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The Effect of Respiration-Driven Flow Waveforms on Hemodynamic Metrics Used in Fontan Surgical Planning

Elaine Tang, PhD^{1,‡}, Zhenglun Alan Wei, PhD², Phillip M. Trusty, MS², Kevin K. Whitehead, MD, PhD³, Lucia Mirabella, PhD^{2,‡}, Alessandro Veneziani, PhD⁴, Mark A. Fogel, MD³, and Ajit P. Yoganathan, PhD^{1,2,*}

¹School of Chemical and Biomolecular Engineering, Georgia Institute of Technology, Atlanta, GA

²Wallace H. Coulter Department of Biomedical Engineering, Georgia Institute of Technology & Emory University, Atlanta, GA

³Division of Cardiology, Children's Hospital of Philadelphia, Philadelphia, PA

⁴Department of Mathematics and Computer Science, Emory University, Atlanta, GA

Abstract

Objective: Poor total cavopulmonary connection (TCPC) hemodynamics have been hypothesized to be associated with long-term complications in Fontan patients. Image-based Fontan surgical planning has shown great potential as a clinical tool because it can pre-operatively evaluate patient-specific hemodynamics. Current surgical planning paradigms commonly utilize cardiac-gated phase contrast magnetic resonance (MR) imaging to acquire vessel flows. These acquisitions are often taken under breath-held (BH) conditions and ignore the effect of respiration on blood flow waveforms. This study investigates the effect of respiration-driven flow waveforms on patient-specific hemodynamics using real-time MR acquisitions.

Methods: Patient-specific TCPCs were reconstructed from cardiovascular MR images. Real-time phase contrast MR images were acquired under both free-breathing (FB) and breath-held conditions for 9 patients. Numerical simulations were employed to assess flow structures and hemodynamics used in Fontan surgical planning including hepatic flow distribution (HFD) and indexed power loss (iPL), which were then compared between FB and BH conditions.

Results: Differences in TCPC flow structures between FB and BH conditions were observed throughout the respiratory cycle. However, the average differences (BH – FB values for each patient, which are then averaged) in iPL and HFD between these conditions were 0.002 ± 0.011 (p

*Corresponding Author: Ajit P. Yoganathan, Wallace H. Coulter Department of Biomedical Engineering, Georgia Institute of Technology & Emory University, Technology Enterprise Park, Suite 200, 387 Technology Circle, Atlanta, GA 30313-2412, Tel: +1 404-894-2849, Fax: +1 404-385-1268, ajit.yoganathan@bme.gatech.edu.

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Conflict of Interest

No benefits in any form have been or will be received from a commercial party related directly or indirectly to the subject of this manuscript.

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= 0.40) and $1\pm 3\%$ ($p = 0.28$), respectively, indicating no significant difference in clinically important hemodynamic metrics.

Conclusions: Respiration affects blood flow waveforms and flow structures, but might not significantly influence the values of iPL or HFD. Therefore, breath-held MR acquisition can be adequate for Fontan surgical planning when focusing on iPL and HFD.

1 Introduction

Fontan patients suffer from various long-term complications which are hypothesized to be attributed to an adverse hemodynamic environment in the total cavopulmonary connection (TCPC). Elevated TCPC power loss (PL) under exercise conditions has been associated with diminished patient exercise performance (Khiabani et al., 2015; Tang et al., 2017; Whitehead et al., 2007). Unbalanced hepatic blood flow distribution (HFD) between the two lungs has been associated with increased risk of pulmonary arteriovenous malformations (PAVM) in Fontan patients (Pike et al., 2004; Sundareswaran et al., 2009). These two TCPC hemodynamic metrics are often involved in image-based Fontan surgical planning, which is progressively making its way from a pure research tool to clinical use to improve Fontan surgical outcomes (Corsini et al., 2015; de Zélicourt and Kurtcuoglu, 2016; Trusty et al., 2018; Wei et al., 2017). Historically, the majority of surgical planning cases utilized blood flow waveforms segmented from magnetic resonance images (MRI) acquired under breath-held (BH) conditions. The BH acquisition is preferable because it reduces image artifacts due to respiratory motion (Wei et al., 2016) but it ignores the impact of respiration on the flow waveforms.

