How Similar Are the Mice to Men? Between-Species Comparison of Left Ventricular Mechanics Using Strain Imaging

Kenya Kusunose¹, Marc S. Penn¹, Youhua Zhang², Yuanna Cheng², James D. Thomas¹, Thomas H. Marwick¹, Zoran B. Popović¹*

¹Department of Cardiovascular Medicine, Cleveland Clinic, Cleveland, Ohio, United States of America, ²Department of Molecular Cardiology, Cleveland Clinic, Cleveland, Ohio, United States of America

Abstract

Background: While mammalian heart size maintains constant proportion to whole body size, scaling of left ventricular (LV) function parameters shows a more complex scaling pattern. We used 2-D speckle tracking strain imaging to determine whether LV myocardial strains and strain rates scale to heart size.

Methods: We studied 18 mice, 15 rats, 6 rabbits, 12 dogs and 20 human volunteers by 2-D echocardiography. Relationship between longitudinal or circumferential strains/strain rates (S_long/S_Rlong, S_circ/S_Rcirc), and LV end-diastolic volume (EDV) or mass were assessed by the allometric (power-law) equation $Y = kM^b$.

Results: Mean LV mass in individual species varied from 0.038 to 134 g, LV EDV varied from 0.015 to 102 ml, while RR interval varied from 81 to 1090 ms. While $S_{long}$ increased with increasing LV EDV or mass ($b$ values $0.047 \pm 0.006$ and $0.051 \pm 0.005$, $p<0.0001$ vs. 0 for both) $S_{circ}$ was unchanged ($p=N.S$ for both LV EDV or mass). Systolic and diastolic $S_{Rlong}$ and $S_{Rcirc}$ showed inverse correlations to LV EDV or mass ($p<0.0001$ vs. 0 for all comparisons). The ratio between $S_{long}$ and $S_{circ}$ increased with increasing values of LV EDV or mass ($b$ values $0.039 \pm 0.010$ and $0.040 \pm 0.011$, $p>0.0003$ for both).

Conclusions: While $S_{circ}$ is unchanged, $S_{long}$ increases with increasing heart size, indicating that large mammals rely more on long axis contribution to systolic function. $S_{Rlong}$ and $S_{Rcirc}$, both diastolic and systolic, show an expected decrease with increasing heart size.

Introduction

The structure of mammalian hearts is grossly invariant between the species. Heart size maintains constant proportion to whole body size, while left ventricular (LV) myocardial histology shows a stable pattern of distribution of fibers that are oriented obliquely in subendo- and subepicardium, and circumferentially in mid-myocardium in all species. [1–2]. In contrast, myocardial function parameters show complex scaling relationships to body size. These relationships are driven by the fact that smaller animals consume more energy per body weight. [3] Because of this, scaling of functional indices, such as heart rate intervals, myocardial long axis velocities, or intraventricular pressure differences, can be expected. While the link between organ function and body (or organ) size can be modeled by several nonlinear mathematical relationships, allometric (power law) equation is most widely used as it is parsimonious (i.e. with a smallest number of parameters), and because the value of scaling coefficient has an intrinsic significance. [4].

What is less understood is the impact of heart size on LV deformational mechanics, namely myocardial strain and strain rate. Normal myocardial strains quantify myocardial deformation occurring in the direction of the three principal axis of the heart, with longitudinal strain ($S_{long}$) reflecting long axis, and circumferential ($S_{circ}$) and radial ($S_{rad}$) strains reflecting corresponding components of short-axis deformation. Liu et al. have shown that $S_{circ}$ and $S_{rad}$ are decreased in mice and rats. [5] We have previously shown that ratio of long axis versus short axis displacement (which grossly represents ratio of $S_{long}$ to $S_{circ}$) decreases with decreasing heart size. [6] However, studies that quantitate the relationship between various strain components and body/organ size are lacking.

Systolic and diastolic strain rates, in turn, represent the rate of deformation. Strain rates are purportedly linked to contraction and relaxation at a more basic level. [7–8] Scaling of strain rates should be dependent on the interaction between scaling of myocardial velocities and left ventricular dimensions. Several papers, using both tissue Doppler and speckle tracking imaging, showed considerably higher strain rate in small rodents [9,10,11].

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* E-mail: popoviz@ccf.org
and in smaller individuals within the single species [12,13,14]. However, no systematic assessment of the impact of heart size on strain rates has ever been performed.

