Left atrial volume changes during exercise stress echocardiography in heart failure and hypertrophic cardiomyopathy

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PII: S1109-9666(22)00018-5

DOI: https://doi.org/10.1016/j.hjc.2022.01.003

Reference: HJC 670

To appear in: Hellenic Journal of Cardiology

Received Date: 15 August 2021
Revised Date: 25 January 2022
Accepted Date: 26 January 2022

Please cite this article as: Wierzbowska-Drabik K, Kasprzak JD, Haberka M, Peteiro J, Re F, D'Alfonso MG, Mori F, Palinkas ED, Agoston G, Varga A, Djordjevic-Dikic A, Tesic M, Zagatina A, Rodriguez-Zanella H, Simova I, Merli E, Morrone D, D'Andrea A, Camarozano AC, Reisenhofer B, Prota C, Citro R, Celutkiene J, Boshchenko A, Ciampi Q, Picano E, Left atrial volume changes during exercise stress echocardiography in heart failure and hypertrophic cardiomyopathy, *Hellenic Journal of Cardiology*, https://doi.org/10.1016/j.hjc.2022.01.003.

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Exercise echocardiography and left atrial changes

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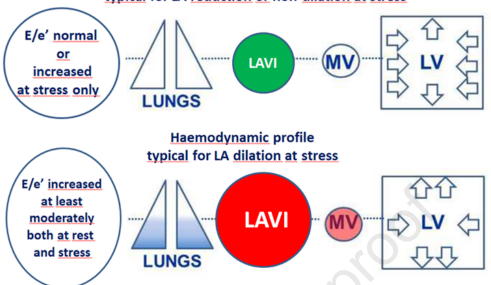
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Relationships with industry and Other Entities: none (for any author)

Wordcount of the whole manuscript (with abstract): 5 226 words.

<u>Haemodynamic</u> profile <u>typical</u> for LA <u>reduction or non-dilation at stress</u>



Main results in short:

- 1. Prevalence of reduction, stabilisation and dilation profiles of LAVI during stress test: 29%, 41% and 30%
 - 2.Dilation more frequent (near 3-fold) in HFrEF and HCM than in HFpEF
 - 3. LAVI change distribution the same with bike and treadmill exercise test
- 4. Resting predictors of stress LAVI dilation: E/e' > 14, LVEF < 50% and LAVI < 35 ml/m²

Structured Abstract

Background

We assessed feasibility and functional correlates of LAVI (left atrial volume index) changes during exercise stress echocardiography (ESE).

Methods

ESE on bike or treadmill was performed in 363 patients with heart failure with preserved ejection fraction (HFpEF, n=173), reduced ejection fraction (HFrEF, n=59) or hypertrophic cardiomyopathy (HCM, n=131). LAVI stress-rest increase ≥ 6.8 ml/m² was defined as dilation.

Results

LAVI measurements were feasible in 100%. LAVI did not change in HFrEF being at rest 32 (25-45) vs. at stress 36 (24 – 54) ml/m², P = NS and in HCM at rest 35 (26 – 48) vs. at stress 38 (28 – 48) ml/m², P = NS whereas it decreased in HFpEF from 30 (24 -40) to 29 (21 – 37) ml/m² at stress, P = 0.007. LA dilation occurred in 107 (30%) patients (27% with treadmill vs. 33% with bike ESE, P = NS): 26 with HFpEF (15%), 26 with HFrEF (44%) and 55 with HCM (42%) with P < 0.001 for HFrEF and HCM vs. HFpEF.

Multivariate analysis revealed as the predictors for LAVI dilation $E/e^{\circ} > 14$ at rest with OR 4.4, LVEF < 50% with OR 2.9, and LAVI at rest < 35 ml/m² with OR 2.7.

Conclusion

LAVI assessment during ESE was highly feasible and dilation equally frequent with treadmill or bike. LA dilation was threefold more frequent in HCM and HFrEF and could be predicted by increased resting E/e' and impaired EF as well as smaller baseline LAVI.

Key words: E/e' ratio, exercise stress echocardiography, left atrial volume index, left ventricular ejection fraction, mitral regurgitation.

Abbreviations List

ESE - exercise stress echocardiography

HCM - hypertrophic cardiomyopathy

HFpEF - heart failure with preserved ejection fraction

HFrEF - heart failure with reduced ejection fraction

LA- left atrium

LAP- left atrial pressure

LAVI - left atrial volume index

LVCR- left ventricular contractile reserve

MR- mitral regurgitation

SPAP - systolic pulmonary artery pressure

INTRODUCTION

Left atrial volume index (LAVI) is a well-recognized parameter, used not only for left atrium (LA) quantification but strongly recommended over LA diameter for left ventricular diastolic function stratification and used as a potent prognostic marker in a wide range of clinical conditions. ^{1,2,3,4} Its acute changes during stress echocardiography have been so far described in sparse studies with more focus given on long-term remodeling. ^{5,6,7} LAVI in the absence of significant mitral valve stenosis or regurgitation, atrial fibrillation and frequent arrhythmia, especially related to atrio-ventricular dissociation, reflects in the proportional manner the chronically increased filling pressure of the left ventricle, the advancement of diastolic impairment and the myocardial hypertrophy acting as a "barometer of the heart".