Recent clinical studies have demonstrated that passive flow in the TCPC is affected by respiration in single ventricle patients. Fogel et al. has shown that flow in the systemic venous pathway of Fontan patients is phasic to both cardiac and respiratory cycles; approximately 70% of the flow is cardiac-dependent, while the remainder depends on respiratory influence (Fogel et al., 1997). Hsia et al. reported that hepatic venous flow (measured with Doppler ultrasonography) depends on respiration in Fontan subjects more so than in normal subjects and is larger during inspiration than expiration (Hsia et al., 2001; Hsia et al., 2000). Hjortdal et al. also observed increased inferior vena cava (IVC) flow during inspiration in Fontan patients (Hjortdal et al., 2003). During inspiration, the negative intra-thoracic pressure assists the forward flow in the vena cavae and the pulmonary arteries (PAs), increasing venous return as compared to expiration. Recently, Wei et al. showed increased pulsatility in systemic return flows during free-breathing (FB) conditions compared to the BH setting. (Wei et al., 2016). These respiratory flow effects may have important implications in the evaluation of TCPC hemodynamics and require further investigation.

Recognizing this shortcoming, Marsden et al. simulated TCPC hemodynamics by including a respiratory influence on the IVC and SVC flow waveforms (Marsden et al., 2007). This was done by superimposing the patient-specific flow waveform of a cardiac cycle with the flow waveform of a respiratory cycle from Hjortdal et al. (Hjortdal et al., 2003). When compared with a steady flow model, their respiration model showed reduced energetic

efficiency. While this study effectively elucidates the effect of respiration on TCPC energetics, it is not clear whether the respiratory influence is patient-specific or statistically significant for a larger population and other clinically important hemodynamic metrics. Thus, further investigation using patient-specific respiratory flow waveforms is necessary. To bridge this knowledge gap, the current study utilizes real-time phase contrast MRI (PC-MRI) patient-specific flow waveforms acquired under BH and FB conditions, which serve as boundary conditions for numerical simulations. These simulations were conducted to assess TCPC flow structure and clinically important hemodynamic metrics that are commonly involved in Fontan surgical planning including PL and HFD (de Zélicourt and Kurtcuoglu, 2016; Marsden, 2013; Wei et al., 2017). These metrics were compared between BH and FB conditions to demonstrate the impact of respiratory flow on assessing TCPC hemodynamics in Fontan surgical planning.

2 Materials and Methods

2.1 Patient Cohort

Single ventricle patients with a TCPC anatomy were selected from the Georgia Tech/Children's Hospital of Philadelphia Fontan database. With informed consent and Institutional Review Board approval, prospective cardiovascular magnetic resonance (CMR) data at resting BH and FB conditions were acquired. Patients included in this study had: (i) a TCPC completion, and (ii) real time PC-MRI acquired at all TCPC inlets. Patients were excluded if: (i) a pacemaker or implanted metal was present, or (ii) severe CMR image artifacts distorted the images. A total of nine consecutive patients were included after two were excluded due to severe image artifacts.

2.2 CMR Acquisition

Anatomic CMR and real time PC-MRI acquisitions were performed on each patient. Static steady-state free precession imaging was acquired, and 3D anatomies were reconstructed (Frakes et al., 2003; Frakes et al., 2005) (figure illustrated in Supplementary Material 1). An echoplanar (EPI) sequence utilizing shared velocity encoding was applied to acquire real time PC-MR images (Lin et al., 2012) across the Fontan pathway (FP) at the IVC superior to the hepatic vein confluence and the superior vena cavae (SVC) as separate velocity maps for at least 10 seconds (approximately 20 frames per second). The same imaging protocol was performed under BH (acquired at end expiration) and FB resting conditions. The velocity of the diaphragm under BH conditions was more constant at end expiration than at end inspiration (Holland et al., 1998). Therefore, it was suggested to acquire MRI under end expiration (Fratz et al., 2013). This sequence has been validated in phantom models and shown to be reliable for low-velocity flows (Lin et al., 2012). Conventional PC-MR images were also acquired at the left (LPA) and right (RPA) pulmonary arteries during resting BH conditions at the end of expiration. The patient demographic and image acquisition parameters are summarized in Table 1.

