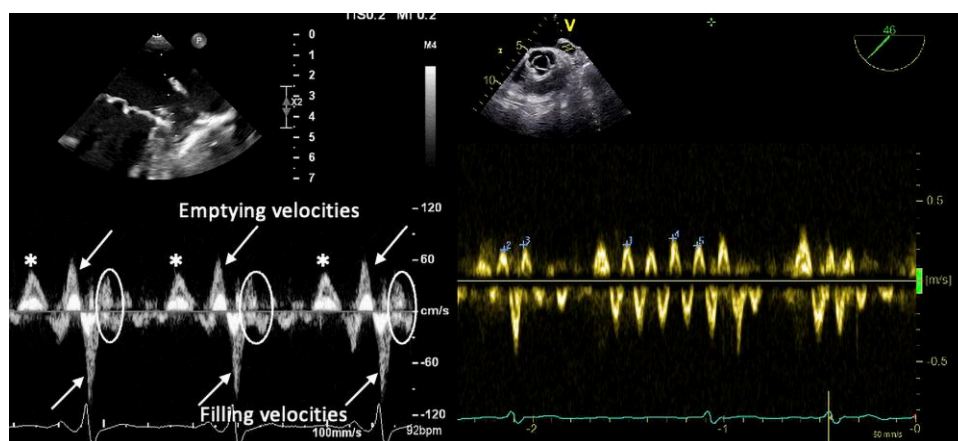


Figure 9 Normal quadriphasic wave pattern of LAA flow. Early diastolic emptying (asterisk), systolic reflection (circles) waves.

Table 4 Reference values of LA by CMR³⁴

	Men		Women	
	Mean ± SD	Lower–upper limits	Mean ± SD	Lower–upper limits
Bi-plane LAVImax (mL/m ²)	38 ± 11	17–59	39 ± 11	17–61
Bi-plane LAVImin (mL/m ²)	14 ± 5	3–24	13 ± 5	4–23
Bi-plane LA EF (%)	62 ± 8	46–77	63 ± 8	48–78
Simpson LAVImax (mL/m ²)	41 ± 8	24–57	44 ± 8	28–60
Simpson LAVImin (mL/m ²)	19 ± 5	9–28	19 ± 4	11–27
Simpson LA EF (%)	54 ± 8	38–70	57 ± 6	45–69

Abbreviation: SD, standard deviation.

Table 5 LA strain reference values by CMR feature tracking⁴⁸

	Mean	95% CI
LAS reservoir %	34.9	29.6–40.2
LAS conduit %	–21.3	–16.6 to –26.1
LAS contraction %	–14.3	–11.8 to –16.8

Assessment of function

With ECG-triggered acquisition, functional series can be obtained in order to perform LA strain quantification. However, currently, data regarding LA strain assessment using functional CCT images are scarce. LA strain is usually obtained by tracing endocardial LA borders on multiple apical two-chamber views excluding LAA and PVs. Szilveszter *et al.*⁶² demonstrated excellent intra-observer reproducibility for this approach with an intraclass correlation of 0.95, a correlation coefficient of 0.87, and 5.6 points of underestimation as compared to 2D echo. Yet, relatively low temporal resolution and radiation exposure limit the use of CCT for LA strain quantification in clinical practice.

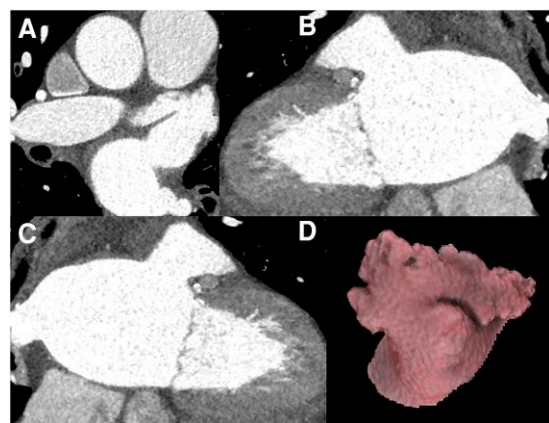


Figure 11 LAA. (A) Axial, B) sagittal oblique, C) coronal oblique views, and D) 3D volume-rendered image.

method to assess the PVs, although no differences are described between CCT and CMR in terms of diagnostic accuracy of PV pattern (Figure 13).