In the previous study from our stress echo 2020 network, Morrone et al. showed ⁸ that a subset of patients with suspected or diagnosed coronary artery disease undergoing pharmacological or exercise stress echocardiography (ESE) presents LAVI dilation. The pattern of stress-induced LAVI dilation correlated with abnormal left ventricle contractile reserve (LVCR) as well as with the increase of B-lines, a direct sign of pulmonary congestion correlated with an increase in pulmonary capillary wedge pressure.

However, LAVI plays also an important role outside chronic coronary syndromes and ESE is the recommended test modality for applications beyond coronary artery disease.

Our aim was to assess the feasibility and functional correlations of LAVI changes during ESE performed with treadmill or semi-supine bike in patients with heart failure with reduced ejection fraction (HFrEF), heart failure with preserved ejection fraction (HFpEF) and hypertrophic cardiomyopathy (HCM).

METHODS

The study was approved by the Ethical Committee as a part of the SE 2020 study (148-Comitato Etico Lazio-1, July 16,2016; Clinical trials.Gov Identifier NCT 030.49995). All patients gave their informed consent to enter the study.

Study group

We included 363 consecutive subjects undergoing clinically-indicated ESE in 17 centers of 10 countries from Europa, Russian Federation and America participating in SE 2020 study described in earlier papers. 9,10

Three subgroups were analysed: HFpEF, n=173, HFrEF, n=59, and HCM, n=131. Detailed inclusion/exclusion criteria were formulated in SE 2020 protocol in respective subprojects of the SE 2020 study. For the present study, the following inclusion/exclusion criteria were accepted:

For all three groups:

- patients aged > 18 years
- good visualization for at least 14 LV segments as the condition for qualification to stress echocardiography
- both sinus rhythm and atrial fibrillation at the baseline assessment were allowed

As exclusion criteria served for all groups:

- presence of prognosis-limiting comorbidities, such as advanced cancer, reducing life expectancy to < 1 year
- pregnancy/lactation
- unwillingness to give informed consent and to enter a regular follow-up program.

Specifically, to HFrEF group were included:

- patients with reported diminished exercise tolerance and proved or suspected heart failure with symptoms ranging from I to III NYHA class
- LV ejection fraction < 50% as assessed at baseline echocardiography (etiology of CAD and DCM of heart failure were allowed, significant mitral and aortic valve stenosis and severe primary valve regurgitation excluded)

To HFpEF group:

- patients with known and suspected heart failure from I to III NYHA class
- LV ejection fraction ≥ 50% at baseline echocardiography, significant mitral and aortic stenosis as well as severe primary valve regurgitation (organic) excluded
- patients with no alternative causes of dyspnea such as anemia and/or chronic obstructive lung disease

To HCM group:

- patients with LV myocardial thickness ≥ 15 mm in any segment in the absence of another cause of LV hypertrophy and resting (or after Valsalva maneuver) left ventricular outflow tract gradient < 50 mmHg
- specific exclusion criteria for this group included phenocopies of HCM such as infiltrative/storage disease (eg, Fabry disease), ejection fraction <45% and history of coronary artery disease.

Echocardiographic rest and stress assessment

Transthoracic echocardiography at rest and ESE was performed with high-end echocardiographic systems. An electrocardiogram (ECG) tracing was displayed on the monitor during the examination. The echocardiographic measurements were acquired following the recommendations. All echocardiographers in Stress Echo 2020 had passed quality control of reading examinations with interobserver reproducibility exceeding 90%. ESE modality was semi-supine bike in 178 and treadmill in 185 patients. Criteria for interrupting the test were chest pain, induced wall motion abnormalities, significant rhythm disturbances, excessive blood pressure increase or hypotonia, limiting dyspnoea and fatigue, legs pain or predicted heart rate.

The definition of LAVI dilation was based on a well validated statistic called reference change value, taking into account biological, analytical and observer variability with LAVI change of ≥ 6.8 ml/m² between rest and stress considered as a change above background variation and used as a cut-off to identify a LAVI – dilator cohort. ^{14,5,15}

Statistical analysis

The distribution of variables was assessed with the D'Agostino-Pearson test and adequate parametric or non-parametric tests were used. Accordingly to distribution data was expressed as mean \pm standard deviation or median and interquartile range or frequency for categorical data. Multiple-samples comparison was performed with analysis of variance and Newmann-Keuls test or Kruskal-Wallis test respectively. Frequency of categorical data was compared with ch² test. One-sample comparisons were performed with paired t-test or Wilcoxon test. For correlation Pearson's or Spearman coefficients were calculated. Statistical significance was set at P < 0.05. Multivariate stepwise logistic regression analysis was conducted on the variables significant in univariate analysis with P value set at < 0.05 for entering and > 0.1 for removing variable. Analyses were conducted with MedCalc V. 12.1.4. (Frank Schoonjans, Belgium).

RESULTS

Rest and stress characteristic of patients

Main characteristics of the study patients at rest and stress are summarized in Table 1 and Table 2. LAVI reducers prevalence was around 30% and similar in all groups, whereas LAVI dilators were 3-times more frequent in HFrEF (44%) and HCM (42%) as compared to HFpEF patients (15%), Table 2.

As far as the comparison between type of ESE is concerned in group examined with ergometer LAVI profiles prevalence was as follow: 26% reducers, 41% with stable LAVI and 33% dilators, whereas in group examined with treadmill there were 32% reducers, 41% with stable parameter and 27% dilators and this proportions did not differ statistically.