In this paper we use echocardiography to assess three normal components of strains and corresponding strain rates in five mammalian species that differ in their body size by more than three orders of magnitude. The aim of this study was to determine the presence and magnitude of the impact of heart size on strains and strain rates using an allometric scaling model. Our research hypotheses were that 1) normal myocardial strains show a small but significant decrease with decreasing heart size [5] (See Appendix S1 and Figure S1 for further details), 2) strain rates show a large increase with decreasing heart size (see Appendix S1), and 3) $S_{long}$ to $S_{circ}$ scale differently to changing heart size. [6].

### Methods

#### Study Population

Out of our database of 117 healthy volunteers free of any known cardiovascular disease, with normal physical examination, normal electrocardiogram, and not taking any cardio-active medication, we randomly selected 20 subjects whose age was between 21 and 45 years.

The studies in human subjects were performed in accordance with the Declaration of Helsinki. The Institutional Review Board of the Cleveland Clinic approved that study, and all participants gave written informed consent. The animal data were collected during echocardiographic evaluation of control (normal) animals used in several different studies in the period of 2003–2011. The data collection in each of these studies was prospective and specified by the respective study protocols. Importantly, the evaluation of the reported echocardiographic data was performed in each study and animal by using compatible equipment and techniques. All the animal experiments reported in this article were approved by the Institutional Animal Care and Use Committee of Cleveland Clinic and were in compliance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals.

#### Procedures

Subjects were studied in left lateral decubitus position. Due to the specificity of the animals’ behavior, the procedures differed slightly among the species. Twelve mongrel dogs were trained to lie down calmly for echocardiography. Six white New Zealand rabbits were lightly sedated with 35 mg/kg ketamine administered intramuscularly, with the handler keeping the head covered with soft cloth in a dark room. [15] Fifteen Lewis rats were sedated with 85 mg/kg of ketamine intraperitoneally [16] After ketamine administration, both rats and rabbits responded to transducer application to the chest wall, and they had to be gently restrained by the handler. Finally, eighteen C57/BL6 mice were assessed in a conscious state, with each one of them having at least one previous echocardiography session with the same handler. All animals were of adult age.

#### Data Collection

Echocardiography was performed using Vivid 7 echocardiography machine (GE Medical, Milwaukee, WI, USA). M-mode echocardiography, two dimensional echocardiography, and two-dimensional color tissue Doppler echocardiography data were collected using a dual harmonic 1.7/3.4 MHz or 2.0/4.2 MHz sector transducer (humans, dogs), 6 MHz or 11.5 MHz pediatric sector transducer (rabbits, rats) and 14 MHz epicardial linear transducer (mice). The minimal frame rates acquired during standard two-dimensional echocardiography in humans, dogs, rabbits, rats and mice were 30, 50, 70, 90 and 160 frames s$^{-1}$, respectively. The minimal frame rates acquired during color tissue Doppler two-dimensional echocardiography in humans, dogs, rabbits, rats and mice were 125, 125, 189, 200 and 208 frames s$^{-1}$, respectively. Data were digitized in a proprietary format for further analysis. Long axis views were obtained from parasternal window in mice, rats and rabbits, and from apical window in dogs and human.

#### Data Analysis

Data were analyzed using Echopac PC (GE Medical Systems, Milwaukee, WI). LV ejection time was measured directly from the digital pulsed-wave Doppler tracing of the LV outflow tract with the sweep speed set to 200 mm s$^{-1}$. LV end-diastolic and end-systolic volumes were measured from the long-axis view by the single-plane Simpson equation. LV mass was calculated from two-dimensional echocardiography data by the bullet equation:

$$\text{LV mass (g)} = 1.05(5/6)[A_{\text{Epi/L}} + (A_{\text{Epi}}/\pi - (A_{\text{Endo}}/\pi) - A_{\text{Endo}})$$

where $1.05$ is the specific gravity of muscle, $A_{\text{Epi}}$ and $A_{\text{Endo}}$ are the epi- and endocardial parasternal short-axis areas, respectively, and L is the parasternal long-axis LV length. LV mass was calculated using both end-systolic and end-diastolic data, and the two estimates were then averaged. Previous studies have validated LV mass measurement based on two-dimensional echocardiography in mice [17], rabbits [18], dogs [19] and humans [20]. Additionally, the validation of LV mass estimation by the bullet equation in mice, rats, dogs and humans from our laboratory was already published. [6].