2.3 Selection of Respiratory Cycle

Patient-specific flow conditions were obtained by segmenting real time PC-MR images at the cross section of each vessel. The segmentation was done by using a freely available

scientific package Mediviso Segment (<http://medviso.com/products/segment/>) (Heiberg et al., 2010). The details of the real time PC-MRI processing are described in Wei *et al.* (Wei et al., 2016) The change of chest wall area, tracked in the same images of vessel flows, assists in identifying the respiratory cycle and its relation with the vessel flow waveforms. It is not trivial to determine the cardiac cycle because real time PC-MRI acquisition does not contain any electrocardiograms (ECG). Therefore, this study segmented the ascending aorta and/or descending aorta that present on the same images as the SVC and FP, respectively. The aortic flow waveform is then used to define the cardiac cycle. An example of the segmented waveforms is illustrated in Figure 1. For each inlet vessel (FP and SVC) in all data sets, respiratory cycles were identified in FB conditions. An approximately the same duration with the same number of cardiac cycles under BH conditions was selected as a surrogate “respiratory cycle” under BH conditions for numerical modeling and comparison purposes. FB and BH flow waveforms of the patients included in this study were illustrated in Supplementary Material 2.

2.4 Hemodynamic Comparison

Computational fluid dynamics (CFD) was used to simulate patient specific hemodynamics. The FP and SVC waveforms were utilized as inlet boundary conditions in the CFD. The outlet flow waveforms (LPA and RPA) under FB condition were not available (due to the more notable vessel motion of the PAs during FB condition). Therefore, resistance boundary conditions based on flows obtained from conventional BH MRI were prescribed at the outlets (Peiró and Veneziani, 2009; Quarteroni et al., 2016). A similar setting was validated by an *in vitro* experiment (Tree et al., 2018). The finite element solver LifeV (www.lifev.org) was used (Mirabella et al., 2012). A three-element Windkessel (resistor, capacitor, and resistor) model was employed to prescribe lung resistances at the end of the LPA and RPA (Alastruey et al., 2008). Simulations were performed for each patient under pulsatile FB and BH conditions.

For all simulations, PL was computed at each time step using the following equation:

$$PL = \sum_{inlet} \left(p + \frac{1}{2} \rho v^2 \right) Q - \sum_{outlet} \left(p + \frac{1}{2} \rho v^2 \right) Q \quad (1)$$

Indexed power loss (iPL), which represents the non-dimensional power loss through the TCPC (Dasi et al., 2009; Wei et al., 2018), was computed by normalizing time-averaged PL by flow and patient body surface area (BSA) ($iPL = PL / (\rho Q_s^3 / BSA^2)$), where Q_s is the total systemic venous return and ρ is blood density = 1060 kg m⁻³. Pulsatile PL and iPL were computed using the pulsatile FB and BH waveforms. Non-pulsatile PL and iPL were computed using time-averaged FB and BH waveforms as boundary conditions for the CFD simulations.

Particle tracking was performed with PARAVIEW software (Kitware Inc., Clifton Park, NY, USA) to visualize the flow field. HFD was computed by seeding particles at the FP and counting the number of particles leaving through the LPA and RPA:

$$\%HFD(LPA) = \frac{n_{FP-LPA}}{n_{FP-LPA} + n_{FP-RPA}} \times 100\% \quad (2)$$

where %HFD(LPA) is the percentage of particles seeded at the FP leaving through the LPA, and n_{FP-LPA} and n_{FP-RPA} are the total number of particles exiting through the LPA and RPA, respectively. The difference in %HFD (LPA) between BH and FB conditions was then defined as %HFD = %HFD_{BH}-%HFD_{FB}. Details of the numerical simulations, Windkessel model parameters and particle tracking were described in Supplementary Material 3.