CCT is accurate and reproducible to assess epicardial fat tissue (EFT) by either manual or semi-automated volume quantification. This latter algorithm defines all structures with contrast attenuation ranging between -195 and -45 HU as adipose tissue.⁵⁹ A volumetric quantification is advised rather than area measurements.⁵⁹ CCT also allows visualization of the peri-atrial anatomy for safe guidance of procedures and helps to avoid phrenic nerve injury or atrio-oesophageal fistula.^{60,61}

Advice table 4 Use of CMR and CT

CMR is the gold standard for the quantification of peri-atrial adipose tissue volume.

CCT is advised for assessing the anatomy of PVs.

CMR and CCT are alternatives to echocardiography for the assessment of LA, LAA morphology, volume, and thrombus. Using the same imaging modality is strongly advised for intra- and inter-subject comparisons.

CMR can be used as an alternative to CCT for assessing the anatomy of PVs.

CMR with automated feature tracking can be used for LA strain quantification.

CMR with LGE can be used for visualization of LA wall fibrosis with good-quality imaging, by the experts.

Nuclear imaging

Molecular positron emission tomography (PET) imaging as a tool to assess atrial inflammation and fibrosis is in early stages. Increased atrial 18F-FDG uptake has been shown in patients with AF and in those with sarcoidosis as a predictor of subsequent AF.^{63,64} More recently, 68Ga-fibroblast activation protein inhibitor (FAPI-PET) was used to detect increased fibroblast activation in the atria of patients with AF or after PV isolation.^{65,66} Atrial activity is rarely reported but seems to be clinically relevant. Oncologic patients frequently have atrial uptake on their PET/CT which is also linked to the risk of AF and pro-thrombotic state with cardio-oncologic consequences.⁶⁴ Further investigations are awaited to consolidate clinical implications.

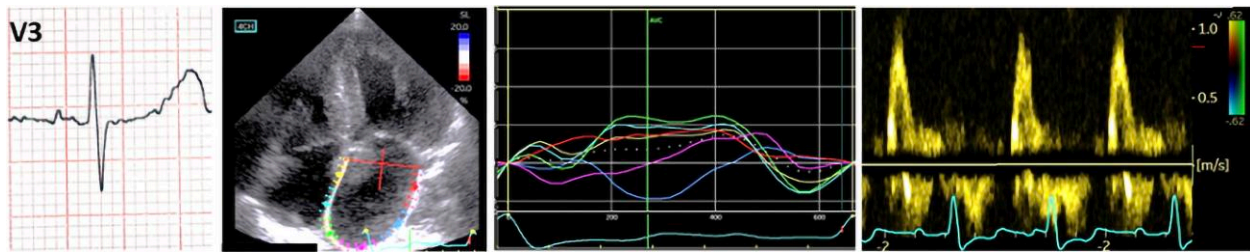


Figure 14 Atrial electromechanical dissociation in CA. Poor LA strain, absent LA contraction strain, and A wave, despite sinus rhythm.

abnormal LASr is associated with dyspnoea, NYHA class, and HF hospitalization and is a useful adjunct to the evaluation of DD and estimation of LV filling pressure algorithms in indeterminate cases.^{73,79}

HF with mildly reduced and reduced EF

LA enlargement in HF with mildly reduced EF (HFmrEF) and HF with reduced EF (HFrEF) is associated with adverse cardiovascular events.⁸⁰ However, the impact of LASr on outcomes, in these patients, has been less studied. The best relationship between LASr and filling pressure is found in patients with reduced systolic function.^{77,81} HFmrEF (EF = 41–49%) by definition needs the presence of symptoms and/or signs of HF. The presence of increased LAVI, elevated natriuretic peptides, and evidence of structural heart disease make the diagnosis more likely, but are not mandatory for diagnosis.⁸²

