#### **ORIGINAL PAPER**



# Scan-rescan reproducibility of diastolic left ventricular kinetic energy, viscous energy loss and vorticity assessment using 4D flow MRI: analysis in healthy subjects

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#### Abstract

The aim of the current study was to assess the scan–rescan reproducibility of left ventricular (LV) kinetic energy (KE), viscous energy loss (EL) and vorticity during diastole from four-dimensional flow magnetic resonance imaging (4D flow MRI) in healthy subjects. Twelve volunteers (age  $27 \pm 3$  years) underwent whole-heart 4D flow MRI twice in one session. In-scan consistency was evaluated by correlation between KE and EL.  $\text{EL}_{index}$  was computed to measure the amount of EL relative to KE over diastole. Scan–rescan analysis was performed to test reproducibility of volumetric measurements of KE, EL,  $\text{EL}_{index}$  and vorticity in the LV over early (E) and late (A) diastolic filling. In-scan consistency between KE and EL was strong-excellent (E-filling scan1: r=0.92, P<0.001; scan2:  $\rho=0.96$ , P<0.001 and A-filling scan1:  $\rho=0.87$ , P<0.001; scan2: r=0.99, P<0.001). For the majority of subjects (10 out of 12), KE and EL measures showed good to strong reproducibility. However, with a wide range of agreement [intraclass correlation (ICC): 0.64-0.95] and coefficients of variation (CV)  $\leq 25\%$ . EL<sub>index</sub> showed strong reproducibility for all 12 subjects with a strong ICC (0.94, P<0.001) and a CV of 9%. Scan–rescan reproducibility of volumetric vorticity showed good–excellent ICCs (0.83-0.95) with CVs  $\leq 11\%$ . In conclusion, the current study shows strong–excellent in-scan consistency and overall good agreement between scans for 4D flow MRI assessment of left ventricular kinetic energy, energy loss and vorticity over diastole. However, substantial differences between the scans were also found in some parameters in two out of twelve subjects. Strong reproducibility was found in the dimensionless EL<sub>index</sub>, which measures the amount of viscous energy loss relative to the average kinetic energy over diastole.

Keywords 4D flow MRI · Magnetic resonance imaging · Kinetic energy · Viscous energy loss · Vorticity

# Introduction

Congenital and acquired heart diseases affect the efficacy of intracardiac flow patterns and energy distribution [1, 2]. Given the three-dimensional (3D) time-varying nature of these flow patterns and energetics, specialized in vivo

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imaging and analysis techniques are needed to evaluate volumetric changes in such complex hemodynamic parameters. Four-dimensional flow magnetic resonance imaging (4D flow MRI) allows for comprehensive non-invasive assessment of 3D time-varying blood flow properties in the heart and great vessels in all three velocity encoding directions and spatial dimensions over the cardiac cycle [3].

4D flow MRI has recently emerged as a novel tool for in vivo quantification of intracardiac flow energetics, associated energy losses and vortical flow patterns by means of kinetic energy (KE) [4–7], viscous energy loss (EL) [1] and vorticity [8, 9]. KE is the energy contained in the flow of the bloodstream due to motion and EL is the KE that is irreversibly lost due to viscosity-induced frictional forces within the blood flow [1]. In acquired heart disease, remodelling occurs which can lead to alterations in intracardiac hemodynamics [4]. Alteration in

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intraventricular KE derived from 4D flow MRI has been used to assess left ventricular (LV) and right ventricular (RV) (dys)function in patients with different stages of heart failure (HF) [4-7]. This is also the case in various congenital heart diseases. Even after correction, patients may develop systolic and/or diastolic dysfunction, leading to changes in intracardiac flow energetics [1]. In addition, intracardiac anatomy may not be restored. In corrected atrioventricular septal defect patients, elevated EL was associated with altered 3D vortex ring formation in the LV filling pattern [1]. Vorticity, the curl of velocity, is a fundamental quantity in fluid mechanics that describes the local spinning rate of fluid particles and can characterize vortex flow [10]. Quantitative vortex parameters, such as vorticity, have been used to assess diastolic (dys) function in several patient groups [8, 9, 11, 12]. Furthermore, in patients with complex congenital intracardiac deformations such as after the Fontan operation, flow collision with remaining septal structures and stagnation of flow through a ventricular septal defect may result in altered EL and vortex formation [13].

Recently, good reproducibility of inflow- and outflowassessment from 4D flow MRI was shown [14]. However, there is a lack of studies validating the reproducibility of intracardiac energy and quantitative vorticity parameters from 4D flow MRI in a scan–rescan setting. Scan–rescan reproducibility is important for clinical applicability as it expresses the reliability in repeated quantitation, for example during serial follow-up or in case of a rest–stress protocol. Therefore, the aim of this study was to assess the scan–rescan reproducibility of 4D flow MRI measurements of kinetic energy, viscous energy loss and vorticity within the LV during diastolic filling in healthy subjects.

# Materials and methods

#### **Study population**

The study protocol was approved by the local Medical Ethical Committee of the Leiden University Medical Center and informed consent was obtained from all participants. Twelve healthy volunteers with no history of cardiac disease were included. All subjects underwent an MRI scan including whole-heart 4D flow MRI between July 2015 and April 2017. The same scanning protocol was performed twice in the same session with 10-min breaks between the scans and repositioning and replanning for every volunteer. Ten of these volunteers were included in a recent study [14]. That study did not report assessment of KE, EL or vorticity.

# Cardiovascular magnetic resonance acquisition and data preparation

Whole-heart 4D flow MRI was obtained on a 3 T scanner (Ingenia, Philips Medical Systems, The Netherlands) with maximal amplitude of 45 mT/m for each axis, slew rate of 200 T/m/s and a combination of FlexCoverage Posterior coil in the table top with a dStream Torso coil, providing up to 32 coil elements for signal reception. The orientation of the acquisition of 4D flow data was identical to the 4-chamber orientation (usually double-oblique axial or coronal). Velocity-encoding of 150 cm/s in all three directions was used in a standard four-point encoding scheme, spatial resolution  $3.0 \times 3.0 \times 3.0$  mm<sup>3</sup>, fieldof-view 400 mm, flip angle 10°, echo time (TE) 3.7 ms, repetition time (TR) 10 ms, true temporal resolution 40 ms, SENSitivity Encoding (SENSE) factor 2 in anterior-posterior direction and Echo Planar Imaging (EPI) readout with a factor 5. Free breathing was allowed and no respiratory motion compensation was performed. Retrospective gating was used with 30 phases reconstructed to represent one cardiac cycle. Expected scan-time for the 4D flow MRI acquisition for a patient with a heart rate of 60 bpm and 39 slices would be 9 min and 11 s. This 4D flow MRI sequence with EPI readout has been validated in vivo and in vitro [15] and compared to other 4D flow MRI sequences [16]. Concomitant gradient correction was performed from standard available scanner software. Cine two-dimensional (2D) left 2-chamber, 4-chamber, coronal and sagittal aorta views and a cine multi-2D short-axis stack of slices were acquired, using steady-state free-precession (SSFP) sequences with TE/TR 1.5/3.0, 350 mm field-of-view, 45° flip angle, acquisition resolution  $1.9 \times 2.0 \times 8.0$  mm<sup>3</sup>. Retrospective gating was used with 30 phases reconstructed to represent one cardiac cycle. Expected scan-time for the cine multi-2D short-axis acquisition for a patient with a heart rate of 60 bpm and 12 slices would be 1 min. Free breathing was allowed without using motion suppression, three signal averages were taken to minimize effects of breathing motion, which makes the expected scan-time 3 min. Image analysis was performed by one observer (VPK) with over 2 years of experience in MRI and verified by one observer (JJMW) with over 15 years of experience in MRI. The endocardial border was manually traced in all slices and phases in the multislice 2D cine short-axis images and ventricular volume was calculated at the end-diastolic and end-systolic phases using in-house developed MASS software. Papillary muscles were disregarded and assumed to be included in the ventricular volume. LV in- and outflow was assessed using the 4D flow MRI data with retrospective valve tracking of the mitral and aortic valve, as shown in a recent study

[14]. Cardiac output (CO) was computed from the 4D flow data as LV outflow × Heart rate (HR). Beginning and ending of diastolic phases [early diastolic filling (E-filling) and late diastolic filling (A-filling)] were derived from the mitral valve flow-time curves that resulted from retrospective valve tracking. Segmentation of the LV cavity in the 4D flow MRI acquisition, that is required for the energy analysis, was obtained by transforming the available time-varying segmentation of multi-slice cine short-axis acquisition to the 4D flow MRI data. To correct for patient motion related misalignment between the two acquisitions, automated image-based 3D rigid registration was performed using the phase with optimal depiction of the LV cavity in both scans with the Elastix image registration toolbox [17]. Kinetic energy, viscous energy loss and vorticity analysis of segmented LV volumes was done by one investigator (MSME) using in-house developed software.