2.5 Statistical Analysis

Statistical comparisons were conducted to identify differences between FB and BH conditions. Paired sample t-tests were utilized for normally distributed data, while Wilcoxon signed ranks test was used for non-normal distributions. For all statistical analysis, a p-value 0.05 is considered significant.

3 Results

3.1 Flow Field Comparison

To better understand the hemodynamic differences between the TCPC flow fields under FB and BH conditions, velocity fields across a coronal plane within the TCPC were compared for each patient. Particle tracking videos were also created to visualize the instantaneous flow fields (see Supplementary Material 4-7). In general, larger variations in the TCPC flow field (velocity magnitude) within the duration of a respiratory cycle were observed qualitatively under FB conditions compared to BH conditions. In all patients, the maximum velocity magnitude was higher during inspiration under FB conditions than during the corresponding time point under BH conditions. During expiration, retrograde flow (as observed from the PC-MRI data and prescribed in the flow boundary conditions) in the FP and SVC was observed in some patients under FB conditions, while no retrograde flow was seen in the FP under BH conditions for any patient. For most patients, the FB and BH flow fields were most similar at end expiration. The following sections illustrate the flow fields for two patients as representative cases. Note that in the flow field figures, the same velocity contour scale was used for a given patient throughout the four phases (mid-inspiration, end inspiration, mid-expiration, and end expiration).

The velocity field in the FP of Patient A throughout different phases of the respiratory cycle is visualized in Figure 2. During mid-inspiration (Figure 2(a)), forward flow was noted in both conditions, while higher peak velocity magnitudes were found under the FB condition. During end-inspiration (Figure 2(b)), maximum flow and maximum velocity magnitudes were observed in the FP under the FB condition. During mid-expiration, retrograde flow existed in the FP under the FB condition, resulting in different flow fields between the FB and BH conditions (Figure 2(c)). Finally, similar velocity magnitudes and directions were observed in both the FB and BH conditions at end expiration (Figure 2(d)).

The TCPC velocity field for Patient I throughout different phases of the respiratory cycle is visualized in Figure 3(a-d). The maximum velocity magnitude of the FB condition under inspiration was higher than that of the BH condition. From the particle tracking videos of Patient I (Supplemental Materials 4 (FB) and 5 (BH)), different flow recirculation patterns were observed between the two conditions. During the FB condition, flow recirculation occurred at the inferior part of the FP. This recirculation could have been caused by higher SVC flow compared to FP flow around $t = 3.84\text{--}3.97$ s (as demonstrated in the flow waveforms in Figure 3). Figure 4 confirmed the cause of this flow recirculation by showing that flow particles seeded at the SVC flushed the FP. Under the BH condition however the recirculation caused by the SVC irrigation to the FP was not observed, while flow recirculation was observed at the center of the TCPC. The particle tracking videos of Patient I are shown in Supplemental Materials 6 (FB) and 7 (BH)).

3.2 Hepatic Flow Distribution

Global pulmonary flow distribution (PFD) and HFD under FB and BH conditions were computed and summarized in Table 2. Comparing the two conditions, the average difference observed in time-averaged HFD were 1%, with a range of -3% to 7%. The preference of hepatic flow to the left/right lungs is not statistically different ($p = 0.28$) between the FB and BH conditions. This was also reflected by the Bland–Altman plot of HFD shown in Figure 5. The “2SD” lines in indicate the 95% limits of agreement.