Predictors of dilated LAVI during ESE

We analyzed patients with reduced (LAVI diminished during stress by $> 6.8 \text{ ml/m}^2$), stable (stress-rest changes \pm 6.8 ml/m²) and dilated LAVI (rest-stress increase $> 6.8 \text{ ml/m}^2$). LAVI-reducers displayed higher resting LAVI and less frequent advanced mitral regurgitation (Table 3). At peak ESE dilators showed more frequent MR, lowest LVCR and higher E/e', Table 4.

Abnormal (>34 ml/m²) LAVI values measured at the peak of ESE better then resting LAVI separated patients with abnormal LVCR, increased E/e², greater then mild mitral regurgitation and similarly with increased SPAP (Figure 1).

At individual patient analysis based on rest LAVI values 209 patients showed normal (<35 ml/m²), 103 moderately abnormal (35-50 ml/m²) and 51 severely abnormal (>50 ml/m²) LAVI. During stress 170 patients (47%) were reclassified with 88 (24%) increasing their LAVI of at least one grade (Figure 2). Figure 3 displays an example of patient with small reduction of LAVI during ESE.

Predictors of LAVI changes during ESE

Predictors of LAVI dilation detected in univariate and multivariate logistic regression analysis are shown in Table 5. Multivariate stepwise analysis revealed as the predictors for LAVI exercise-related dilation 3 parameters: E/E' > 14 at rest with OR 4.4, 95% CI (1.9-9.9), rest LVEF < 50% with OR 2.9, 95% CI (1.2 – 6.9) and LAVI at rest < 35 ml/m² with OR 2.7, 95% CI (1.2 - 6.2).

Finally, we analyzed exercise-related LAVI changes while going from normal filling pressure with E/e' < 12 at both rest and stress trough mild stage of diastolic dysfunction with E/e' normal at rest but elevated at stress, towards E/e' elevation \geq 12 at both rest and stress and the most advanced stage with E/e' > 15 at rest and stress. We found that "LAVI volumetric behavior during ESE" reflected consistently LAP pressure burden (revealed as

noninvasive proxy E/e' ratio) showing transition from reduction through stabilization towards dilation pattern, see Table 6 and Figure 4.

DISCUSSION

LAVI is a highly dynamic variable and may change substantially during ESE in HFrEF, HFpEF and HCM. In all these conditions, LAVI assessment is feasible with high success rate showing a heterogeneous response at individual patient analysis. LAVI-dilator pattern is threefold more frequent in HFrEF and HCM compared to HFpEF patients. Some functional variables are associated with LAVI-dilator pattern. These variables are the increase in E/e' (a widely-accepted proxy of increased left ventricular filling pressure), the reduction of LVEF and more frequent and advanced mitral regurgitation. All these variables potentially concur in determining of the increased left atrial pressure through abnormalities of LV diastolic, systolic or mitral valve function. Although to some extent the LAVI dilatation during exercise may act as physiological increasing of atrial reservoir function at a later stage a decompensation occurs with signs of backward failure with possible B-lines appearance, as observed in HFpEF subgroup.

Dilators showed the highest value of E/e' at rest as well as the higher percentage of moderate and severe MR as compared to reducers. Moreover, at peak stage of ESE, higher E/e' with more frequent MR (especially advanced grades) as well as lower LVCR and LVEF were observed consistently in dilators group, see Figure 5 (central figure).

One of the most important finding is the relationship between E/e' values (at rest and peak ESE) and LAVI at rest and peak as well as its dynamics. Achieved results are displayed in Figure 4 and reveal reduction of LAVI in subjects with normal E/e' (and the lowest LAP pressure) stable LAVI behaviour at early stages of diastolic dysfunction, followed by small and large dilation of median LA (by circa 10 ml/m²) in patients with moderately and severely elevated left atrial pressure. On the other hand, the pathologic dilation may be limited in most advanced stage by the increasing stiffness of severely remodelled LA. This seems to be supported by the relationship between basal LAVI and LAVI values at peak exercise in our group. According

to Figure 2 the typical reaction for exercise was in the smallest LA group the stabilisation or increase of volume whereas in the patients with the most enlarged atria more often the reduction of LAVI.

Finally, in multivariable analysis to the predictors of significant increase of LAVI during ESE belonged E/E' ratio >14 and LVEF lowered < 50%, as well as LAVI < 35 ml/m² at rest offering still greater potential or reserve for dilation, whereas atria with LAVI > 50 ml/m² at rest tended to diminished at peak stage.

Comparison with previous studies

Our findings confirm and expand previous, limited experiences showing the high success rate of LAVI imaging during ESE already documented in chronic coronary syndromes, hypertrophic cardiomyopathy, valvular heart disease and heart failure. ¹⁶

Atrial volume and function during exercise were examined by Schnell F et al. in group of 45 subjects including normal individuals, endurance athletes and patients with chronic thromboembolic pulmonary hypertension (CTEPH).¹⁷ The authors observed larger LAVI at rest in athletes (56 ml/m²) then in controls (40 ml/m²) and CTEPH (31 ml/m²) which decreased consistently during exercise. Contrary, right atrial volume index (RAVI) increased during stress in patients with CTEPH (from 64 to 79 ml/m²) while decreasing in controls and athletes. The authors hypothesised that increase of reservoir function of both atria provides adequate filling for ventricles and maintenance of cardiac output. In their group reservoir function of left atrium was enhanced mainly by diminishing of minimal LAVI (end-diastolic volume) without significant increasing of maximal LAVI (reflecting maximal LA volume during ventricular end systole), nevertheless in more pathological states this may require also the increase of maximal LAVI, what probably took place in our small and large dilators. Pathologic character of significant atrial dilation is suggested by Schnell observation concerning RAVI enlargement in CTEPH patients.