#### Speckle tracking imaging

We used a speckle-tracking algorithm incorporated into Echopac PC (GE Medical Systems, Milwaukee, WI). The speckle-tracking analysis was performed by the same trained observer. The region of interest

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**Table 1. Basic morphometric and echocardiography data of different species.**

| Specie   | n  | Body weight (kg) | RR interval (ms) | ET (ms) | LV D/L ratio | EDV (mL) | ESV (mL) | EF | LV mass (g) |
|----------|----|------------------|-----------------|--------|--------------|----------|----------|----|-------------|
| Mouse    | 18 | 0.033±0.006      | 92.5±11.1       | 39±6   | 0.49±0.08    | 0.032±0.017 | 0.009±0.004 | 0.68±0.06 | 0.068±0.02 |
| Rat      | 15 | 0.264±0.025      | 143.2±17.1      | 63±8   | 0.52±0.10    | 0.255±0.09  | 0.076±0.028 | 0.69±0.07 | 0.728±0.17 |
| Rabbit   | 6  | 3.01±0.39        | 296.8±36.1      | 125±17 | 0.66±0.11    | 2.9±0.8   | 1.1±0.5   | 0.61±0.08 | 3.6±0.5    |
| Dog      | 12 | 26.3±5.9         | 501.3±38.8      | 157±12 | 0.65±0.10    | 57.4±5.9  | 22±6.9   | 0.62±0.06 | 86.1±19    |
| Human    | 20 | 61.7±9.8         | 891.1±116.1     | 297±28 | 0.55±0.05    | 83.5±13   | 32.9±8.5  | 0.61±0.07 | 100.3±23.6 |

Abbreviations: ET: ejection time; D/L ratio: diameter/length ratio (diastole); EDV: end diastolic volume; ESV: end systolic volume; EF: ejection fraction; LV: left ventricular.

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was overlaid across a cross section of the ventricular silhouette at the image corresponding to the minimal endocardial area. The software algorithm then automatically divided the LV long-axis view and LV short-axis view into six segments for speckle tracking throughout the cardiac cycle. The tracking quality was then visually inspected, and, if it was satisfactory for at least five segments, the tracing was accepted. In mice, since our experiments were done in conscious setting without the possibility of registering the electrocardiogram, we manually defined end-systole and end-diastole. [21] Segmental longitudinal strain (S_{long}) and strain rate (SR_{long}), circumferential strain (S_{circ}) and strain rate (SR_{circ}), and radial strain (S_{rad}) and strain rate (SR_{rad}) curves were then constructed and then averaged to obtain global segmental strain and strain rate curves. End-systolic strains were then obtained from the global strain curves, while peak systolic and diastolic strain rates were calculated from global strain rate curves. For the analysis of human, dog and rabbit data, at least three beats were measured. For small rodents, at least six beats (for rats) or nine beats (for mice) were averaged. The mean value was used for statistical analysis. Long – short axis strain ratio (L/C strain ratio) was calculated by dividing longitudinal strain by circumferential strain.

**Tissue Doppler Imaging.** Tissue Doppler Imaging, while carrying higher noise and larger operator variability, has a higher sampling rate and therefore potentially better detection of short-duration, high frequency peaks. [22] Therefore, in order to check for possible underestimation of strain rates, we also assessed Tissue Doppler imaging data obtained in a 4 chamber view and analyzed using EchoPAC PC (GE Medical Systems, Milwaukee, WI). [10,11] Diastolic and systolic longitudinal strain rates (TDI-SR_{long}) were measured in mid interventricular septum using a strain rate length/region of interest of 0.6 mm/0.6 mm, 1.5 mm/1 mm, 3 mm/3 mm, 12 mm/6 mm and 15 mm/6 mm in mice, [23] rats, rabbits, dogs and humans, [24] respectively. The temporal smoothing filters were turned off for all measurements. The values obtained in 6 consecutive (mice and rats) or 3 consecutive (rabbits, dogs, and humans) cardiac cycles were averaged.