3.3 TCPC Power Loss

Table 3 and Table 4 tabulate PL and iPL respectively under FB and BH conditions. For each patient, power loss was the highest under the FB (pulsatile) condition. Power loss for the other conditions was found in the following descending order for all patients: FB (pulsatile), FB (non-pulsatile), BH (pulsatile), and BH (non-pulsatile). In all patients and at all conditions, pulsatile TCPC power loss was higher than non-pulsatile flow TCPC power loss. When comparing between FB and BH pulsatile iPL, the average difference in iPL was 0.002 ± 0.011 . There was no significant difference in iPL between the two conditions ($p = 0.40$). Figure 6 shows the Bland-Altman plots of PL and iPL, reinforcing the insignificant differences between FB and BH conditions. Since the differences of PL and iPL were not normally distributed, corresponding log scales were utilized to form the Bland-Altman plots.

4 Discussion

Image-based surgical planning of the Fontan procedure has emerged in the past few decades to provide surgeons additional insights to the possible hemodynamic outcomes of a given surgical option. Patient specific hemodynamics, power loss, and hepatic flow distribution are common metrics used to evaluate the performance of patients' current Fontan connection (Trusty et al., 2016) or that of potential surgical options (Sundareswaran et al., 2009). However, these metrics are often evaluated using flow waveforms acquired under breath-held conditions. Wei *et al.* concluded that respiration significantly affects the properties of TCPC flow waveforms, e.g. pulsilities, but not time-averaged venous flow rates (Wei et al., 2016). The effect of respiration on flow waveforms has the potential to influence the simulated TCPC hemodynamic metrics (Khiabani et al., 2012; Wei et al., 2017), which are

important to Fontan surgical planning. This current work serves as an extension of Wei *et al.* (Wei et al., 2016) to investigate how the respiratory flow waveform influences simulated TCPC hemodynamics.

In this study, the patient-specific effect of respiration on flow waveforms was evaluated with the use of real time PC-MRI data acquired under FB and BH conditions. The flow waveforms under BH conditions were driven by cardiac pulsation, while those under FB conditions were driven by both cardiac pulsation and respiration. Therefore, comparison between FB and BH conditions could isolate the effect of respiration-driven flow waveform on TCPC hemodynamics. CFD simulations utilizing patient-specific flow waveforms under FB and BH conditions demonstrated differences in the TCPC flow field between these conditions. As observed in the velocity fields of the CFD results, during mid-inspiration and end inspiration under the FB condition, higher flow rates through the TCPC resulted in increased peak velocity magnitude within the TCPC. This finding agrees with previous studies that showed increased IVC flow during inspiration in Fontan patients (Hjortdal et al., 2003). During inspiration, the negative intra-thoracic pressure in the chest cavity assists the forward flow in the TCPC, increasing venous return. During mid-expiration and end expiration, the velocity magnitude and direction of velocity vectors show more similarity between the two conditions. This resemblance is expected since the BH data were acquired during end-expiration, and the caval flow rates were of similar magnitude in the two conditions.

In addition to comparisons of instantaneous flow fields, the impact of respiratory flow was also explored for clinically important hemodynamic metrics including HFD and iPL. Statistical analysis demonstrates no significant difference in neither HFD nor iPL between FB and BH conditions. Moreover, Bland-Altman plots conclude the agreement between FB and BH conditions in assessing TCPC hemodynamic metrics.

Even though instantaneous flow structures are different between FB and BH conditions, the time-averaged hemodynamic metrics used in Fontan surgical planning, HFD and iPL, are not statistically significantly different between the two conditions. The major flow structure differences between FB and BH conditions are observed during mid-inspiration, which FB condition have higher velocity magnitudes; however the velocity directions are in general consistent between the two conditions during mid-inspiration. In addition, the presence of retrograde flow in the vena cava may differ between the FB and BH conditions during expiration (e.g. patient A). However, these differences did not have a major impact on the resulting time-averaged HFD and iPL. In most patients, the flow mixing patterns between FP and SVC were similar between FB and BH conditions. Different FP-SVC flow mixing patterns were observed in Patient I, but the resulting HFD only differs by 7% and iPL only differs by 0.0002 between the two conditions. This suggests the difference in FB and BH flow waveforms contributes to differences in TCPC flow structures and also instantaneous HFD and iPL values, but has only a minor impact on the resulting time-averaged HFD and iPL.