Ischemic heart disease

Patients with ischaemic heart disease or after myocardial infarction (MI) make up the largest Stage B HF group. Accordingly, LA function and remodelling could be a marker of abnormal cardiac function with a diagnostic value. Additionally, LA function has been shown to predict HF hospitalizations after MI⁸³ and was incremental to LAVI.⁸⁴ A recent large CMR study showed that LAEF is independently associated with increased mortality in patients with ischaemic cardiopathy (LVEF <50%) even after adjusting for infarct size and MR severity.⁸⁵ LASr, assessed within 48 h of acute MI, was associated with the composite outcome of death and HF⁸⁶ and provided incremental value to LAVI in patients treated with percutaneous coronary interventions.⁸⁷ Data from multicentre prospective CMR studies [AIDA STEMI (NCT00712101) and TATORT NSTEMI (NCT01612312)] also showed that LASr (cut-off of 18.8%) is an independent predictor of outcome and incremental to LVEF, GLS, microvascular obstruction, and infarct size.⁸⁸ LAVI predicted morbidity and mortality after acute MI as well.^{89,90} However, LA dilatation reflects a chronic process therefore may not be an ideal marker shortly after an acute MI in contrast to the indices of LA function that correlate more strongly to LV filling pressure after acute MI. Additionally, reduced LASr was shown to predict an increased risk of new-onset AF after coronary artery bypass graft surgery.⁹¹

Athlete's heart

LA dilatation is triggered by the increase in preload during athletic training as an adaptive mechanism.^{92,93} Age, type of sport, and duration and intensity of training influence the degree of atrial remodelling. LAVI is associated with higher cardiorespiratory fitness and maximal oxygen consumption during exercise in both men and women.⁹⁴ A systematic review including 7189 elite athletes and 1375 controls described increased LAVI in athletes with an upper limit of normal 35.8 mL/m² compared to <34 mL/m² in the general population.⁹⁵ Spencer *et al.*⁹⁶ reported LAVI exceeding 48 mL/m² in 40% of male and 32% of female

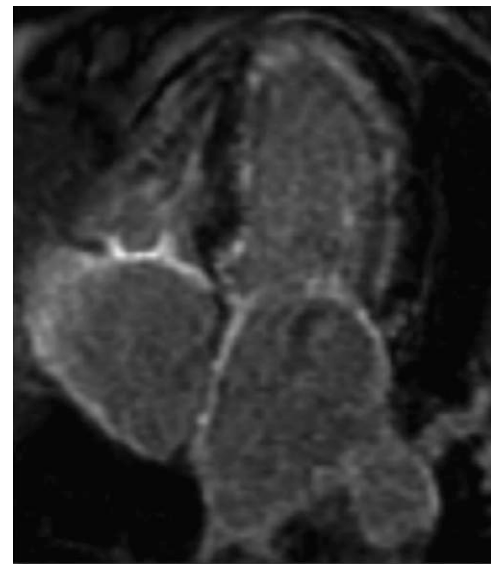


Figure 15 Diffuse LGE on the LA wall in CA.

athletes. Importantly, there is a balanced adaptation with global remodelling in both atria and ventricles. Despite LA enlargement, E/e' remains normal by means of increased LA and LV compliances and bradycardia and maintains LA pressure within normal range.⁹⁶ Conflicting evidence from relatively small cohorts exists about reversal of LA dilation with detraining.^{93,97} In athletes, LASr is either preserved or mildly reduced (39%; 95% CI, 38–41%) compared to untrained controls,⁹² and LA active emptying is lower in athletes (17%; 95% CI, 16–19%). Athletic atrial remodelling seems to be dependent on the intensity of training.^{98,99} Adaptation of phasic volume changes during exercise enables distinction between physiological and pathological atrial remodelling.¹⁰⁰ Moderate exercise appears to protect against AF, whereas strenuous exercise increases the risk of AF which is postulated to be mediated by atrial dilatation, vagal tone, exercise-related adrenergic stimulation, and augmented LA pressure during exercise.^{101,102} Increasing intensity and duration of athletic training leads to atrial enlargement and reduced atrial strain; only subtle further changes occur with AF. Therefore, prediction of AF in athletes by LA volume and strain is challenging as evidenced by conflicting results.^{103–105}