#### **KE analysis over LV diastole**

The amount of KE during diastolic filling was computed following previously published methods [1]. KE for each voxel within the LV was computed as  $1/2 \text{ mv}^2$ , with (m) as the mass representing the voxel volume multiplied by the density of blood (1.025 g/mL) and (v) as the 3-directional velocity from 4D flow MRI. For each acquired time-phase, volumetric KE was then computed by integrating (by cumulative sum) the computed KE over the segmented 3D LV volume. In order to quantify KE during diastolic filling, the time-averaged KE during diastolic phases (KE<sub>E-avg</sub> and KE<sub>A-avg</sub>) and peak KE (KE<sub>E-peak</sub> and KE<sub>A-peak</sub>), in Joule, were assessed.

## Viscous EL analysis over LV diastole

Following recently published methods [1], we have computed EL from 4D flow MRI using the dissipation terms from the Navier-Stokes energy equations, assuming blood as a Newtonian fluid. The formulae that were used to calculate EL are summarized in "Appendix 1". The time-averaged viscous energy loss rate [EL, in Watt (W)] during E-filling  $(\overset{\bullet}{\text{FLE-avg}})$ and A-filling  $(\overset{\bullet}{ELA-avg})$  and  $\overset{\bullet}{EL}$  peaks  $(\overset{\bullet}{ELE-peak}$  and  $\overset{\bullet}{ELA-peak})$ were assessed. We have used the previously reported correlation between KE and  $\mathbf{\dot{FI}}$  [1] as a measure of in-scan consistency. Furthermore, EL over the total diastole ( $EL_{total diastole}$ ) in Joule (J) was computed. Given that the amount of viscous energy lost is proportional to the amount of kinetic energy, we computed a dimensionless energy loss parameter, EL<sub>index</sub>, that reflects the amount of viscous energy loss indexed for the average kinetic energy over diastole. EL<sub>index</sub> is a dimensionless index that was used in an earlier echo particle image velocimetry study by Agati et al. [18] to indicate the relative

amount of kinetic energy lost to that available over cardiac cycle. In this study,  $EL_{index}$  was computed over diastole as  $EL_{total diastole}/KE_{average diastole}$ , with  $KE_{average diastole}$  being the average KE during total diastole. In order to compare our results with a previous study reporting EL values normalized by stroke volume (SV) [1], we also report EL values in the current study as normalized by SV (reported as norm\_ELE-avg, norm\_ELE-peak, norm\_ELA-avg, norm\_ELA-peak and norm\_EL\_total diastole). To be consistent with the previous study SV was derived from cine SSFP short-axis slices.

# Integral vorticity magnitude over LV diastole (vorticity\_LV)

The formulae that were used to calculate the integral vorticity magnitude are shown in "Appendix 1". In short, following previously published work [8, 9], for each acquired time-phase, voxel-wise vorticity magnitude (1/s) was first computed over the segmented LV volume. Then, the instantaneous integral vorticity magnitude was computed as the cumulative sum of voxel-wise vorticity and multiplied by voxel volume to give the integral in (milliliter  $\times$  1/second) i.e. (mL/s). Note that the computed vorticity integral parameter is a scalar quantity and therefore does not take the vorticity direction into account. We will refer to this vorticity integral over the LV as vorticity LV throughout the text, to differentiate it from voxel-wise vorticity. In order to quantify the integral vorticity\_LV over diastolic filling, the time-average and peak vorticity\_LV during E-filling (vorticity\_LV<sub>E-avg</sub>, vorticity\_LV<sub>E-peak</sub>, respectively) and A-filling (vorticity\_ $LV_{A-avg}$ , voriticity\_ $LV_{A-peak}$ , respectively) were computed.

# Scan-rescan analysis

For the scan–rescan analysis, all data was blinded by one observer (VPK) and presented in a random order to the observer (MSME) that performed the energy and vorticity analysis. Scan–rescan analysis was performed to test the reproducibility of (1) KE over E-filling and A-filling; (2) EL over E-filling, A-filling and total diastolic filling and (3) vorticity\_LV over E-filling and A-filling.

#### **Statistical analysis**

Data analysis was performed using SPSS Statistics software (v. 23.0 IBM SPSS, Chicago, IL). Variables were tested for normal distribution using the Shapiro–Wilk test. Continuous data was expressed as mean  $\pm$  standard deviation (SD) with minimum and maximum values or as median [interquartile range] where suitable. Mean differences were determined for inter-scan comparison and significance was tested by a paired samples *t* test or, in case of non-normality, the Wilcoxon signed-rank test. Differences were computed as: measurement in scan 2—measurement in scan 1. The coefficient of variation (CV) was calculated with the root mean square method [19]. Correlation between the in-scan and inter-scan measurements done in repeated scans was tested by the Pearson correlation coefficient (r), or the Spearman correlation coefficient ( $\rho$ ) in case of non-normality of the data. The approach described by Bland and Altman [20] was used to study systematic differences between measurements obtained from the two scans. Agreement between these measurements was assessed by determining the intra-class correlation (ICC) coefficient. Correlation and agreement were classified as follows: r/ $\rho$  and ICC > 0.95: excellent, 0.95–0.85: strong, 0.85–0.70: good, 0.70–0.5: moderate, < 0.5: poor. A *P* value < 0.05 was considered statistically significant.

# Results

Table 1 Baseline characte

Volunteer characteristics are shown in Table 1. Heart rate and cardiac output were not significantly different between the two scans (HR:  $60.8 \pm 7.8$  vs.  $59.9 \pm 6.9$  bpm, P = 0.52 and CO:  $5.5 \pm 0.9$  vs.  $5.6 \pm 1.3$  L/min, P = 0.75). 4D flow MRI data acquisition was successful in all volunteers. Figure 1a–c shows cross-sectional mapping of the volumetric measurements of KE, EL and vorticity inside the LV at peak E-filling in a standard 4-chambers view. An example of the temporal evolution of KE and  $\frac{1}{EL}$  over LV diastole is shown in one subject in Fig. 1d.

# Intra-scan comparison of energetics (KE vs. EL)

Figure 2 shows the in-scan comparison of KE versus EL. Comparison of  $KE_{E-avg}$  to  $\frac{1}{ELE-avg}$  showed strong-excellent correlation (scan 1: r = 0.92, P < 0.001; scan 2:  $\rho = 0.96$ , P < 0.001). Also, correlation between  $KE_{A-avg}$  and  $\cdot EL_{A-avg}$  was strong-excellent in both scans (scan 1:  $\rho = 0.87$ , P < 0.001; scan 2: r = 0.99, P < 0.001).

# Scan-rescan analysis of kinetic energy and viscous energy loss rate over early diastolic filling

Detailed results of the scan-rescan tests of KE and  $_{EL}$  assessment over early diastolic filling are shown in Tables 2 and 3 and Fig. 3. Scan-rescan assessment showed poor results for early diastolic filling, as shown in Table 2. Of note, Fig. 3 shows that for KE and  $_{EL}$  assessment over early diastolic filling two subjects showed more distinct differences. Throughout the text we will refer to these two subjects as "Subject 1" and "Subject 2". Detailed scan-rescan information of these two subjects is provided in "Appendix 2". In Fig. 3a, Subject 1 and 2 are indicated as dark ( $KE_{E-avg}$ ) and light red triangles ( $KE_{E-peak}$ ).