Valuable observation of mechanism of exercise-related lung congestion in patients with HFpEF has been recently published by Reddy et al. ¹⁸ The authors noticed that patients with HFpEF who developed lung congestion during exercise test, confirmed with B-lines appearance, were

characterised not only by increased pulmonary capillary wedge pressure but also by increased pulmonary artery and right atrial pressure as well as by indices of deranged right ventricle - pulmonary artery coupling i.e. lower ratios of TAPSE, FAC (fractional area change) and RV S' to mean pulmonary artery pressure. Nevertheless, in our group only a small subgroup developed pulmonary congestion during ESE and we did not observe significant and consistent impairment of pulmonary pressure and RV function in the LAVI dilators.

We pooled data from semi-supine and treadmill ESE which have different hemodynamic effect and could have influence on volume changes and stress LAV changes to some extent. According to the literature bicycle increases the blood pressure significantly but not so much the heart rate comparing to the treadmill. The final double product is however similar. Semi-supine exercise increases pulmonary capillary wedge pressure more than upright exercise, and treadmill increases end-diastolic volume of left ventricle more than semi-supine exercise in healthy subjects. 20,21 The observational study design did not interfere with the individual choice of the referring physician, which is a matter of personal experience, awareness of the individual patient indications and local practice. However, the prevalence of LAV dilation, reduction and stabilization was similar with the both tests used in our population, suggesting their comparable impact on left atrium.

Study limitations

Each group had a moderate sample size limiting the statistical strength of subgroup analyses. However, the findings were consistent suggesting that observed functional correlations act similarly across various cardiovascular conditions.

Concentration on volumetric data limited to LA in its reservoir phase neglected more comprehensive analysis of LA function, which however is especially challenging in time-limited settings of ESE.

We used different exercise modalities, such as treadmill and supine bike with some difference in type of hemodynamic stress induced, which are however both widely accepted for ESE beyond coronary artery diseases.

Despite observed in our study similar prevalence of reduced, stable and dilated LAVI pattern in both bicycle and treadmill tests, more detailed, quantitative comparisons of LAVI reaction to various types of exercise loading in various groups of patients, as well as the dependence from the achieved workload is needed in the future.

Finally, the analysis of clinical outcomes should be advocated in an longitudinal study to understand better the prognostic significance of LAVI changes during ESE.

Conclusions:

LAVI is a relatively simple parameter which can be obtained with high success rate in various cardiovascular conditions from HFrEF to HFpEF and HCM and measured at peak stress reflects the functional status more closely than rest values.

The LAVI dilator pattern is associated with more frequent and severe mitral regurgitation, diastolic dysfunction and pulmonary congestion all known as adverse predictors of outcome. With only minimal increase on analysis time LAVI can be a further useful adjunct to comprehensive SE also beyond coronary artery disease.

Competency in medical knowledge

The echocardiographic assessment of LAVI exercise changes and their classification to proposed patterns of reducers, stable LAVI and dilators provides a novel and integral tool reflecting status and function of both circulations. High feasibility, simplicity, time

efficiency and non-invasiveness of LAVI assessment supports its wider use in the settings of various ESE and clinical entities.

Translational outlook

Although widely appreciated in resting assessment of left ventricular function LAVI still requires further detailed diagnostic and prognostic evaluation as far as its dynamic changes are concerned. Presented study may be seen as the preliminary confirmation of high feasibility in both kinds of ESE, bike and treadmill based, wide diagnostic potential of LAVI changes assessment in patients with different types of heart failure as well as the indicator of the underestimation of this patient and operator friendly parameter in present clinical practice.

Tables and Figures:

Figure legend:

Figure 1. Functional correlates of LAVI at rest and peak ESE. The comparison of global systolic left ventricular function parameter LVCR (left upper panel), frequency of mitral regurgitation (left lower panel), E/e' (right upper panel) and SPAP values (right lower panel) between groups with normal/mildly enlarged (white bars) and significantly enlarged LAVI (red bars) assessed at rest (left columns in each panel) and at peak stress (right columns). The patients with larger LAVI at stress displayed lower LVCR, higher E/e', SPAP and MR frequency. Stress LAVI was more closely related to impaired hemodynamics.

E/e' – the ratio of maximal velocity of early wave of mitral inflow (E) to maximal velocity of mitral annulus motion early phase (e'), LAVI- left atrial volume index, LVCR- left ventricular contractility reserve, MR- mitral regurgitation, SPAP- systolic pulmonary artery pressure

Central panel displays the numbers of patients transferred between categories of normal or mildly (<35 ml/m²), moderately (35-50 ml/m²) and severely enlarged (>50 ml/m²) LAVI.

During stress 170 patients (47%) were reclassified with 88 (24%) increasing their LAVI of at least one grade. 138 patients from normal resting LAVI group (66%) did not dilate significantly during ESE.

In the group comparison median of normal LAVI increased significantly from 25.8 to 28.9 ml/m², P < 0.001, whereas the largest LAVI decreased from 60.5 to 38.4 ml/m², P < 0.0001, and intermediate from 40.2 to 36.4 ml/m², P = 0.002.

Figure 3. The examples of LAVI measured in 4 chamber and 2 chamber views in patient showing LAVI reduction during ESE.