HCM

LA remodelling is promoted by impaired LV filling and raised LA filling pressures in HCM. HCM may also cause direct LA cardiopathy as



Figure 16 LA strain in AF. Note the lack of contraction; only the reservoir strain can be quantified which is significantly reduced.

evidenced by reduced passive and active LA emptying in the preclinical stage with positive genotype but without evident LV hypertrophy.¹⁰⁶ LA imaging and identification of AF risk are important because HCM is associated with a five-fold higher risk of AF incidence as compared with the general population and an increased rate of cardioembolism.¹⁰⁷ Increased LA volume, reduced atrial EF, and reduced LASr have been found to predict incident AF in the HCM populations.^{108,109} A high burden of atrial LGE on CMR was reported in patients with HCM and AF.¹¹⁰ Adverse LA remodelling in HCM has been shown to be a marker of poor outcome.^{111,112} LA diameter is a component of the sudden cardiac death risk scoring system in HCM patients as validated in 2014.¹¹³ The utility of more novel LA metrics has not been tested in identifying sudden cardiac death risk in large cohorts. Treatment of HCM is associated with LA structural and functional changes. Hegde *et al.*¹¹⁴ documented reductions in LA volumes and improvement in LV diastolic function and natriuretic peptide levels after treatment with mavacamten. Finally, regarding the controversy of exercise training in HCM, a similar LAVI increase was observed with competitive exercise in athletes with and without HCM.¹¹⁵

CA

Both primary LA cardiopathy from amyloid accumulation-mediated damage and secondary involvement due to increased LV filling pressure, MR, and AF occur in CA.¹¹⁶ Amyloid infiltration typically increases the thickness and stiffness of the atrial wall and IAS preventing excessive dilatation. Consequently, deformation-based parameters become more relevant than LA size for risk stratification in this population. Poor LASr and poor or absent LASct are typical findings in CA.¹¹⁷ The increase in LA stiffness can be estimated by the ratio $E/e'/\text{LASr}$.²⁸ During ventricular systole, the LA acts as a non-distensible (stiff) reservoir causing increased LA pressures and reducing the energy stored in the walls which affects the conduit phase. Finally, a lack of atrial mechanical contraction can be observed in a proportion of the patients

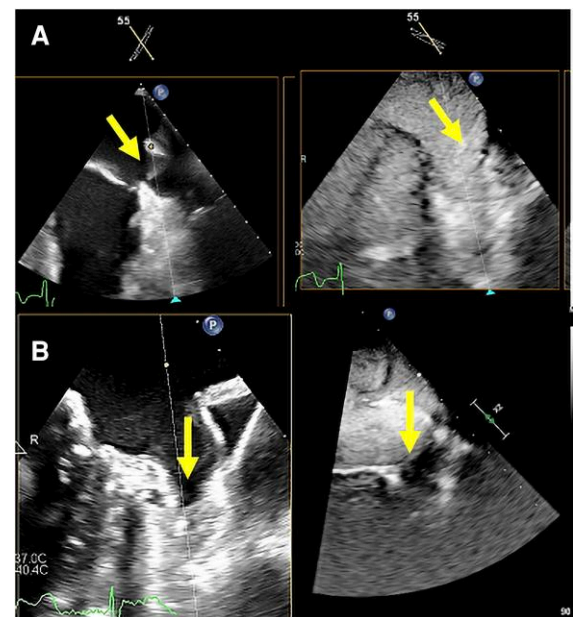
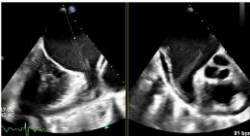
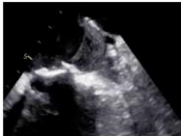
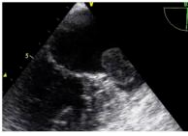


Figure 17 Ultrasound contrast for LAA opacification: (A) artefact mimicking thrombus, washing out with contrast, (B) thrombus producing a filling defect with contrast.