Possibly some factors related to the 4D flow MRI acquisition or physiological factors have resulted in these marked differences, therefore we performed an evaluation for a subcohort without Subject 1 and 2. Scan–rescan correlations were much stronger and variation was less for this sub-cohort, as shown in Table 3. KE<sub>E-avg</sub> and ELE-avg assessment showed non-significant differences between the two scans, strong ICCs (KE<sub>E-avg</sub>: 0.95, P < 0.001 and ELE-avg: 0.91, P=0.03) and CVs  $\leq 11\%$  (KE<sub>E-avg</sub>: 10% and ELE-avg: 11%). norm\_ELE-avg showed a strong ICC (0.90, P=0.001) and a CV of 12%. Scan–rescan assessment of KE<sub>E-peak</sub> and ELE-peak showed non-significant differences. The ICC of KE<sub>E-peak</sub> was good (0.82,

Scan-independent characteristics			
N	12		
Male (%)	6/12 (50%)		
Age (years)	$27 \pm 3$		
Height (cm)	$175 \pm 7$		
Weight (kg)	$69 \pm 12$		
BSA (m <sup>2</sup> )	$1.8 \pm 0.2$		
BMI (kg/m <sup>2</sup> )	$22 \pm 3$		
Characteristics per scan	Scan 1	Scan 2	P valu
HR (bpm)	$60.8 \pm 7.8$	$59.9 \pm 6.9$	0.52
End-diastolic volume (mL)	143.8 (132.4–183.8)	$158.6 \pm 30.3$	0.64
End-systolic volume (mL)	$59.0 \pm 12.8$	$56.2 \pm 12.9$	0.27
Stroke volume (mL)	$100.8 \pm 21.8$	$102.4 \pm 21.0$	0.44
Ejection fraction (%)	$62.8 \pm 3.6$	63.1 (61.6–65.4)	0.21
CO (L/min)	$5.5 \pm 0.9$	$5.6 \pm 1.3$	0.75

*bpm* beat per minute, *BMI* Body Mass Index, *BSA* body surface area, *CO* cardiac output, *HR* heart rate, LV left ventricular

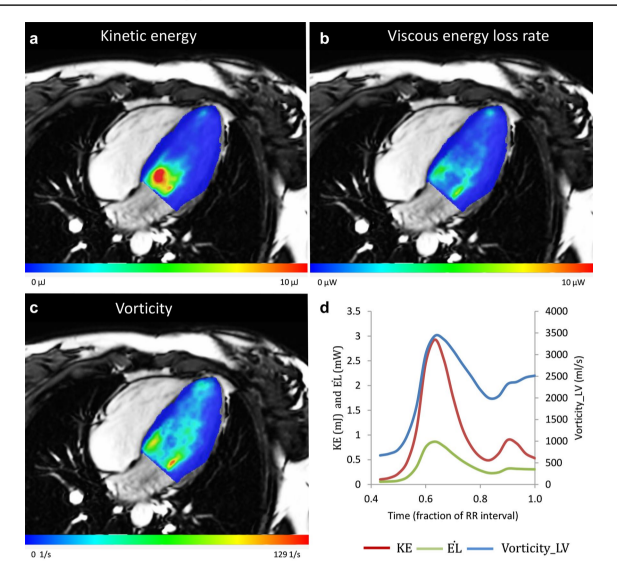


Fig. 1 Maps of left ventricular kinetic energy (KE), viscous energy loss rate (EL) and vorticity over the LV of a healthy female subject (age 20 years) in a standard 4-chambers MRI cross-sectional view, **a** left ventricular kinetic energy at peak early diastolic filling, **b** left

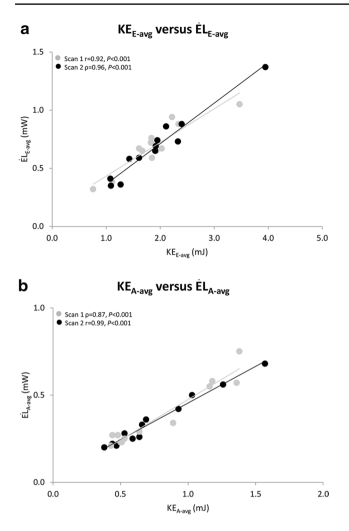
ventricular viscous energy loss rate at peak early diastolic filling, **c** left ventricular voxel-wise vorticity at peak early diastolic filling, **d** temporal evolution of volumetric kinetic energy, viscous energy loss rate and vorticity over LV diastole

P=0.01) with a CV of 15%. Similarly the ICC of ELE-peak was good (0.76, P=0.03) with a CV of 17%. Furthermore, norm\_ELE-peak showed a strong ICC (0.86, P=0.005) and a CV of 16%.

# Scan-rescan analysis of kinetic energy and viscous energy loss rate over late diastolic filling

Detailed results of the scan–rescan tests of KE and  $_{EL}$  assessment over late diastolic filling are shown in Tables 2 and 3 and Fig. 3. Scan–rescan assessment showed good results for late diastolic filling, as shown in Table 2. Subject 1 and 2 are

now within the same range as the other values [Fig. 3c, indicated as dark ( $KE_{A-avg}$ ) or light red triangles ( $KE_{A-peak}$ )]. The same was observed when assessing  $E_{LA-avg}$  and  $E_{LA-peak}$  [Fig. 3d, indicated as dark ( $E_{LA-avg}$ ) or light red triangles ( $E_{LA-peak}$ )]. However, to be consistent we repeated the evaluation in the sub-cohort without Subject 1 and Subject 2. Scan-rescan correlations and variation were similar for the sub-cohort without Subject 1 and Subject 2, as shown in Table 3. Reproducibility of  $KE_{A-avg}$  and  $E_{LA-avg}$  assessment showed non-significant differences, good ICCs ( $KE_{A-avg}$ : 0.77, P = 0.02 and  $E_{LA-avg}$ : 0.75, P = 0.03) and CVs up to 24% ( $KE_{A-avg}$ : 23% and  $E_{LA-avg}$ : 24%). norm\_ $-E_{LA-avg}$ 



**Fig.2** Scatter plots of kinetic energy (KE) versus viscous energy loss rate (EL) during early diastolic filling and KE versus EL during late diastolic filling. **a** Scatter plot depicting the correlation between KE<sub>E-avg</sub> and EL<sub>E-avg</sub> measured in scan 1 (grey) and scan 2 (black), in all 12 subjects, **b** scatter plot depicting the correlation between KE<sub>A-avg</sub> and EL<sub>A-avg</sub> measured in scan 1 (grey) and scan 2 (black), in all 12 subjects

showed a non-significant difference between the scans, a good ICC (0.70, P = 0.048) and a CV of 22%. Scan–rescan assessment of KE<sub>A–peak</sub> and  $\dot{E}_{LA-peak}$  showed non-significant differences, good ICCs (KE<sub>A–peak</sub>: 0.83, P = 0.01 and  $\dot{E}_{LA-peak}$ : 0.79, P = 0.02) and CVs up to 25% (KE<sub>A–peak</sub>: 23% and  $\dot{E}_{LA-peak}$ : 25%). Lastly, scan–rescan assessment of norm\_ $EL_{A-peak}$  showed a non-significant difference between the scans, a moderate ICC (0.64, P = 0.08) and a CV of 24%.

# Scan-rescan analysis of kinetic energy and viscous energy loss over total diastole

Detailed results of the scan-rescan tests of KE and  $\dot{EL}$  assessment over total diastole (EL<sub>total diastole</sub> and

norm\_EL<sub>total diastole</sub>) and EL<sub>index</sub> are shown in Tables 2 and 3 and Fig. 3. Scan-rescan assessment showed poor results for total diastole, but strong results for EL<sub>index</sub>, as shown in Table 2. Scan-rescan correlations were much stronger and variation was less for the sub-cohort without Subject 1 and Subject 2 (Table 3). Scan-rescan assessment of EL<sub>total diastole</sub> showed a non-significant difference between the scans, a strong ICC of 0.91 (P = 0.001) and a CV of 11%. norm EL<sub>total diastole</sub> showed a non-significant difference between the scans, a good ICC (0.81, P = 0.01) and a CV of 12%. Figure 3f shows the Bland-Altman plot of EL<sub>index</sub>. For all subjects, scan-rescan assessment of EL<sub>index</sub> showed excellent reproducibility with a small non-significant difference between the scans, a strong ICC of 0.94 (P < 0.001) and a CV of 9%. When evaluating the sub-cohort without Subject 1 and 2 the results remained similar (ICC: 0.95, P < 0.001and CV: 8%).

# Scan-rescan analysis of volumetric vorticity\_LV over early diastolic filling

Detailed results of the scan–rescan tests of vorticity\_LV assessment over early diastolic filling are shown in Tables 4 and 5 and Fig. 4. Scan–rescan assessment showed moderate results for early diastolic filling as shown in Table 4. The Bland–Altman plots of the assessment of vorticity\_LV<sub>E-avg</sub> and vorticity\_LV<sub>E-peak</sub> showed higher differences between scan and rescan measurements for Subject 1 and 2 (Fig. 4a, indicated as dark (vorticity\_LV<sub>E-avg</sub>) or light red triangles (vorticity\_LV<sub>E-peak</sub>)).