Patient with BSA 2.2 m^2 and slight reduction of LAVI (Δ 5 ml/m², which is < 6.8 ml/m² cut-off value) during ESE: from 27 ml/m² to 22 ml/m² as calculated for biplane LAVI (given in the Figure) .

Figure 4. Evolution of LAVI ESE - induced changes in relations to E/e' values at rest and at stress. LAVI seems to compensate the short intervals of elevated E/e' related to exercise only, but dilates significantly in more advanced stages, while E/e' is elevated also at rest.

E/e' – the ratio of maximal velocity of early wave of mitral inflow (E) to maximal velocity of mitral annulus motion early phase (E'), LAVI- left atrial volume index,

Figure 5. (Graphical abstract) Graphic presentation of pathophysiologic changes of LV function and circulation's hemodynamics as typical for LAVI reductors or non-dilators (upper panel) and dilators (lower panel).

The upper panel displays normal function of left ventricle (good contractility represented by preserved LVEF or LVCR expressed as triple inward arrows inside the box) as well as not-elevated LV pressure (or its proxy E/e', illustrated with outward single arrow), normal or elevated only at stress E/e' ratio (regular circle) and absent B-lines in lungs and MR (clear lungs and mitral valve icons) in the presence of small LA at stress (green LAVI). The opposite situation related to LAVI dilator pattern (red, increased LAVI) is rendered in the lower panel.

E/e'- ratio of early mitral inflow velocity to early mitral annulus motion velocity, LAVI- left atrial volume index, LUNGS- with eicon represents absence of B-lines, partially blue-filled represents water congestion with present B-lines, LV- left ventricle, arrows directed inwardly represent LVCR (left ventricular contractile reserve) or LVEF (left ventricular ejection fraction) - MV- mitral valve, white circle represent absend or mild MR, red one — the presence of MR above second grade, MR- mitral regurgitation.

Table 1. Demographic and clinical characteristic as well as resting echo data

Variable	Group 1	Group 2	Group 3	P value	P value	P value
	HFrEF,	HFpEF,	нсм,	HFrEF	HFpEF	HFrEF
	N = 59	N = 173	N = 131	vs	vs HCM	vs HCM
				HFpEF		
Age (years)	61 (51 – 71)	69 (63-75)	52 (42-61)	< 0.05	<0.05	<0.05
Sex,	42/17	63/110	84/47	<0.001	<0.001	ns
M/F, (M %)	(71.2%)	(36.4%)	(64.1%)	Ċ		
BSA (m ²)	1.91	1.93	1.89	ns	<0.05	ns
	(1.8 - 2.1)	(1.8 - 2.1)	(1.7 - 2.0)			
BMI (kg/m ²)	26.5	29.4	25.7	< 0.05	<0.05	ns
	(24.2 - 30.1)	(26.1 – 32.6)	(23.3 - 28.1)			
NYHA class 1	10 (17.9%)	23 (13.5%)	81 (61.8%)	=0.03	<0.001	< 0.001
2	37 (66.1%)	138 (80.7%)	44 (33.6%)			
3	9 (16.1%)	10 (5.8%)	6 (4.6%)			
НА	35 (59.3%)	153 (88.4%)	43 (32.8%)	< 0.001	<0.001	=0.001
Diabetes	15 (25.4%)	55 (31.8%)	3 (2.3%)	ns	<0.001	=0.001
Dyslipidemia	35 (59.3%)	92 (53.2%)	32 (24.4%)	ns	<0.001	<0.001
Smoking	22 (40%)	23 (13.3%)	31 (23.8%)	< 0.001	=0.03	ns
HR rest (bpm)	69 (60 – 81)	72 (64 – 80)	65 (60 – 73)	ns	<0.05	<0.05
DBP rest	75 ± 10	77 ± 10	76 ± 9	ns	ns	ns
(mmHg)						
SBP rest	115 ± 19	129 ± 18	120 ± 16	< 0.05	<0.05	<0.05
(mmHg)						
EF rest (%)	38 (32- 46)	62 (55- 67)	67 (62-72)	<0.05	<0.05	<0.05

WMSI rest	2.0 (1.4-2.3)	1.0 (1.0-1.0)	1.0 (1.0-1.0)	< 0.05	< 0.05	< 0.05
E/e' rest	12.9	10.2	9.9	< 0.05	ns	<0.05
	(9.7 - 16.3)	(8.3 - 12.6)	(7.6 - 12.9)			
SPAP rest	29 (21-39)	27 (23-36)	28 (25-33)	ns	ns	ns
TAPSE rest	20 (16-24)	24 (21-28)	24 (21-26)	< 0.05	ns	<0.05
B lines rest (n)	0 (0-4)	1 (0-3)	0 (0-0)	ns	< 0.05	< 0.05
LAVI rest	32 (25 - 45)	30 (24 – 40)	35 (26 – 48)	ns	<0.05	ns
(ml/m ²)						
MR rest 0	17 (28.8%)	115 (66.9%)	61 (47.3%)	<0.001	=0.001	=0.006
1	25 (42.2%)	49 (28.5%)	51 (39.5%)			
2	12 (20.3%)	6 (3.5%)	16 (12.4%)	>		
3	5 (8.5%)	2 (1.2%)	1 (0.8%)			

HR- heart rate, DPB- diastolic blood pressure, SBP- systolic blood pressure, EF- ejection fraction, WMSI- wall motion score index, LAD- left anterior descending coronary artery, LAVI- left atrial volume index