despite sinus rhythm, i.e. atrial electromechanical dissociation as a distinct feature and poor feature and poor prognosticator in CA (Figure 14).¹¹⁶ LA strain is independently associated with high thrombotic risk in patients with CA.¹¹⁸ LGE due to amyloid deposition or fibrosis in the LA wall can be detected by CMR⁴² (Figure 15) and is

Table 7 Clinical characteristics of SEC, sludge, and thrombus⁵⁶

	SEC	Sludge	Thrombus
Prevalence	 ≈50%	 1–14%	 13%
Echocardiographic characteristics	Smoke-like echogenicity with variable density. Grade 1: minimal dynamic echogenicity in the LAA or sparsely distributed in the LA; transient during cardiac cycle; Grade 2, swirling pattern with similar distribution to Grade 1; Grade 3, constantly detectable dense swirling pattern in the LAA that spills into the LA with less dense intensity; Grade 4, very slow swirling dense smoke-like echoes in the LAA, extending with similar density into the LA. Full, opacification with contrast, no filling defect with colour Doppler^a	Echo density with viscid gelatinous features but without a solid component. Opacification with swirling contrast, no filling defect with colour Doppler^a	Echo dense mass with margins and motion distinct from the atrial wall. Filling defect with colour Doppler^a, echo-free area with contrast
Thromboembolic risk	↑	↑↑	↑↑↑

^aLow Nyquist limit, 25–35 cm/s.

Table 8 Imaging markers of cardioembolic risk

TTE	TOE	CMR	CCT
LA volumes	LAA emptying velocity SEC in LA, LAA	LA volumes	LA volumes
LA strain	Sludge/thrombus in LA or LAA	LA fibrosis	SEC in LA, LAA
SEC in LA	LAA non-chicken wing morphology	LA strain	LA or LAA thrombus
Thrombus in LA	PFO	LA 4D flow	LAA non-chicken wing morphology

associated with reduced LA function.^{119–122} LA cardiopathy holds diagnostic¹²² and prognostic significance for all-cause mortality and increased risk of AF development and cardioembolic events in CA.^{118,122}

MR

LA dilation is an adaptive response to volume overload in patients with progressive MR.^{123,124} Furthermore, enlarged LAVI identifies individuals at increased risk of mortality, independent of the severity of MR or AF.¹²⁵ The 2021 European Society of Cardiology (ESC)/European Association of Cardiothoracic Surgery (EACTS) Guidelines for the management of valvular heart diseases recommend early surgical mitral valve repair in low-risk asymptomatic patients with severe primary MR when LAVI ≥ 60 mL/m² or LA diameter ≥ 55 mm.¹²³ In addition to LA dilation, reduced LASr has been independently associated with all-cause mortality in patients with significant primary and secondary MR and has shown incremental prognostic value over LAVI and LV GLS.^{126,127} LA fibrosis that occurs in the process of MR also reduces LASr.¹²⁸ The impact of mitral valve repair on the reversibility of LA fibrosis is currently investigated by LGE CMR (NCT05345730).⁴¹ In atrial functional MR, which occurs most commonly in the setting of chronic HFpEF or AF, LA dilatation is deemed to be the main driver of MR through annular dilatation.¹²⁹ LA reverse remodelling after mitral valve repair is a favourable prognosticator

but depends on several factors including pre-operative LAVI, MR severity, post-operative trans-mitral pressure gradient,¹³⁰ and intrinsic atrial cardiopathy. Recently, bi-leaflet prolapse was found to be associated with reduced LA function regardless of MR severity, suggesting a primary cardiopathy in these patients.¹³¹