Scan–rescan correlations were much stronger and variation was less for the sub-cohort without Subject 1 and 2 (Table 5). Scan–rescan assessment of vorticity\_LV<sub>E-peak</sub> and vorticity\_LV<sub>E-avg</sub> showed a non-significant difference between the scans, good-strong ICCs (vorticity\_LV<sub>E-peak</sub>: 0.83, P = 0.01 and vorticity\_LV<sub>E-avg</sub>: 0.95, P < 0.001) and CVs up to 11% (vorticity\_LV<sub>E-peak</sub>: 11% and vorticity\_LV<sub>E-avg</sub>: 7%).

# Scan-rescan analysis of volumetric vorticity\_LV over late diastolic filling

Detailed results of the scan–rescan tests of vorticity\_LV assessment over late diastolic filling are shown in Tables 4 and 5 and Fig. 4. Scan–rescan assessment showed good results for late diastolic filling as shown in Table 4. Subject 1 and 2 are now within the same range as the other values (Fig. 4b, indicated as dark (vorticity\_LV<sub>A-avg</sub>) and light red triangles (vorticity\_LV<sub>A-peak</sub>)). Scan–rescan variation was similar for the sub-cohort without Subject 1 and 2 (Table 5). Scan–rescan assessment of vorticity\_LV<sub>A-peak</sub> and vorticity\_LV<sub>A-avg</sub> showed non-significant differences between the scans, good-strong ICCs (vorticity\_LV<sub>A-peak</sub>: 0.91, P=0.001 and vorticity\_LV<sub>A-avg</sub>: 0.89, P=0.002) and CVs of 11%.

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Table 2 Scan-rescan compa	rison of energy variables for th	Table 2 Scan-rescan comparison of energy variables for the cohort with all subjects $(N=12)$	12)			
	Scan 1	Scan 2	Difference (scan 2-scan 1)	Pearson correlation coefficient	Intraclass correlation coef- ficient	Coefficient of variation (%)
	Mean±SD (min–max) or median [IQR] (min–max)	Mean±SD (min-max) or median [IQR] (min-max)	Mean±SD or (min–max) median [IQR] (min–max)		ICC (95% CI)	
Early filling						
$KE_{F-neak}$ (mJ)	$4.9 \pm 1.9 \ (2.6 - 9.6)$	$4.8 \pm 1.4 \ (2.0 - 7.6)$	$0.1 \pm 2.3 \ (-4.8 \text{ to } 5.0)$	0.06	0.12 (-2.78 to 0.76)	28
$KE_{E-avg}$ (mJ)	$1.9 \pm 0.7 \; (0.8 - 3.5)$	$1.9 [1.3-2.3] (1.1-4.0)^{\circ}$	$-0.01 [-0.1 \text{ to } 0.3] (-2.2 \text{ to } 2.1)^{\circ}$	0.52	0.31 (-1.85 to 0.81)	26
$\dot{EL}_{P-mak}(mW)$	$1.5 \pm 0.4 \ (0.8 - 2.1)$	$1.5 \pm 0.6 \ (0.7 - 2.9)$	$0.04 \pm 0.7 \ (-1.4 \text{ to } 1.7)$	0.02	0.04 (-3.25 to 0.74)	31
norm_EL <sub>F-neak</sub> (W/m <sup>3</sup> )	15.2±5.1 (9.4–24.9)	$14.9 \pm 5.1 \ (6.0 - 21.8)$	$-0.2\pm6.2$ ( $-12.6$ to $12.3$ )	0.27	0.45 (-1.17 to 0.85)	30
ELE-avg (mW)	$0.7 \pm 0.2 \; (0.3 - 1.1)$	$0.7 \pm 0.3 \ (0.4 - 1.4)$	-0.01 [ $-0.1$ to $0.3$ ] ( $-0.7$ to $0.8$ ) <sup>c</sup>	0.11	0.21 (-2.36 to 0.78)	28
norm_EL <sub>E-avg</sub> (W/m <sup>3</sup> ) Late filling	7.0±2.3 (4.2–11.4)	<b>6.6±2.1</b> (3.2−10.4)	$-0.4\pm2.8(-6.4\text{ to }5.8)$	0.20	0.35 (-1.55 to 0.82)	28
$\mathrm{KE}_{\mathrm{A-peak}}$ (mJ)	0.9 [0.7–2.0] (0.7–2.5)°	1.0 [0.9–1.6] (0.6–2.6) <sup>c</sup>	$0.1 [-0.2 \text{ to } 0.3] (-1.2 \text{ to} 0.5)^{\circ}$	0.62 <sup>b</sup>	0.87 (0.55–0.96) <sup>b</sup>	23
${\rm KE}_{{\rm A-avg}}  ({ m mJ})$	$0.6 [0.5 - 1.2] (0.4 - 1.4)^{c}$	$0.8 \pm 0.4 \ (0.4 - 1.6)$	$-0.03 \pm 0.3 (-0.7 \text{ to } 0.4)$	$0.68^{\mathrm{b}}$	$0.84 \ (0.42 - 0.95)^{\rm b}$	23
EL <sub>A-peak</sub> (mW)	$0.5 \pm 0.3 \ (0.3 - 1.1)$	$0.5 \pm 0.2 \ (0.3 - 1.0)$	$-0.007 \pm 0.2 \ (-0.5 \text{ to } 0.3)$	$0.67^{\rm b}$	$0.81 (0.33 - 0.95)^{b}$	24
norm_EL <sub>A-reak</sub> (W/m <sup>3</sup> )	$5.3 \pm 1.8 \ (2.5 - 8.3)$	$5.1 \pm 1.5 \ (3.3 - 7.9)$	$-0.2 \pm 1.7 (-3.2 \text{ to } 2.2)$	0.50	0.68 (-0.19 to 0.91) <sup>b</sup>	22
ELA-ave (mW)	0.3 [0.2–0.6] (0.2–0.7) <sup>c</sup>	$0.4 \pm 0.2 \ (0.2 - 0.8)$	$-0.02 \pm 0.1 \ (-0.4 \text{ to } 0.2)$	0.50	$0.80 (0.29 - 0.94)^{\rm b}$	23
norm_EL <sub>A-avg</sub> (W/m <sup>3</sup> ) Total diastole	3.7±1.3 (2.1−6.6)	3.4±1.1 (2.2−5.2)	$-0.3 \pm 1.1 (-2.8 \text{ to } 1.5)$	0.58 <sup>b</sup>	0.73 (0.07–0.92) <sup>b</sup>	21
$\mathrm{EL}_{\mathrm{total diastole}}\left(\mathrm{mJ} ight)$	$0.5 \pm 0.2 \ (0.3 - 1.0)$	0.5 [0.4–0.6] (0.3–1.3)°	-0.006 [-0.07 to 0.06] $(-0.4 to 0.7)^{c}$	0.81 <sup>b</sup>	0.58 (-0.60 to 0.88)	22
norm_EL <sub>total diastole</sub> (J/m <sup>3</sup> )	$5.3 \pm 1.5 \ (3.4 - 8.1)$	4.8 [4.2-5.3] (3.1-9.8) <sup>c</sup>	$-0.1 \pm 2.2 (-3.9 \text{ to } 5.3)$	0.30	0.17 (-2.52 to 0.77)	23
$EL_{index}$	$0.3 \pm 0.07 \ (0.2 - 0.4)$	$0.3 \pm 0.06 \; (0.2 - 0.4)$	$-0.001 \pm 0.03 (-0.04 \text{ to} 0.06)$	$0.89^{a}$	$0.94 \ (0.80-0.98)^{a}$	6
KF kinetic energy E early di	KF binatic anarrov E aarlo diastolic fillino - 4 lata diastolic fillino - FI viscons anarrov loss	flling FI viscous anaray loss				

KE kinetic energy, E early diastolic filling, A late diastolic filling, EL viscous energy loss

 $<sup>{}^{</sup>a}P < 0.001$ 

 $<sup>^{</sup>b}P < 0.05$ 

 $<sup>^{\</sup>circ}P\!<\!0.05$  for non-normality (Shapiro–Wilk test)