Table 2. Hemodynamic and echo data at peak

Variable	Group 1	Group 2	Group 3	P value	P value	P value
	HFrEF,	HFpEF,	нсм,	HFrEF	HFpEF	HFrEF
	N = 59	N = 173	N = 131	vs	vs HCM	vs
				HFpEF		нсм
HR peak (bpm)	120 ± 26	124 ±23	132±26	ns	<0.05	<0.05
SBP peak (mmHg)	144 ± 34	165 ± 27	164 ± 28	< 0.05	ns	<0.05
DBP peak (mmHg)	82 ± 12	83 ± 16	85 ± 12	ns	ns	ns
EF peak (%)	37 (29-48)	70 (63-76)	73 (67-77)	< 0.05	< 0.05	<0.05
WMSI peak	2.0 (1.15-	1.0 (1.0-1.0)	1.0 (1.0-1.0)	< 0.05	ns	<0.05
	2.3)		0			
E/e' peak	14	12	10	ns	<0.05	<0.05
	(10.2 - 20)	(9.7 - 15)	(7.6 - 13.4)			
SPAP peak	41 (25-59)	38 (30 - 55)	40 (31-50)	ns	ns	ns
TAPSE peak	26 (21-30)	27 (24-31)	30 (25-34)	ns	ns	ns
B lines peak	1 (0 - 10)	3 (1 - 5)	0 (0 - 1)	< 0.05	<0.05	<0.05
LVCR	1.4	1.7	1.6	< 0.05	ns	ns
	(1.1 - 1.8)	(1.3 - 2.2)	(1.3 - 2.1)			
LAVI peak (ml/m ²)	36 (24 – 54)	29 (21 – 37)	38 (28 – 48)	< 0.05	< 0.05	ns
ΔLAVI	4.8 (-6.5 –	-2.3 (-7.7 –	2.2 (-15.3 –	< 0.05	<0.05	ns
	15)	3.4)	18.9)			
MR peak 0	14 (28.6%)	124 (77%)	51 (40.5%)	< 0.001	<0.001	ns
1	16 (32.7%)	30 (18.6%)	50 (39.7%)			
2	12 (24.5%)	4 (2.5%)	19 (15.1%)			
3	6 (12,2%)	3 (1.9%)	4 (3.2%)			

4	1 (2%)	0 (0%)	2 (1.6%)			
Reducers LAVI	15 (25.4%)	48 (27.7%)	43 (32.8%)	< 0.001	< 0.001	ns
Stable LAVI	18 (30.5%)	99 (57.2%)	33 (25.2%)			
Dilators LAVI	26 (44.1%)	26 (15%)	55 (42%)			
Positive SE (visual)	16 (27.1%)	7 (7.5%)	5 (3.8%)	0.002	ns	< 0.001
, , ,						

HR- heart rate, DPB- diastolic blood pressure, SBP- systolic blood pressure, EF- ejection fraction,

WMSI- wall motion score index, LVCR- left ventricular contractile reserve (Force at peak/Force rest ratio; Force = SBP/LVESV, LVESV

Table 3. Resting characteristic of groups with reduced, stable and dilated LAVI.

Variable	Group 1	Group 2	Group 3	P value	P value	P value
	Reduced LAVI		Dilated LAVI	reduced	stable	reduced
	N = 106	LAVI,	N = 107	vs	vs	vs
		N = 150		stable	dilated	dilated
Age	62 (44 – 71)	66 (55 – 71)	61 (49 – 70)	ns	ns	ns
Sex M (%)	54 (50.9%)	70 (46.7%)	65 (60.7%)	ns	=0.04	ns
BMI	26.9	28.1	26.8	ns	ns	ns
	(24.2 - 30.9)	(25.1 – 30.8)	(24.4 - 30.4)			
BSA	1.9 (1.72 – 2.05)	1.9	1.92	ns	ns	ns
		(1.78 - 2.06)	(1.76 - 2.08)			
NYHA class				ns	ns	ns
1	38 (36.5%)	39 (26.5%)	37 (34.6%)			
2	61 (58.7%)	100 (68%)	58 (54.2%)			
3	5 (4.8%)	8 (5.4%)	12 (11.2%)			
Hypertension	58 (54.7%)	113 (75.3%)	60 (56.1%)	=0.0009	=0.002	ns
Diabetes	19 (17.9%)	36 (24.0%)	18 (16.8%)	ns	ns	ns
Dyslipidemia	42 (39.6%)	72 (48%)	45 (42.1%)	ns	ns	ns
Smoking	14 (13.2%)	32 (21.6%)	30 (28.8%)	ns	ns	=0.009