MS

The pressure overload in MS promotes excessive dilatation of the LA with decreasing deformability, compliance, and contraction. In rheumatic MS, rheumatic atrial cardiopathy further exacerbates LA enlargement leading to one of the largest LAs observed in humans. The assessment of LA remodelling in MS includes LA size, EF, emptying fraction, PV flow patterns, LAA function, and LA deformation. From a clinical standpoint, LA compliance rather than the size is instrumental for mitigating pulmonary hypertension and increasing stroke volume downstream, whereby modulating symptoms in MS.¹³² LA reservoir function, quantified by longitudinal strain, reflects LA compliance and is a function of pure MS in young subjects. Of note, concomitantly reduced LV compliance would affect LA compliance, LASr, conduit, and pump strain in the elderly⁷⁵ which is typical for degenerative MS with mitral annular calcification. LA dilatation and reduced LASr predict symptoms, hospitalizations, valve intervention, recurrence of functional tricuspid regurgitation after tricuspid valvuloplasty,

and AF and thromboembolic complications.^{133–140} Anticoagulation with vitamin K antagonists should be considered if LAVI exceeds 60 mL/m² in patients with rheumatic MS even in sinus rhythm according to the 2021 ESC/EACTS Guidelines for the management of valvular heart disease.¹²³ Importantly, LASr and LA compliance were shown to improve following balloon mitral valvuloplasty, and this improvement translates into functional capacity.^{141,142} Similarly,

LAA contraction was shown to improve after balloon mitral valvuloplasty.¹⁴³

AF

Atrial contraction is abolished during AF. Atrial volume increases, while LA reservoir function decreases (Figure 16). LA remodelling and fibrosis

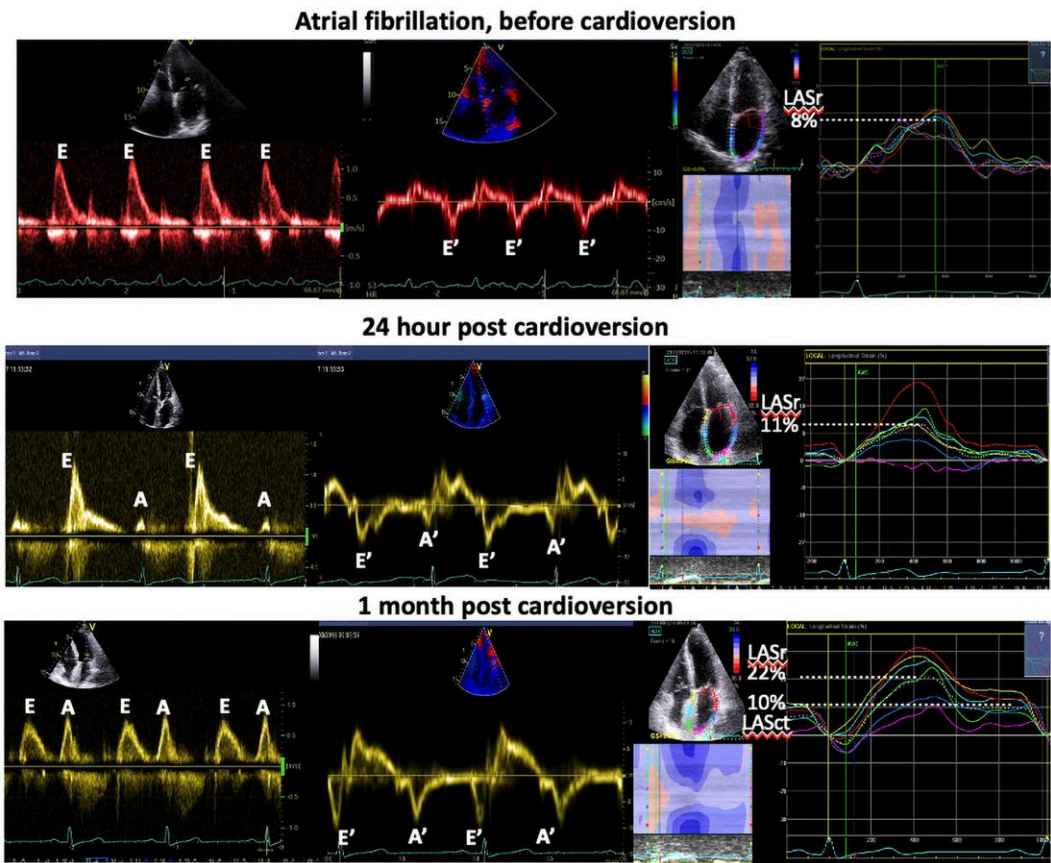


Figure 20 LA function before, early after, and late after cardioversion. Note the regeneration of LA contraction (A, a', LASct).