	Scan 1	Scan 2	Difference (scan 2-scan 1)	Pearson correlation		Coefficient of
	Mean±SD (min-max) or median [IQR] (min-max)	Mean±SD (min–max) or median [IQR] (min–max)	Mean±SD or (min–max) median [IOR] (min–max)	coefficient	ncient ICC (95% CI)	variation (%)
Harly filling						
Early mung VE (mT)	5 0 E 1 0 5 31 7 0 5 4VC	111066M	$0.1 \pm 0.04 \pm 0.045$	0 67		15
KE. (mI)	$1.7 \pm 0.5(0.8-2.3)$	$1.8 \pm 0.5$ (1 1–2 4)	$0.1 \pm 0.5 (-0.5 \text{ (0} 2.0))$	0.02 0.91 <sup>a</sup>	0.95 (0.82–0.90) 0.95 (0.82–0.99) <sup>a</sup>	10
FI E much (mW)	$1.5 \pm 0.4 \ (0.8 - 2.1)$	$1.5 \pm 0.4 \ (0.7 - 2.2)$	$0.01 \pm 0.4 \ (-0.5 \ to \ 0.9)$	0.58	$0.76 (-0.07 \text{ to } 0.94)^{\text{b}}$	17
norm_FI F_mark (W/m <sup>3</sup> )	15.4±5.2 (9.6–24.9)	$15.2 \pm 4.2 \ (8.9 - 21.8)$	$-0.3 \pm 3.5 (-4.7 \text{ to } 6.9)$	0.75 <sup>b</sup>	0.86 (0.40–0.97) <sup>b</sup>	16
FI H-ave (mW)	$0.7 \pm 0.2 \ (0.3 - 0.9)$	$0.6 \pm 0.2 \ (0.4 - 0.9)$	$-0.02\pm0.1\ (-0.2\ \text{to}\ 0.2)$	0.83 <sup>b</sup>	$0.91 (0.64-0.98)^{b}$	11
norm_EL <sub>E-avg</sub> (W/m <sup>3</sup> ) 1 ate fillino	7.0±2.3 (4.2–11.4)	$6.6 \pm 1.6 \; (4.2 - 9.0)$	$-0.4 \pm 1.2 (-2.4 \text{ to } 1.6)$	0.87 <sup>b</sup>	0.90 (0.60–0.97) <sup>b</sup>	12
KE <sub>A-peak</sub> (mJ)	0.9 [0.7–1.7] (0.7–2.5) <sup>c</sup>	$1.2 \pm 0.5 \ (0.6 - 2.4)$	$0.2 [-0.1 \text{ to } 0.3] (-1.2 \text{ to } 0.5)^{\circ}$	0.59	0.83 (0.25–0.96) <sup>b</sup>	23
${ m KE}_{ m A-avg}(mJ)$	$0.5 [0.5-1.2] (0.4-1.4)^{c}$	$0.7 \pm 0.3 \ (0.4 - 1.3)$	$-0.02 \pm 0.3 (-0.7 \text{ to } 0.4)$	$0.65^{\mathrm{b}}$	0.77 (0.02–0.94) <sup>b</sup>	23
EL <sub>A-neak</sub> (mW)	0.4 [0.3–0.7] (0.3–1.1) <sup>c</sup>	$0.5 \pm 0.2 \ (0.3 - 0.9)$	$-0.03 \pm 0.2 (-0.5 \text{ to } 0.3)$	0.50	$0.79 (0.11 - 0.95)^{b}$	25
norm_ELA_neak (W/m <sup>3</sup> )	$5.3 \pm 2.0 \ (2.5 - 8.3)$	$5.0 \pm 1.3 \ (3.3 - 7.0)$	$-0.3 \pm 1.7 (-3.2 \text{ to } 2.2)$	0.49	0.64 (-0.56 to 0.91)	24
$\dot{EL}_{A-avg}(mW)$	$0.3 [0.2-0.6] (0.2-0.8)^{\circ}$	$0.3 \pm 0.1 \ (0.2 - 0.6)$	$-0.03 \pm 0.1 (-0.4 \text{ to } 0.2)$	0.42	$0.75 (-0.05 \text{ to } 0.94)^{\text{b}}$	24
norm_EL <sub>A-avg</sub> (W/m <sup>3</sup> ) Total diastole	$3.6 \pm 1.4 \ (2.1 - 6.6)$	3.4±1.0 (2.3-5.2)	$-0.3 \pm 1.2 (-2.8 \text{ to } 1.5)$	0.57	0.70 (- 0.21 to 0.93) <sup>b</sup>	22
$\mathrm{EL}_{\mathrm{total}\mathrm{diastole}}(\mathrm{mJ})$	$0.5 \pm 0.2 \ (0.3 - 1.0)$	$0.5 \pm 0.1 \ (0.3 - 0.7)$	$-0.01 [-0.05 to 0.03] (-0.3 to 0.03)^{\circ}$	0.88 <sup>b</sup>	0.91 (0.64–0.98) <sup>b</sup>	11
norm_EL <sub>total diastole</sub> (J/m <sup>3</sup> )	$5.1 \pm 1.3$ (3.4–7.0)	$4.8 \pm 0.9 (3.1 - 6.4)$	$-0.3 \pm 0.9 (-2.0 \text{ to } 1.2)$	$0.73^{\mathrm{b}}$	$0.81 (0.27 - 0.95)^{\rm b}$	12
EL <sub>index</sub>	$0.3 \pm 0.06 \ (0.2 - 0.4)$	$0.3 \pm 0.07 \ (0.2 - 0.4)$	$-0.01 \pm 0.03 (-0.04 \text{ to } 0.06)$	$0.89^{a}$	$0.95 (0.79 - 0.99)^{a}$	8

KE kinetic energy, E early diastolic filling, A late diastolic filling, EL viscous energy loss

\*Subjects with measurements outside the 95% confidence interval of the respective Bland-Altman plots

 $^{a}P < 0.001$ 

 $^{\rm b}P<0.05$ <br/> $^{\rm c}$ P<0.05 for non-normality (Shapiro–Wilk test)

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## Discussion

In the current study, kinetic energy, viscous energy loss and vorticity inside the left ventricle during diastole are derived from 4D flow MRI and scan-rescan reproducibility of these parameters is tested. Scan-rescan reproducibility is essential for the clinical application of a parameter since it reflects the reliability of a measurement and feasibility of repeated measurement evaluations. The main findings of this study are: (1) internal consistency between kinetic energy and viscous energy loss is strong-excellent in both scans during early and late diastolic filling; (2) In the majority (10 out of 12) of subjects, reproducibility of peak and average kinetic energy and viscous energy loss during early, late and total diastolic filling shows non-significant differences but with a good to excellent agreement (by means of ICC) and CVs up to 25%; (3) In the studies parameters, time-averaged measurements over E- and A-filling show stronger reproducibility than peak measurements. (4) For all subjects, EL<sub>index</sub> shows good reproducibility with a small non-significant difference between the scans, strong agreement and a CV of 9%; (5) Assessment of volumetric vorticity over the left ventricle during early and late diastolic filling shows non-significant differences, good-excellent ICCs and CVs up to 11%.

In the Bland-Altman plots for E-filling parameters, measurements obtained in two subjects (Subject 1 and 2 in Appendix 2) showed distinct higher differences between scan and rescan measurements than the measurements of the majority of the cohort. These high differences had an impact on the scan-rescan reproducibility for KE and  $\mathbf{FI}$  measures. This impact seemed to be largest during early diastolic filling, where reproducibility was much higher for the sub-cohort without Subject 1 and 2. However, during late diastolic filling, reproducibility was similar. There is no obvious explanation for the measurements that showed distinct higher differences between scan and rescan measurement, but this could be related to the technical restrictions of the 4D flow MRI acquisition or to physiological differences between scans, or a combination of both. It is to be expected that physiological differences were small as all subjects are healthy volunteers that were scanned twice under the same circumstances with only a short break between the scans  $(\pm 10 \text{ min})$  and heart rate was not significantly different between both scans. Still, subtle physiological differences could result in poor scan-rescan reproducibility for these few cases. Table 6 in "Appendix 2" shows the HR, LV outflow and CO of all subjects. Subject 1 and 2 present the highest CO differences, however there are other subjects with CO differences within the same range. Table 7 in "Appendix 2" shows subjectspecific scan-rescan information from the mitral valve flow curves. Subject 1 and 2 have the highest difference in area under the curve of the E-filling and peak filling rate of the E-filling. This could indicate that differences may be related to technical restrictions of the acquisition, as the E-filling duration, E peak

filling rate and the area-under-the-curve are sensitive to temporal and spatial resolution. These observations are important to take into consideration when evaluating flow energetics and vorticity in a research or clinical setting.