Time-averaged HFD and iPL are important clinical metrics evaluated in Fontan surgical planning. Impact of pulsatile flow waveforms in Fontan patients has been investigated

previously within a cardiac cycle (Dur et al., 2010; Dur et al., 2012), but not for respiratory cycles. Even though differences in iPL were observed between pulsatile and non-pulsatile conditions for both FB and BH conditions, there were no significant difference between pulsatile iPL between FB and BH conditions. The findings from the current study suggest that, in general, the routine use of BH conditions in Fontan surgical planning is acceptable. This is important, not only because fewer image artifacts exist in MRI under BH conditions (Wei et al., 2016), but also because numerical simulations with BH flow waveforms require less computational time (Wei et al., 2017). This is particularly important in image-based surgical planning of Fontan surgery, when simulations of multiple surgical options have to be completed in a short time frame.

5 Limitations

The major limitations of this study are due to the assumptions made in the computational model. The first limitation refers to the outlet boundary conditions. We approximated the downstream circulation with a Windkessel model, and the parameters were empirically tuned to the available patient-specific data. Even though this approach gives a reliable representation of the effects of the peripheral vasculature, the assumptions made may introduce some inaccuracies. Second, this study focuses on investigating the effect of respiratory flow on evaluating TCPC hemodynamics. In order to isolate this effect, all CFD simulations adopt rigid vessel wall assumptions. A previous study suggested that wall compliance has little impact on time-averaged HFD and energy efficiency (Long et al., 2012). Future study should investigate whether this holds under the influence of change in intrathoracic pressure under FB condition. Third, the respiratory cycle in this work was identified by tracing the change in chest wall area from a transverse CMR image. Note that chest wall motion, in reality, is 3-dimensional. iPL was used as a major hemodynamic metric, given its relevance to Fontan surgery planning. However, this metric was originally formulated for time-averaged flow. Future work shall address the pulsatile contribution of power loss to iPL. In addition, a uniform spatial distribution was used to seed particles at the FP plane for HFD quantification. This method does not account for the spatial velocity profile at the FP and therefore may be less accurate than using an appropriately distributed particle flux. Nevertheless, all these limitations should not affect the comparison of FB and BH conditions significantly. Finally, while the current study only included nine patients (mainly intra-atrial and adult patients) due to data availability, it serves as the first study to investigate the effect of BH MRI acquisition on TCPC hemodynamic metrics using patient-specific FB waveforms acquired via real-time MRI. Future work should extend this investigation to include extra-cardiac and also younger patients. It would also be worthwhile to explore the potential difference between intra-atrial and extra-cardiac patients and how respiration effects would be different in the two templates.

6 Conclusions

This study demonstrated the effects of respiration-driven flow waveforms in the evaluation of TCPC hemodynamics. The use of BH acquired flow waveforms appears to have negligible effects on both HFD and iPL, but does affect the TCPC's instantaneous flow fields. These findings support that the routine use of BH MRI acquisition to obtain flow