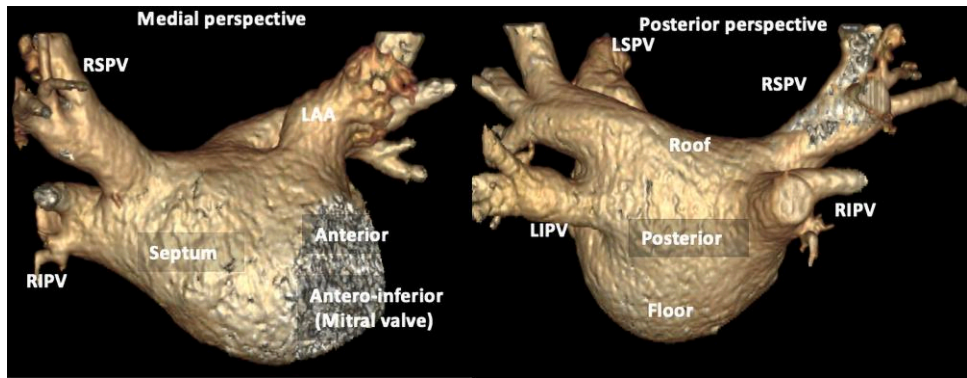


Figure 21 Demonstration of different parts of LA from a 3D volume-rendered reconstruction by contrast CCT performed for the assessment of PVs prior to PV isolation.

contribute to the perpetuation of AF.^{144,145} Dilatation of the LA predicts the development of AF in the elderly and its predictive value is incremental to linear measurements.¹⁴⁶ Likewise, LA enlargement predicts AF recurrence after radiofrequency ablation or cardioversion.¹⁴⁷ Total atrial conduction time which is the time interval from the onset of P wave to atrial contraction (tissue Doppler d') has been considered as a marker of LA fibrosis and shown to predict AF recurrence in patients following rhythm control strategy.¹⁴⁸

LASr is an early marker of altered structure and impaired function.¹⁴⁹ In AF, measurements are averaged from five consecutive cycles. Only LASr can be computed during AF.^{24,25} LASr predicted AF development in patients at risk in the general population,¹⁵⁰ in patients undergoing cardiac surgery, and in patients after a first stroke while in sinus rhythm.^{133,151,152} Hence, LASr has the potential to improve risk stratification for AF development and monitoring strategies.¹⁵³ From a prognostic standpoint, improvement in LASr is associated with a higher rate of sinus rhythm maintenance after cardioversion or ablation.^{154–156} Furthermore, an inverse relationship has been shown between LA strain and the extent of fibrosis measured by LGE CMR.¹⁵⁷ Of note, for diagnostic and prognostic purposes, in other diseases such as HFpEF and MR, LA enlargement and LASr should be interpreted with caution in the presence of AF.

Several studies showed that larger amounts of epicardial fat were associated with an increased risk of AF.¹⁵⁸ Moreover, Wong et al.¹⁵⁹

showed a more robust association of AF with epicardial fat as compared to abdominal or overall adiposity.

Thromboembolism

Visualization of thrombus

Thrombi can be found free-floating or attached to the LA wall, in-transit across the patent foramen ovale (PFO), on prosthetic materials (valves, occluders), or in the LAA. LAA is the most common location of thrombi (98%) in non-valvular AF because of stasis in the blinded pouch.^{160,161}

TOE is the most frequently used tool to diagnose thrombus (sensitivity 93–100% and specificity 99%) in the LAA.¹⁶² The use of ultrasound contrast agents improves the diagnostic accuracy of TOE for detecting thrombus (Figure 17).¹⁶³ False-positive results are frequently due to the misdiagnosis of pectinate muscles as thrombi. Differential diagnosis also includes masses with a potential for embolization such as fibroelastomas, tumours, or vegetations. Contrast-enhanced CMR is the method of choice to make the differential diagnosis of intracardiac masses.¹⁶⁴