#### Kinetic energy (KE) over LV diastole

Multiple 4D flow MRI studies showed that patients with LV dysfunction present altered flow patterns through the LV with impaired preservation of inflow KE to the end of diastole and altered KE-time curves (the amount of KE inside the LV during each time step over the total cardiac cycle) [4, 6, 7,21], even in patients with normal to mild LV remodeling and normal to mildly depressed LV systolic function [7]. These KE changes in the LV could be a valuable diagnostic marker to evaluate diastolic function and might be useful for early detection of deteriorating ventricular function [1, 4–7, 21, 22], which could reduce patient morbidity and mortality [23]. However, there is a lack of studies validating LV KE derived from 4D flow MRI in a scan-rescan setting. Therefore, reliability and reproducibility of KE measurements from 4D flow MRI in a repeated scan setting remains largely unknown. Kanski et al. [24] compared mean KE and peak KE between two scans (with and without respiratory gating) with the aim to evaluate the impact of respiratory gating on KE measurements. They found a strong correlation between the KE measurements in both scans. The current study differs from the study by Kanski et al. [24] in that we used the same protocol for both scans. In the current study, moderate-strong correlation was found between KE measurements in both scans. Absolute values of KE<sub>E-peak</sub> and KE<sub>A-peak</sub> reported in the current study are in agreement with previous studies [1, 25, 26].

In the current study we showed the reproducibility of the KE measurements assessed with 4D flow MRI with good-strong agreement (by means of ICCs), however substantial CVs up to 23% were also found. Based on the Bland-Altman analysis the variability of  $KE_{E-avg}$ and  $KE_{A-avg}$  was less than the variability of  $KE_{E-peak}$  and  $KE_{A-peak}$  as shown by the smaller limits of agreement. This might be expected, given that the definition of average is computed over multiple time points and therefore evens out the variations more than a single-time measure such as the peak, especially when the definition of the peak also is affected by the temporal resolution of the 4D flow MRI data.

#### Viscous EL over LV diastole

The assessment of intracardiac EL could provide crucial details on the function of the heart apart from the standard MRI parameters and could be used to further unravel the influence of complex surgery for congenital heart defects [1].

To our knowledge, this is the first study assessing scan-rescan reproducibility of in vivo LV EL over diastole

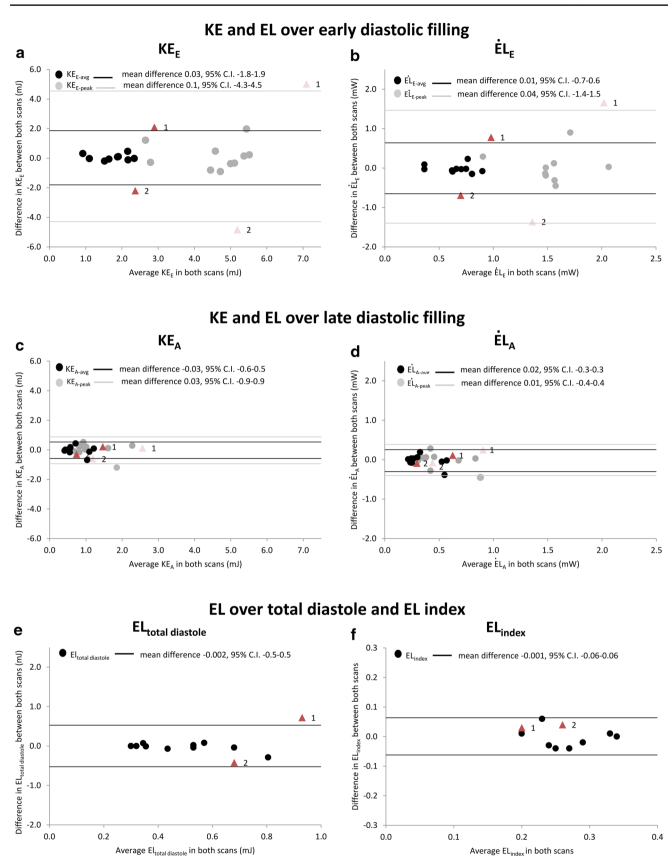


Fig. 3 Bland–Altman plots of kinetic energy (KE) and viscous energy loss rate (EL) at peak early diastolic filling ( $KE_{E-peak}$  and  $EL_{E-peak}$ ) and peak late diastolic filling (KE<sub>A-peak</sub> and  $EL_{A-peak}$ ), EL<sub>total diastole</sub> and ELindex. a Bland-Altman plot depicting the agreement between KE<sub>E-avg</sub> (black) and KE<sub>E-peak</sub> (grey) in scan 1 and scan 2. The subjects with distinct higher differences between scan and rescan measurements (Subject 1 and 2) are depicted as red triangles (KE<sub>E-avg</sub> dark red; KE<sub>E-neak</sub> light red). b Bland-Altman plot depicting the agreement between  $EL_{E-avg}$  (black) and  $EL_{E-peak}$  (grey) in scan 1 and scan 2. Subject 1 and 2 are depicted as red triangles (KE<sub>E-avg</sub> dark red; KE<sub>E-peak</sub> light red). c Bland-Altman plot depicting the agreement between  $\mathrm{KE}_{\mathrm{A-avg}}$  (black) and  $\mathrm{KE}_{\mathrm{A-peak}}$  (grey) in scan 1 and scan 2. Subject 1 and 2 are depicted as red triangles ( $KE_{A-avg}$  dark red;  $KE_{A-peak}$  light red). d Bland–Altman plot depicting the agreement between  $EL_{A-avg}$  (black) and  $EL_{A-peak}$  (grey) in scan 1 and scan 2. Subject 1 and 2 are depicted as red triangles (ELA-avg dark red; ELA-peak light red). e Bland-Altman plot depicting the agreement between EL<sub>total diastole</sub> in scan 1 and scan 2. Subject 1 and 2 are depicted as red triangles. f Bland-Altman plot depicting the agreement between EL<sub>index</sub> in scan 1 and scan 2. Subject 1 and 2 are depicted as red triangles

assessed by 4D flow MRI. Results of norm\_ELE-peak in the current study correspond well with that of a different healthy controls cohort of a recent 4D flow MRI study by Elbaz et al. [1] Also norm\_ELE-avg values are consistent. Furthermore, both results of norm\_ELA-peak, as well as norm\_ELA-avg are similar to previously reported numbers [1].

Both  $EL_{total diastole}$  and norm\_ $EL_{total diastole}$  are slightly higher in our study than in the study by Elbaz et al. [1]. However, results remain in the same range. These differences in results could be explained in part by differences in heart rate between the volunteers of this study and those of the previous study. This is because the total viscous energy loss over diastole is computed over the time period between the first and the last phase of the diastole (Eq. 3) in "Appendix 1") and heart rate mainly affects the duration of diastole. Still, the EL-time curve over total diastole in the current study (Fig. 1d) agrees well with reported in vivo [1] and in vitro [2] EL time curves.

In this study we demonstrated the reproducibility of the EL parameters with moderate–strong ICCs and substantial CVs of up to 25%. Similar to KE, the Bland–Altman analysis shows that the variability of  $EL_{E-avg}$  and  $EL_{A-avg}$  was less than the variability of  $EL_{E-peak}$  and  $EL_{A-peak}$ 

The amount of energy lost over diastole relative to the average kinetic energy as measured by means of  $EL_{index}$ , shows good reproducibility for all subjects with an ICC of 0.94 (P < 0.001) and an CV of 9%. The Bland–Altman plot shows that all subjects are within the same range of differences. This suggests that among tested parameters  $EL_{index}$  is the least sensitive to subtle physiological variations or discrepancies affected by technical limitations of the 4D flow MRI data in the healthy subjects, which might have affected the lesser reproducibility of KE and EL in some subjects. This observation could be attributed to the fact that  $EL_{index}$  is a dimensionless parameter concerned with the relative changes in EL to KE and not with

their absolute changes as in other tested parameters. As such, the reported reproducibility of  $EL_{index}$  could also be considered as another reflection of a good internal consistency in this study. Although a similar  $EL_{index}$  parameter was reported in a previous study and was shown to be significantly altered in patients with acute myocardial infarction [18],  $EL_{index}$  was computed in that study from 2D echo particle image velocimetry and over the complete cardiac cycle as compared to this study's volumetric measurement from 4D flow MRI and over diastole only. Therefore, it is not possible to perform a direct comparison with the published results of that study.

#### Vorticity inside the LV during diastolic filling

LV vortex quantification parameters, such as vorticity, could be useful in the assessment of LV and RV diastolic (dys-)function [1, 8, 9, 11, 12]. In recent studies, vorticity was shown to be a marker of diastolic dysfunction, both in the LV [9] and the RV [8] of patients with pulmonary hypertension.