HR rest (bpm)	71 (63 – 80)	70 (62 – 77)	69 (60 – 78)	ns	ns	ns
SBP rest	124 ± 18	127 ± 19	119 ± 18	ns	< 0.05	ns
(mmHg)						
DBP rest	80 (70-80)	80 (70-80)	74 (70-80)	ns	ns	ns
(mmHg)						
EF rest (%)	64 (57 – 69)	62 (54- 68)	60 (45 – 69)	ns	ns	ns
WMSI rest	1.0 (1.0-1.0)	1.0 (1.0-1.0)	1.0 (1.0-1.39)	ns	ns	ns
E/e' rest	10 (8- 13.2)	10 (8 – 12)	11.3 (8 – 15.4)	ns	< 0.05	<0.05
SPAP rest	26 (23-30)	28 (22-36)	30 (25-35)	ns	ns	ns
TAPSE rest	24 (20 – 26)	23 (21 – 27)	23 (20 - 26)	ns	ns	ns
B lines rest (n)	0 (0-1.0)	0 (0-2.5)	0 (0-2.0)	ns	ns	ns
LAVI rest	44 (36 – 55)	29 (22- 38)	27 (23 – 34)	< 0.05	ns	< 0.05
(ml/m ²)						
MR rest				< 0.001	0.048	< 0.001
0	75 (70.8%)	69 (46.6%)	49 (46.2%)			
1	28 (26.4%)	61 (41.2%)	36 (34%)			
2	3 (2.8%)	17 (11.5%)	14 (13.2%)			
3	0 (0)	1 (0.7%)	7 (6.6%)			

 $Table \ 4. \ Peak \ stress \ characteristic \ of \ reduced, \ stable \ and \ dilated \ LAVI \ during \ ESE$

Variable	Group 1	Group 2	Group 3	P value	P value	P value
	Reduced LAVI	Stable LAVI,	Dilated LAVI	reduced	stable	reduced
	N = 106	N = 150	N = 107	vs	vs	vs
				stable	dilated	dilated
HR peak (bpm)	130 (116 – 150)	121 (107 – 137)	125 (106 – 140)	< 0.05	ns	<0.05
SBP peak (mmHg)	167 (150 – 180)	160 (145 – 180)	150 (135 – 171)	ns	< 0.05	<0.05
DBP peak (mmHg)	80 (80-90)	80 (79 - 94)	80 (70-90)	ns	ns	ns
EF peak	71 (62 - 77)	68 (60 - 75)	67 (52 - 75)	ns	ns	<0.05
WMSI peak	1.0 (1.0-1.0)	1.0 (1.0-1.0)	1.0 (1.0-1.13)	ns	ns	ns
E/e' peak	10.8 (8.1 – 14.2)	11.1 (8.5 - 14)	13 (9.9-16.6)	ns	< 0.05	<0.05
SPAP peak *	37 (25 - 51)	39 (30 - 53)	42 (35 - 55)	ns	ns	ns
TAPSE peak	29 (25 - 33)	27 (23 - 31)	27 (22 - 31)	ns	ns	ns
B lines peak	0.5 (0 - 2.0)	3.0 (0 - 5.8)	1.0 (0 - 4.0)	ns	ns	ns
LVCR	1.71 (1.4 – 2.47)	1.63 (1.3 – 2.09)	1.4 (1.06 – 1.9)	ns	<0.05	<0.05
LAVI peak	27 (20 - 34)	29 (23 - 37)	47 (37 - 60)	< 0.05	< 0.05	<0.05
(ml/m ²)						
ΔLAVI	-14.6 (-25.05 -	0.0 (-3.05 – 3.11)	16.6 (10.4 –	<0.05	< 0.05	<0.05
	-9.07)		29.7)			
MR peak				ns	ns	<0.001
0	70 (68.6%)	76 (58%)	43 (41.7%)			
1	25 (24.5%)	38 (29%)	33 (32%)			
2	6 (5.9%)	12 (9.2%)	17 (16.5%)			
3	1 (1.0%)	4 (3.1%)	8 (7.8%)			
4	0 (0%)	1 (0.8%)	2 (1.9%)			

[•] SPAP peak feasibile Reduced n=40, stable n=68, dilated n=58

Table 5. Univariate and multivariate analysis of predictors for ESE related LAVI dilation.

Univariate analysis in whole group								N			iate analy ble group	sis	
Parameter	C	R	95% C	EI	P		OR		95	% C	[P	
E/e rest >14	3	.0	1.8 – 5.1		<0.0	001	4.4		1.9	9.9)	<0	.001
E/e stress >1	4 2	.1	1.3 – 3	.5	0.00)3							
MR rest > 2nd grade	1	7.9	2.2 – 1	47.3	0.00)7				C			
MR stress > 2nd grade	4	.1	1.4 – 1	1.5	0.00)8		S			7		
LVEF < 50°	% 2	.0	1.2 - 3	.4	0.01		2.9		1.2	2 – 6.9)	0.01	
LVCR < 2	2	.1	1.2 – 3	.6	0.01	54							
B-lines rest	1	5.4	1.8 - 13	31	0.01	12							
B-lines stres	s 3	.8	1.3 – 1	1.1	0.01	16							
LAVI rest < 35 ml/m ²	< 3	.1	1.9 – 5	.2	<0.0	001	2.7		1.2	2 – 6.2	2	0.0)2
Mult	ivaria in H		nalysis F		Mul		riate a HFrE	-	S	M	ultivariat in H0		alysis
Parameter	OR	_	% CI	P	OR	95%	6 CI	P		OR	95% CI		P
E/e rest >14	na	na		na	na	na	na na		na 5.0		1.5 – 16.	.3	0.007
E/e stress >14	na	na na na		9.0	2.1-	37.5	0.003	0.003 na		na		na	
LAVI rest < 35 ml/m ²	5.2	1.5	-18.2	0.01	na	na		na		9.6	3.9 – 23.	.2	<0.001

Table 6. Natural history of LA dysfunction. Comparison between LAVI in group I: normal or mildly increased E/e' both at rest and stress (<12), group II with substantial increase of E/e' during stress only (<12 at rest and \geq 12 at peak EXE) and group III with E/e' elevated in rest and stress \geq 12. Additionally group IV with very high E/e' at rest and stress (>15 at both stages) was analyzed showing extreme dilation of left atrium during stress.