waveforms for Fontan surgical planning is acceptable when evaluating HFD and iPL. Therefore, BH MRI acquisition may continue to serve as the favorable scanning protocol with the advantage of minimizing image artifacts due to respiratory motion while still offering accurate surgical planning metrics.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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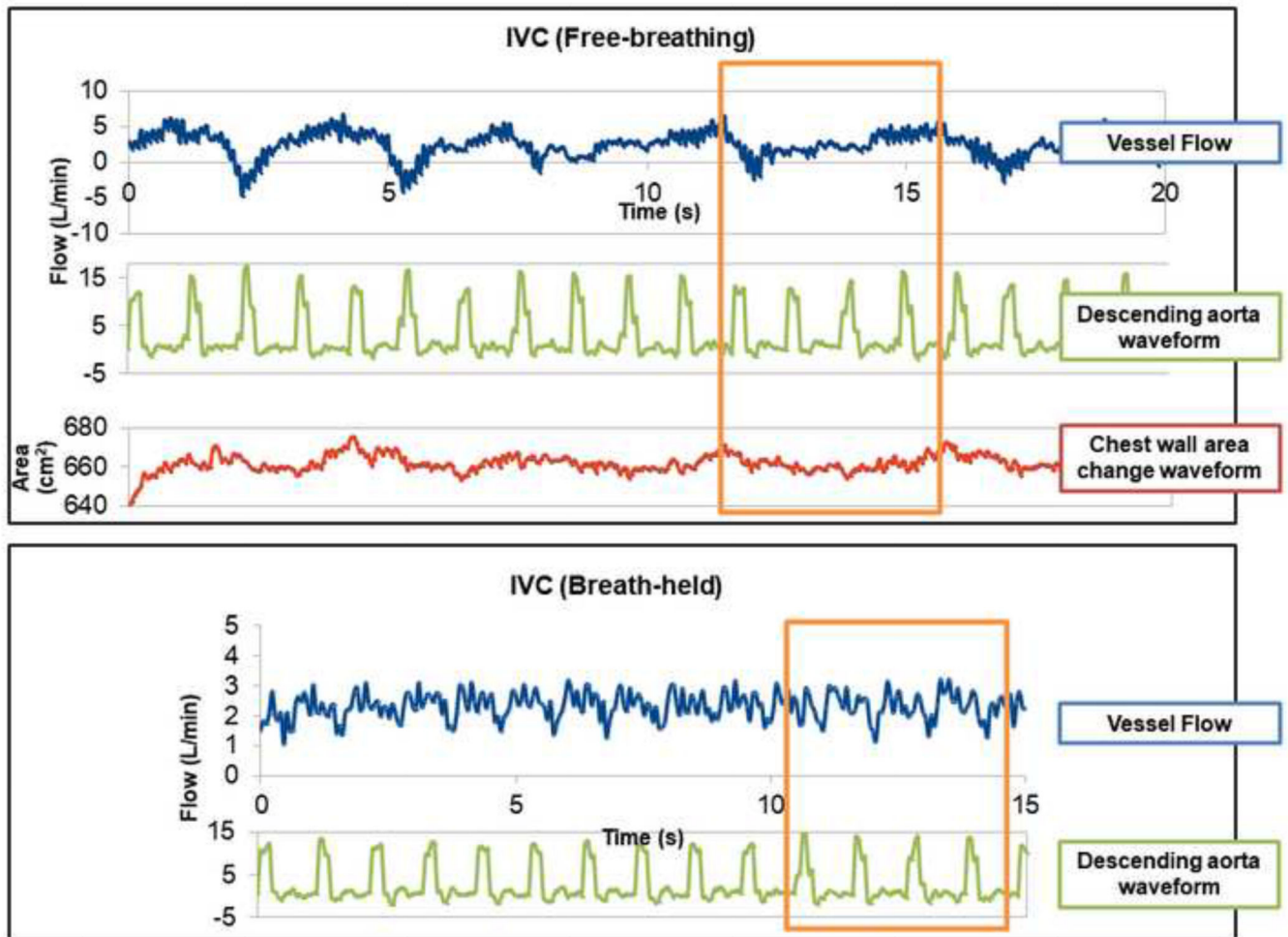


Figure 1. Example of segmented waveforms for vessel flow, descending aorta flow, and chest wall area of Patient A under FB and BH conditions. The selected cycle for simulation is highlighted by the rectangle.