To our knowledge, no previous study is available on assessing scan-rescan reproducibility of in vivo vorticity LV over diastole from 4D flow MRI. Fenster et al. [8] assessed vorticity inside the RV using the integral of vorticity magnitude over the volumes and found results in the same order of magnitude as the results in the current study. In a recent paper by Schafer et al. [9] LV vorticity in healthy subjects was assessed. However, it is not clear whether this was computed over the whole LV volume and therefore we cannot compare results to our measurements. The current study showed scan-rescan reproducibility of integral vorticity\_LV during E-filling as well as during A-filling with good-excellent ICCs and CVs up to 11%. Based on the Bland-Altman analyses, similar to the KE and EL results, the variability of the average of the vorticity\_LV<sub>E-avg</sub> and vorticity\_LV<sub>A-avg</sub> was less than the variability of the peaks vorticity\_LV<sub>E-peak</sub> and vorticity\_LVA-peak. Vorticity\_LVE-avg shows less variability than vorticity\_LV<sub>A-avg</sub> but vorticity\_LV<sub>E-peak</sub> shows slightly more variability than vorticity\_LV<sub>A-peak</sub>.

#### **Technical considerations**

The post-processing steps that are followed for obtaining these flow energetics and vorticity parameters involve manual segmentation, registration and valve tracking. Manual segmentation on cine short axis images can be performed with excellent reproducibility [27]. In this study, manual segmentation was performed by one observer with over 2 years of experience in MRI and verified by another observer with over 15 years of experience, with validated software [28], which warrants high accuracy. Next, to correct for patient motion related misalignment and minimize errors between the cine short axis and the 4D flow acquisitions,

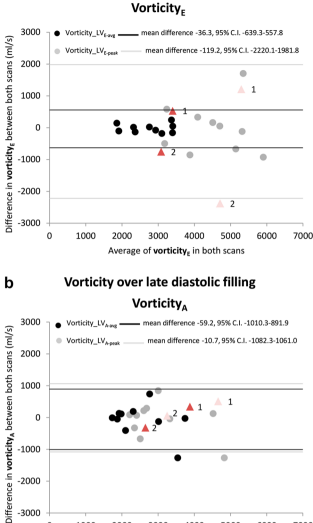
	JCall I	Scan 2	Difference (scan 2-scan 1)	Pearson correla- tion coefficient	correla- Intraclass correla- icient tion coefficient	la-	Coefficient of variation (%)
	Mean±SD (min-max) or median [IQR] (min-max)	Mean±SD (min-max) or median [IQR] (min-max)	Mean ± SD (min-max) or median [IQR] (min-max)	_			
Early nuing Vorticity_LV <sub>E-peak</sub> (mL/s)	4671±992 (2950–6370)	4551±1043 (2930–6200)	$-120 \pm 1075 (-239 \text{ to } 1710)$	0.44	0.63 (-0.37 to 0.90)	Q	15
Vorticity_LV <sub>E-avg</sub> (mL/s) Late filling	2862±636 (1720–3840)	2788±671 (1810–3930)	$-74\pm608$ ( $-1510$ to 1060)	) 0.57	0.74 (0.06–0.93) <sup>b</sup>		13
Vorticity_LVA-peak (mL/s)	2715 [2502–4142] (2176– 5463) <sup>c</sup>	$3193 \pm 945 (2174 - 4914)$	$-10\pm547$ ( $-1270$ to 840)	0.75 <sup>b</sup>	0.92 (0.73–0.98) <sup>a</sup>		11
Vorticity_ $LV_{A-avg}$ (mL/s)	$2660 \pm 842 (1740 - 4180)$	$2599 \pm 766 \ (1730 - 4050)$	$-61 \pm 484 (-1270 \text{ to } 740)$	$0.82^{b}$	$0.91 \ (0.68 - 0.97)^a$		11
	Scan 1 Mean±SD (min-max) or	Scan 1 Scan 2 Difference (sca $Mean \pm SD$ (min-max) or $Mean \pm SD$ (mi	Difference (scan 2-scan 1) Mean ± SD (min-max) or	Pearson correlation coefficient	Intraclass correlation coefficient ICC (95% CI)	tion	Coefficient of variation (%)
	median [IQR] (min-max)	median [IQR] (min-max)	median [IQR] (min-max)				
Early filling							
Vorticity_LV <sub>E-peak</sub> (mL/s)	$4546 \pm 1008 \ (2950 - 6370)$	$4520 \pm 1004 \ (2930 - 6200)$	$-27 \pm 796 (-940 \text{ to } 1710)  0.69^{\text{b}}$	0.69 <sup>b</sup>	0.83 (0.27–0.96) <sup>b</sup>	<u>_</u>	11
Vorticity_LV <sub>E-avg</sub> (mL/s) Late filling	2764±614 (1720–3560)	$2720 \pm 613 (1810 - 3610)$	$-44\pm277$ (-380 to 480)	0.90 <sup>a</sup>	0.95 (0.80–0.99) <sup>a</sup>	-	L
Vorticity_LVA-peak (mL/s)	2563 [2468–3628] (2176– 5463) <sup>c</sup>	$3012 \pm 852 \ (2174 - 4586)$	$-69\pm575$ ( $-1270$ to 840)	0.64 <sup>b</sup>	0.91 (0.64–0.98) <sup>b</sup>		11
Vorticity_LVA-avg (mL/s)	2269 [1889–3251] (1740– 4180) <sup>c</sup>	$2465 \pm 679 \ (1730 - 3730)$	$-75\pm511$ (- 1270 to 740) 0.82 <sup>b</sup>	0.82 <sup>b</sup>	0.89 (0.53–0.97) <sup>b</sup>		11

\*Subjects with measurements outside the 95% confidence interval of the respective Bland–Altman plots

 $^{a}P < 0.001$ 

 $^{b}P < 0.05$ 

 $^{\rm c}P<0.05$  for non-normality (Shapiro–Wilk test)



5000

6000

7000

4000

Average of vorticity, in both scans (ml/s

# а Vorticity over early diastolic filling **Vorticity**<sub>F</sub>

Fig. 4 Bland-Altman plots of volumetric vorticity over peak early filling (vorticity\_LV<sub>E-avg</sub> and vorticity\_LV<sub>E-peak</sub>) and late filling (vorticity\_LV<sub>A-avg</sub> and vorticity\_LV<sub>A-peak</sub>). **a** Bland–Altman plot depicting the agreement between vorticity\_LV<sub>E-avg</sub> (black) and vorticity\_LV<sub>E-peak</sub> (grey) in scan 1 and scan 2. Subject 1 and 2 are depicted as red triangles (vorticity\_LV<sub>E-avg</sub> dark red; vorticity\_  $LV_{E-peak}$  light red). **b** Bland-Altman plot depicting the agreement vorticity\_LV\_{A-avg} (black) and vorticity\_LV\_{A-peak} (grey) in scan 1 and scan 2. Subject 1 and 2 are depicted as red triangles (vorticity\_

3000

0

-1000

-2000

3000

0

1000

2000

LV<sub>A-avg</sub> dark red; vorticity\_LV<sub>A-peak</sub> light red)

automated image-based 3D rigid registration was performed using the validated Elastix image registration toolbox [17]. Another post-processing step requires retrospective valve tracking and mitral flow velocity mapping for assessing the beginning and ending of diastole. The reproducibility and observer variability of this semi-automated method was shown to be excellent [14, 15].

A limitation of this study is the small number of subjects. Furthermore, no patients were scanned in this study as this is difficult to assess in clinical research. In most of our patient studies, 4D flow MRI is part of a clinical CMR evaluation of about 75–90 min, sometimes involving the use of contrast for late gadolinium enhancement and sometimes involving a Dobutamine rest/stress protocol. Scan-rescan evaluation in such cases would imply repeating some of these evaluations as well. Furthermore, repeating 4D flow MRI for scan-rescan purposes would imply an additional scan-time of approximately 30 min (because of replanning), which makes the total scan-time too long to keep the patient in the same physiological state. Another limitation is that the influence of a different scanning protocol or scanner was not assessed. However, it is important to note that in the current study the aim was to assess scan-rescan reproducibility by using the exact same protocol and the same scanner machine twice. The use of different scanners or scanning protocols could result in altered reproducibility, which should be evaluated in future studies.