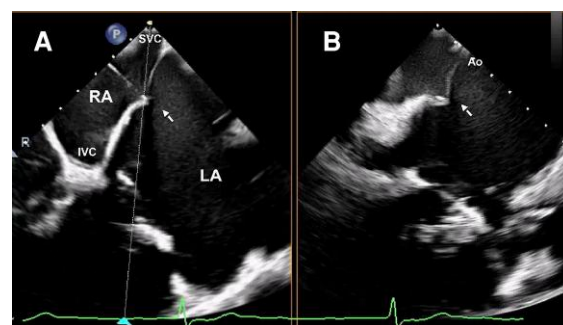


Figure 24 Trans-septal access with 3D ICE catheter using 2D X-plane imaging. A) the inter-atrial septum (IAS) is imaged in the superior and inferior orientation with the ICE catheter tip retroflexed. B) X-plane showing the anterior (Ao) and posterior. SVC- Superior vena cava; IVC = Inferior Vena Cava.

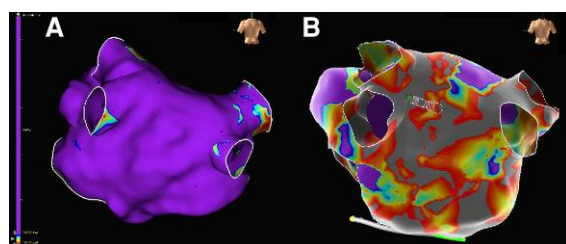


Figure 22 Electro-anatomical colour-coded voltage map (violet voltage >0.5 mV). (A) Normal voltage of the posterior wall. (B) Low-voltage areas and scar (grey colour) in native atrial cardiopathy.

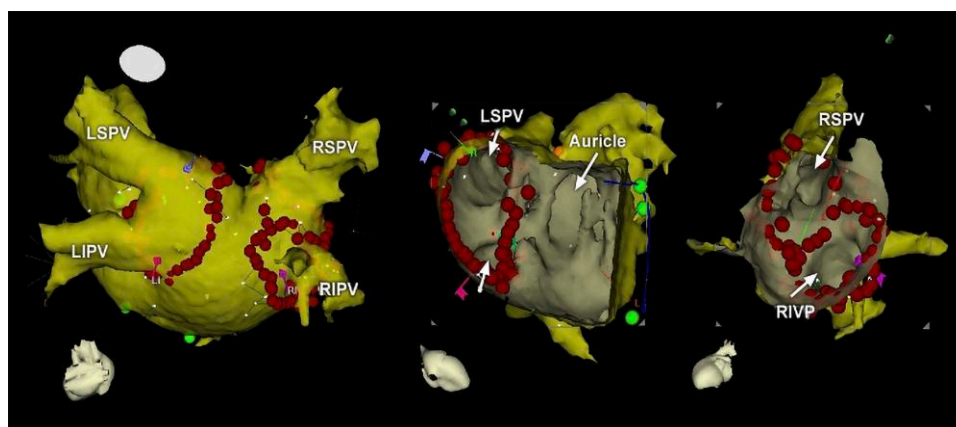
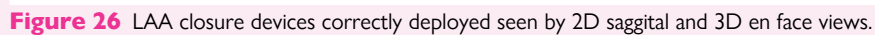
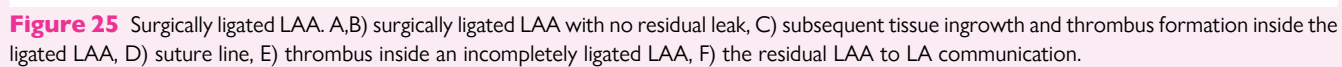


Figure 23 Integration of pre-acquired CCT reconstruction with electro-anatomical map used to guide PV isolation. The circle tags indicate radiofrequency energy delivery sites at the PV ostia. LSPV, left superior PV; LIPV, left inferior PV; RSPV, right superior PV; RIPV, right inferior PV.



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