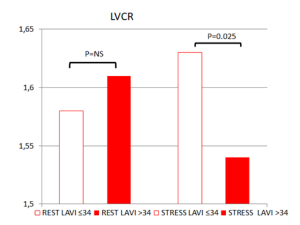
Variable	Group I	Group II	Group III	Group IV	P value					
	N = 157	N = 57	N = 98	N = 41	1 vs 2	2 vs 3	1 vs 3	1 vs 4	2 vs 4	3 vs 4
LAVI rest	31 (23 - 42)	31 (26 - 38)	39 (27- 48)	40 (31- 50)	ns	0.002	<0.001	0.003	0.002	ns
ml/m ²				V bil	9					
LAVI peak	29 (21- 39)	32 (24 - 39)	39 (32- 56)	51 (37 - 69)	ns	< 0.001	< 0.001	< 0.001	< 0.001	0.035
ml/m ²			\(
P value	0.034	ns	0.026	< 0.001	NA	NA	NA	NA	NA	NA
rest vs										
stress										
ESE LAVI	Reduction	Stabilization	Light dilatation	Large dilatation	NA	NA	NA	NA	NA	NA
change										

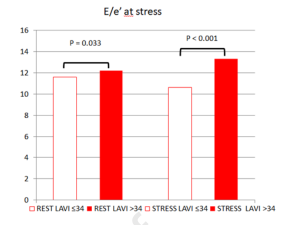
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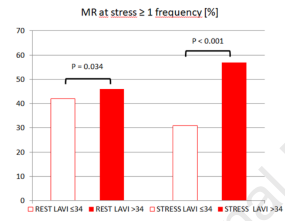
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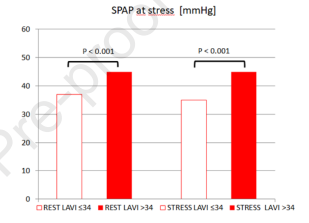
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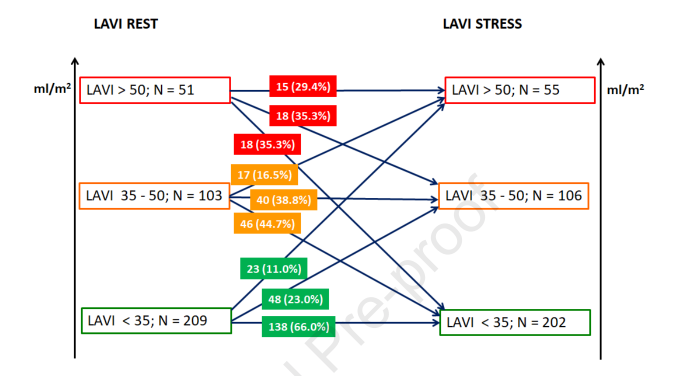
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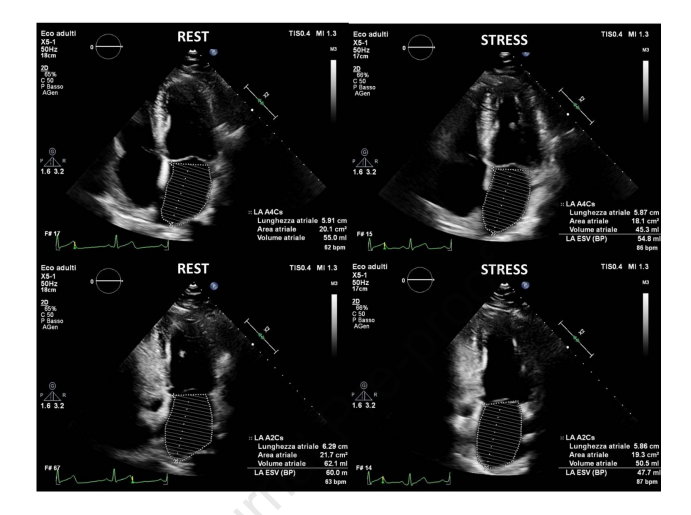


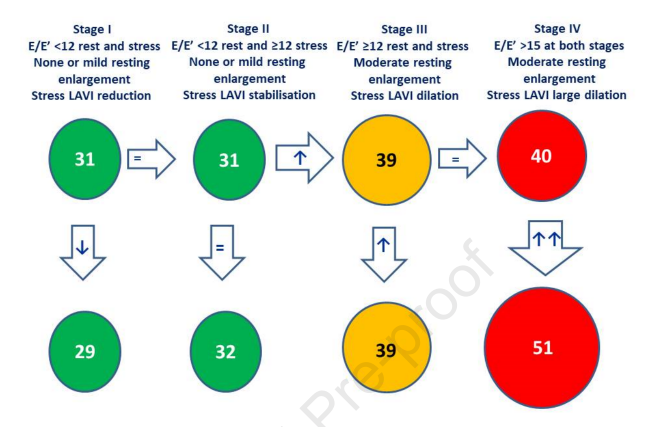












Numbers display median LAVI values, arrows significant changes of LAVI, equality sign lack of change between compared groups; Green = LAVI ≤ 35, Yellow = LAVI > 35 and < 40, Red = LAVI ≥ 40 ml/m²