**Quantifying Observer Variability**
To estimate intraobserver variability, we used data obtained in 2 mice, 2 rats, 2 rabbits, 2 dogs, and 2 humans. The same observer...
Statistical Analysis

We used proposed power-law function to predict the magnitude of the impact of LV size on strain and strain rate [25]:

\[ Y = kM^\beta \]  \hspace{1cm} (1)

where \( M \) is LV heart size parameter (i.e. LV EDV or LV mass), \( Y \) is a dependent variable (i.e. strain, strain rate), \( k \) is constant, and \( \beta \) is the power of the scaling exponent. To fit the data into the power function (1), we used a nonlinear regression method with Levenberg-Marquardt algorithm (SPSS 10.0, SPSS, Inc., Chicago, IL, USA). Confidence intervals (95\%) for exponent \( \beta \) were calculated using an asymptotic error of estimate. A t test statistics was used to compare the values of exponent \( \beta \) compared to zero and obtained by different methods. [26]. Data are presented as mean ± SD or as exponent \( \beta \) ± standard error of estimate. A \( p \) value \(<0.05\) was considered significant.

Results

Basic morphometric and echocardiography parameters of different species are presented in Table 1. In our subjects, LV mass varied from 0.038 to 134 g. LV EDV varied from 0.015 to 102 ml, while RR interval varied from 81 to 1090 ms. Thus, body weight and heart size varied by up to four orders of magnitude among the subjects. Figure 1. represents characteristic individual myocardial strain and strain rate profiles in different species.

Scaling of Long-axis Function

\( S_{\text{Long}} \) showed a small but significant increase with increasing LV EDV and LV mass, with power exponent of \( \beta_{\text{EDV}} 0.047\pm0.006 \), and \( \beta_{\text{LV mass}} 0.051\pm0.007 \), respectively (\( p<0.0001 \) vs. 0 for both).

As expected, absolute strain rates decreased with increased heart size, with systolic SR\(_{\text{Long}}\) scaling with exponent of \( \beta_{\text{EDV}} 0.216\pm0.019 \), and \( \beta_{\text{LV mass}} 0.236\pm0.020 \) (\( p<0.0001 \) vs. 0 for both), and diastolic SR\(_{\text{Long}}\) scaling with exponent of \( \beta_{\text{EDV}} 0.159\pm0.020 \), and \( \beta_{\text{LV mass}} 0.175\pm0.021 \) (\( p<0.0001 \) vs. 0 for both) (Figure 2).

DTI-derived strain rates also showed inverse relationships with the heart size. Systolic-TDI-SR\(_{\text{Long}}\) scaled to LV EDV and mass with exponent of \( \beta_{\text{EDV}} 0.252\pm0.021 \) and of \( \beta_{\text{LV mass}} 0.263\pm0.021 \) (\( p<0.0001 \) vs. 0 for both; \( p = \text{NS} \) when compared to corresponding exponents obtained for speckle tracking-derived
SR), while diastolic-TDI-SR$_{long}$ scaled with exponent of $\beta_{EDV} = -0.241 \pm 0.020$ and of $\beta_{LVmass} = -0.262 \pm 0.020$ ($p < 0.0001$ vs. 0 for both; $p < 0.01$ when compared to corresponding exponents obtained for speckle tracking-derived SR) (Figure 3).

Scaling of Short-axis Function

S$_{circ}$ was not affected with heart size ($\beta_{EDV} = 0.008 \pm 0.008$, and $\beta_{LVmass} = 0.008 \pm 0.009$, $p > 0.1$ vs. 0 for both). Again, systolic S$_{circ}$ decreased with increasing heart size with exponent of $\beta_{EDV} = -0.261 \pm 0.021$, and $\beta_{LVmass} = -0.277 \pm 0.021$ ($p < 0.0001$ vs. 0 for both). Similarly, diastolic S$_{circ}$ scaled with exponent of $\beta_{EDV} = -0.206 \pm 0.023$, and $\beta_{LVmass} = -0.228 \pm 0.023$ ($p < 0.0001$ vs. 0 for both) (Figure 4).

S$_{rad}$ scaled with exponent of $\beta_{EDV} = 0.078 \pm 0.012$, and $\beta_{LVmass} = 0.086 \pm 0.014$ ($p < 0.0001$ vs. 0 for both). Systolic S$_{rad}$ scaled with exponent of $\beta_{EDV} = -0.193 \pm 0.012$, and $\beta_{LVmass} = -0.203 \pm 0.013$ ($p < 0.0001$ vs. 0 for both). Diastolic S$_{rad}$ scaled with exponent of $\beta_{EDV} = -0.130 \pm 0.018$, and $\beta_{LVmass} = -0.142 \pm 0.020$ ($p < 0.0001$ vs. 0 for both) (Figure 5).