In conclusion, left ventricular kinetic energy and viscous energy loss quantification from 4D flow MRI in healthy volunteers shows strong-excellent in-scan consistency. Scan-rescan assessment of left ventricular kinetic energy, viscous energy loss and vorticity shows overall good agreement in the majority of the scanned subjects. Nevertheless, in two out of twelve subjects, considerable variation between the scans was found. Agreement of A-filling measurements is better than E-filling between scans in the studied parameters. Furthermore, time-averaged measurements over early and late filling show better reproducibility compared to peak measurements. Strong reproducibility for all cases is found in the dimensionless index, EL<sub>index</sub>, that measures the ratio of the amount of viscous energy lost relative to the average kinetic energy over diastole. ELindex seems to be less influenced by technical and/ or slight physiological differences between scans and may therefore be a useful parameter of energetics for future studies.

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## **Compliance with ethical standards**

Conflict of interest The authors declare that they have no conflict of interest.

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# **Appendix 1**

### Viscous energy loss computation from 4D flow MRI

Given the acquired velocity field v, the rate of viscous energy loss ( $\stackrel{\bullet}{\text{EL}}$ ) in watt (W) and the total energy loss (EL<sub>total</sub>) in joule (J) over a given period of time *T* can be computed from 4D flow MRI using the viscous dissipation function  $\boldsymbol{\Phi}_{v}$  in the Newtonian Navier–Stokes energy equations:

$$\Phi_{\nu} = \frac{1}{2} \sum_{i=1}^{3} \sum_{j=1}^{3} \left[ \left( \frac{\partial v_i}{\partial x_j} + \frac{\partial v_j}{\partial x_i} \right) - \frac{2}{3} (\nabla \cdot v) \delta_{ij} \right]^2, \\
\left\{ \begin{array}{l} \delta_{ij} = 1, & \text{if } i = j \\ \delta_{ij} = 0, & \text{if } i \neq j \end{array} \right] (1)$$

 $\Phi_v$  represents the rate of viscous energy dissipation per unit volume. *i*, *j* correspond to the principal velocity directions x, y, z.  $\nabla \cdot v$  denotes the divergence of the velocity field. Therefore, the rate of viscous energy loss ( $\stackrel{\bullet}{\text{EL}}$ ) in Watt at an acquired time phase t can be computed as:

$$\dot{EL}_{t} = \mu \sum_{i=1}^{M} \boldsymbol{\Phi}_{v} L_{i} [Watt(W)]$$
(2)

assuming the blood as a Newtonian fluid, the dynamic viscosity is  $\mu = 0.004$  Pa s, N as the total number of voxels in the given domain of interest (e.g. LV), L<sub>i</sub> as the voxel volume.

Table 6 Subject specific scan-rescan assessment

As a result, the total viscous energy loss (EL) in joules over time period T starting at phase  $t_{start}$  and ending at  $t_{end}$  can be computed as:

$$EL_{T} = \sum_{d=t_{start}}^{t_{end}} \dot{EL}_{d} p_{d} [Joule(J)]$$
(3)

with  $p_d$  the time step (temporal resolution) of the acquired 4D flow MRI.

#### Vorticity computation from 4D flow MRI

If *u*,*v*, *w* denote the three velocity field components acquired from 4D flow MRI over the principal velocity directions *x*, *y*, *z*, respectively, the vorticity ( $\omega_{i,t}$ ) at voxel *i* of an acquired time phase *t* 

$$\omega_{i,t} = \left(\frac{\partial w_{i,t}}{\partial y_{i,t}} - \frac{\partial v_{i,t}}{\partial z_{i,t}}, \frac{\partial u_{i,t}}{\partial z_{i,t}} - \frac{\partial w_{i,t}}{\partial x_{i,t}}, \frac{\partial v_{i,t}}{\partial x_{i,t}} - \frac{\partial u_{i,t}}{\partial y_{i,t}}\right) \left[1/s\right]$$
(4)

Then, Vorticity\_LV denoting the integral sum of vorticity over the segmented LV volume at an acquired time phase t in liter (mL) per second (s) can be computed as

Vorticity\_LV<sub>t</sub> = 
$$\sum_{i=1}^{M} |\omega_{i,t}| L_{i,t} [mL/s]$$
 (5)

With  $|\omega_{i,t}|$  as the magnitude of the vorticity vector, M as the total number of voxels in the segmented LV volume and  $L_{i,t}$  as the voxel volume.

# Appendix 2

See Tables 6 and 7.

Subject	HR 1 (bpm)	HR 2 (bpm)	Difference HR (bpm)	LV outflow 1 (mL)	LV outflow 2 (mL)	Difference LV outflow (mL)	CO 1 (L/min)	CO 2 (L/min)	Difference CO (%)
1 <sup>a</sup>	67.1	69.6	2.5	104.1	124.8	20.7	7.0	8.7	24.4
$2^{a}$	53.8	51.6	-2.2	112.9	94.5	-18.4	6.1	4.9	- 19.7
3	58.0	51.8	-6.2	110.0	126.4	16.4	6.4	6.6	2.7
4	61.5	59.1	-2.4	81.5	72.4	-9.1	5.0	4.3	- 14.7
5	71.9	66.2	-5.7	68.4	66.0	-2.4	4.9	4.4	-11.2
6	50.5	52.9	2.4	100.7	108.6	7.9	5.1	5.7	12.9
7	63.7	60.4	-3.3	104.0	107.8	3.8	6.6	6.5	-1.7
8	57.7	53.1	-4.6	85.4	86.4	1.0	4.9	4.6	-6.9
9	71.1	72.1	1.0	75.3	84.4	9.1	5.4	6.1	13.7
10	68.0	61.3	-6.7	85.0	81.8	-3.2	5.8	5.0	-13.3
11	48.5	58.3	9.8	90.3	88.7	-1.5	4.4	5.2	18.2
12	58.0	62.0	4.0	71.1	76.3	5.2	4.1	4.7	14.7

Differences were calculated as: value scan 2-value scan 1

A late diastolic filling, AUC area under the cure, CO cardiac output, E early diastolic filling, HR heart rate

<sup>a</sup>Subjects with marked high differences for the assessment of early filling parameters

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	C Differ-
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Table 7	Subject

Subjec	Subject E AUC 1 (mL)	E AUC 2 (mL)	Differ- ence E AUC (mL)	E PFR 1 (mL/s)	E PFR 2 (mL/s)	Differ- ence E PFR (mL/s)	E dur 1 (ms)	E dur 2 (ms)	Differ- ence E dur (ms)	A AUC 1 (mL)	A AUC 2 (mL)	Differ- ence A AUC (mL)	A PFR 1 (mL/s)	A PFR 2 (mL/s)	Differ- ence A PFR (mL/s)	A dur 1 (ms)	A dur 2 (ms)	Differ- ence A dur (ms)
1 <sup>a</sup>	78.6	94.8	16.2	522.8	738.8	216.0	332	292	-40	28.1	29.3	1.2	342.1	357.7	15.6	166	175	6
$2^{\mathrm{a}}$	84.3	71.3	-13.0	567.4	388.4	-179.0	323	413	06	19.1	14.1	-5.0	243.5	159.6	- 83.9	144	188	4
б	85.7	96.4	10.7	744.5	734.7	- 9.8	309	270	- 39	32.8	28.0	- 4.8	409.5	347.0	-62.5	172	194	22
4	68.5	61.4	-7.1	551.4	537.9	- 13.5	316	263	-53	15.3	12.4	-2.9	161.8	158.0	- 3.8	190	131	- 59
5	52.6	46.8	-5.8	343.6	362.2	18.6	348	297	-51	16.7	15.0	- 1.7	187.0	176.3	-10.7	174	208	34
9	83.9	84.6	0.7	500.2	545.2	45.0	378	328	-50	13.6	22.0	8.4	137.4	231.3	93.9	189	197	8
7	79.3	87.5	8.2	683.6	647.7	-35.9	314	351	37	32.3	29.6	-2.7	409.2	374.7	-34.5	157	160	3
8	72.2	71.6	-0.6	592.5	543.6	- 48.9	384	350	<b>-</b> 34	17.5	17.5	0.0	221.1	217.1	-4.0	175	176	1
6	59.4	66.8	7.4	506.1	563.6	57.5	272	300	28	18.3	19.6	1.3	243.7	245.0	1.3	136	137	1
10	58.7	58.9	0.2	367.5	376.7	9.2	322	373	51	17.6	14.4	-3.2	185.3	173.6	-11.7	162	170	8
11	64.5	6.09	-3.6	398.4	363.9	-34.5	349	343	9-0	17.5	14.4	-3.1	181.8	172.3	-9.5	209	205	-4
12	52.8	47.4	-5.4	380.3	314.4	-65.9	330	313	- 17	8.5	12.9	4.4	114.5	164.0	49.5	165	187	22
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<sup>a</sup>Subjects with marked high differences for the assessment of early filling parameters

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