Relationship between Long- and Short-axis Function

As a result, the longitudinal-to-circumferential strain ratio showed significant scaling exponents ($\beta_{EDV} = 0.038 \pm 0.009$ and $\beta_{EDV} = 0.039 \pm 0.010$, $p = 0.0002$ for both when compared with 0) (Figure 6). These results indicate that long- and short-axis strain scale differently.
Relationship between Systolic and Diastolic Strain Rate

During data review, we noted that diastolic strain rates have consistently lower absolute values of power exponent. To elucidate this, we assessed the way the ratio between systolic and diastolic strain rates scales with LV EDV. All three ratios showed highly significant power exponent for ratio between systolic and diastolic SRlong = $0.07 \pm 0.02$, $p < 0.0001$; for ratio between systolic and diastolic SRcirc = $0.06 \pm 0.01$, $p < 0.0001$; for ratio between systolic and diastolic SRrad = $0.06 \pm 0.02$, $p < 0.0001$.

Intra-observer Variability

Intra-observer variability data are shown in Table 2.

Discussion

In this paper, we show that while $S_{circ}$ is unchanged, $S_{long}$ and $S_{rad}$ increases with increasing heart size. Furthermore, there was a significant decrease of the ratio between $S_{long}$ and $S_{circ}$ with decreasing heart size, indicative that contribution of long axis function decreases with decreasing animal size. We also show that all three components of diastolic and systolic strain rates increased with the decreasing heart size while following an allometric (power law) function, with the power exponents similar to the ones a priori proposed by us.

From a practical viewpoint, our findings are relevant for the accurate translation of research from bench to bedside. Additionally, they are important in the context of evolutionary biology, as we describe the way cardiac structures adopt to size constraints by changing the way they operate.

Strain and Heart Size

Myocardial deformation is a complex phenomenon that can best be represented as a $3 \times 3$ tensor, with diagonal elements representing three normal strains (analyzed in this paper), and off-diagonal elements representing shear strains. LV myocardial deformation during systole leads to expulsion of blood, with this process most often quantified by ejection fraction, a measure of global systolic function. As ejection fraction is constant between the species, it follows that myocardial strains should be largely invariant between the species. Yet Liu et al. showed that anesthetized mice have markedly longitudinal shortening (by 10%) and $S_{irc}$ lower, but by a lesser degree (by 6%), than conscious humans [5]. In contrast Bachner-Hinenzon et al. [27] showed that sedated rats have $S_{irc}$ that is 2% lower, but $S_{long}$ that is similar to humans; of note, this study reported markedly lower human $S_{long}$ than the one reported in the largest clinical multi-center $S_{long}$ study of healthy controls [28]. Our data show similar $S_{irc}$ between species but $S_{long}$ that was 5% lower in mice than in humans. A possible discrepancy between these studies may be due to differences in methods and/or the use of anesthetics/sedatives [21]. On the other hand, our findings of a small but significant increase of $S_{long}/S_{circ}$ ratio with increasing animal size are in accordance with the previously shown increase of the ratio between long- and short-axis displacement with animal size increase, [6] and is supported by findings of Liu et al. that shows

![Figure 4. Scaling of circumferential strain ($S_{circ}$) and corresponding peak systolic and early diastolic strain rates ($SR_{circ}$) to left ventricular end-diastolic volume and mass. Legend as for Figure 2. doi:10.1371/journal.pone.0040061.g004](image-url)
that body size has a stronger effect on long axis function. Finally, Decloedt et al. have shown that trotter horses [29,30] compared to humans reported in our study and previously [28] [27] have higher $S_{24.6\%}$ but similar $S_{19.7\%}$, again consistent with increase in $S_{2}/S_{1}$ ratio with increasing animal size.

### Strain Rates and Heart Size

We confirmed that strain rates scaled to the heart size with power exponents that were similar to the ones proposed in Appendix S1. The largest divergence from the predicted exponent was seen for diastolic and systolic $SR_{rad}$ which could represent a true physiologic phenomenon, or may be due to difficulties in accurately measuring radial strain rate. As a further support for our findings, Decloedt et al. have shown that trotter horses have mean values of all strain rate components lower by almost one order of magnitude when compared to the ones we obtained in mice [29,30].

It is interesting that for all three components of global strain rates, diastolic strain rates showed lower values of the power exponent. This would indicate that decreasing heart size leads to a larger increase in systolic, rather than diastolic strain rate. For example, in our data, the average systolic and diastolic $SR_{24.6\%}$ by speckle tracking were $1.0^{-\circ}$ and $1.5^{-\circ}$ in humans, but $6.5^{-\circ}$ and $4.3^{-\circ}$ in mice. This issue is even more complex given relatively shorter diastolic filling time in small mammals. [16] While the cause of this finding is unknown, it may be linked to between-species differences in gene expression. [31,32] Further studies are needed to elucidate if these findings carry evolutionary importance. [33].

### Limitations

Several confounding factors, such as differences in heart rate to frame rate ratio, impact of sedation, or intrinsic differences in heart geometry) could have influenced the results. However, recent report showed that speckle-tracking LV strain efficiently detects discrete alterations in mice despite being obtained from images acquired at the frame rate identical to one used in this study. [34] Also, mice, the smallest species studied, have been assessed without any sedation. Finally,LV geometry, quantified by the ratio between LV radius and length, while showing between-species variability, [5] appears to vary independently from the species size (see Table 2).

In this study LV mass of our human subjects was lower than reported in our previous studies, which decreased the range of LV mass studied. The reason for this was that 14 out of 20 of our subjects were women with relatively small body size. The average indexed LV mass was $61 \pm 12 \text{ gm/m}^2$, i.e. within normal limits.

Speckle-tracking derived strains obtained by method used in this study were validated only in humans and dogs, [35] which makes
accuracy of assessment of strain in small animals uncertain. However, we have shown that speckle-tracking derived strains derived by the current method are reflective of myocardial scar or fibrosis in rabbits, rats, and mice. [9,15,21] We are also showing here that high-frame tissue-Doppler derived strain rate of the mid-ventricular septum (which was validated in men [36] and in mice, [37] i.e. in species at the two extremes of body size) correlates well with the global speckle-tracking derived strain rate, adding to plausibility of our findings.

We have not assessed all strain components, and the significant impact of shear strains cannot be excluded. Also, strain rate measurements by speckle tracking echocardiography are less accurate than the strain measurements. Yet several authors showed that one can accurately measure strain rates in mice and rats, and that these measures correlate with relevant physiologic parameters. Furthermore, we tried to overcome this issue by averaging strain from multiple segments. Finally, the impact of sedation and the stress of performing echocardiography studies cannot be completely eliminated.

In conclusion, while $S_{circ}$ is unchanged, $S_{long}$ and $S_{rad}$ increase with increasing heart size, indicating that large mammals rely more on long axis contribution to systolic function. While both diastolic and systolic $SR_{long}$, $SR_{circ}$, and $SR_{rad}$ increase with decreasing heart size, the increase is more substantial for systolic $SR$. Further studies are needed to develop species normal ranges for subsequent intervention studies.

**Supporting Information**

Figure S1 Scaling of left ventricular (LV) length and systolic and diastolic mitral annulus (MA) long axis velocities to left ventricular mass derived from previous data [6] (Panels A and B); Predicted scaling of longitudinal systolic and diastolic strain rates ($SR_{long}$) (Panels C and D); Scaling of longitudinal and circumferential strains ($S_{long}$ and $S_{circ}$) derived from data from Liu et al. [5] (Panel E).

**Table 2.** Intra-inter observer variability of reconstructed strain data in the different species.

|                  | Longitudinal Strain (%) | Systolic Longitudinal Strain Rate (s^-1) | Diastolic Longitudinal Strain Rate (s^-1) |
|------------------|-------------------------|----------------------------------------|------------------------------------------|
| Mean             | 23.1±13.5               | 3.6±2.6                                | 3.4±1.8                                  |
| Intraobserver variability |
| Mean absolute difference | 1.84±1.27               | 0.45±0.29                              | 0.49±0.24                               |
| Mean relative difference (%) | 8.4±5.2                 | 12.5±7.4                               | 13.3±7.2                                |
| Correlation      | 0.99                    | 0.99                                   | 0.98                                     |

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Author Contributions
Conceived and designed the experiments: ZBP. Performed the experiments: MSP YZ YC ZBP. Analyzed the data: KK MSP YZ JDT THM

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Author/s:
Kusunose, K; Penn, MS; Zhang, Y; Cheng, Y; Thomas, JD; Marwick, TH; Popovic, ZB

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