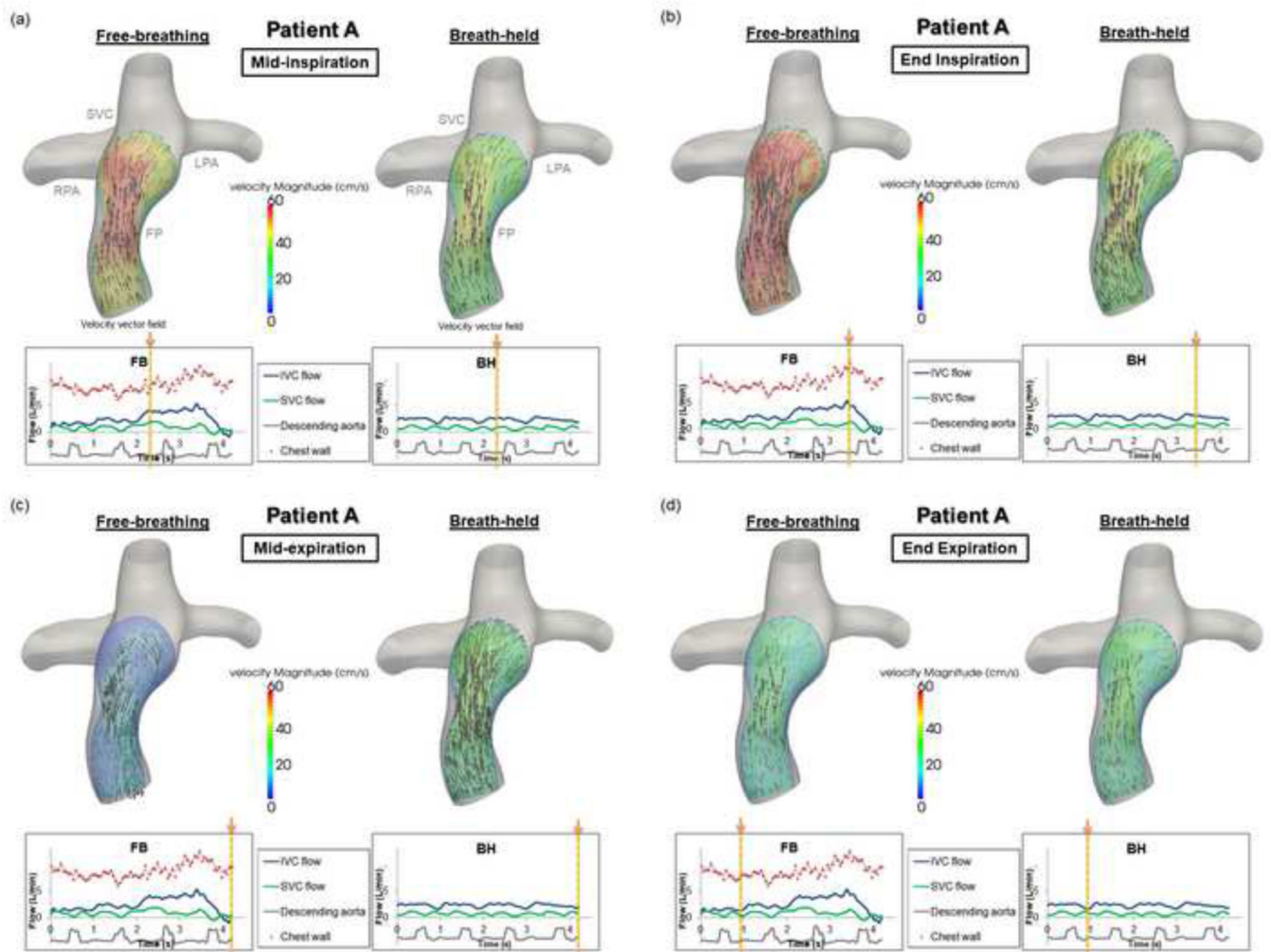


Figure 2. Fontan pathway flow fields under free-breathing and breath-held conditions for Patient A throughout different phases of the respiratory cycle: (a) mid-inspiration, (b) end inspiration, (c) mid-expiration, (d) end expiration. (Velocity magnitude unit: cm/s)

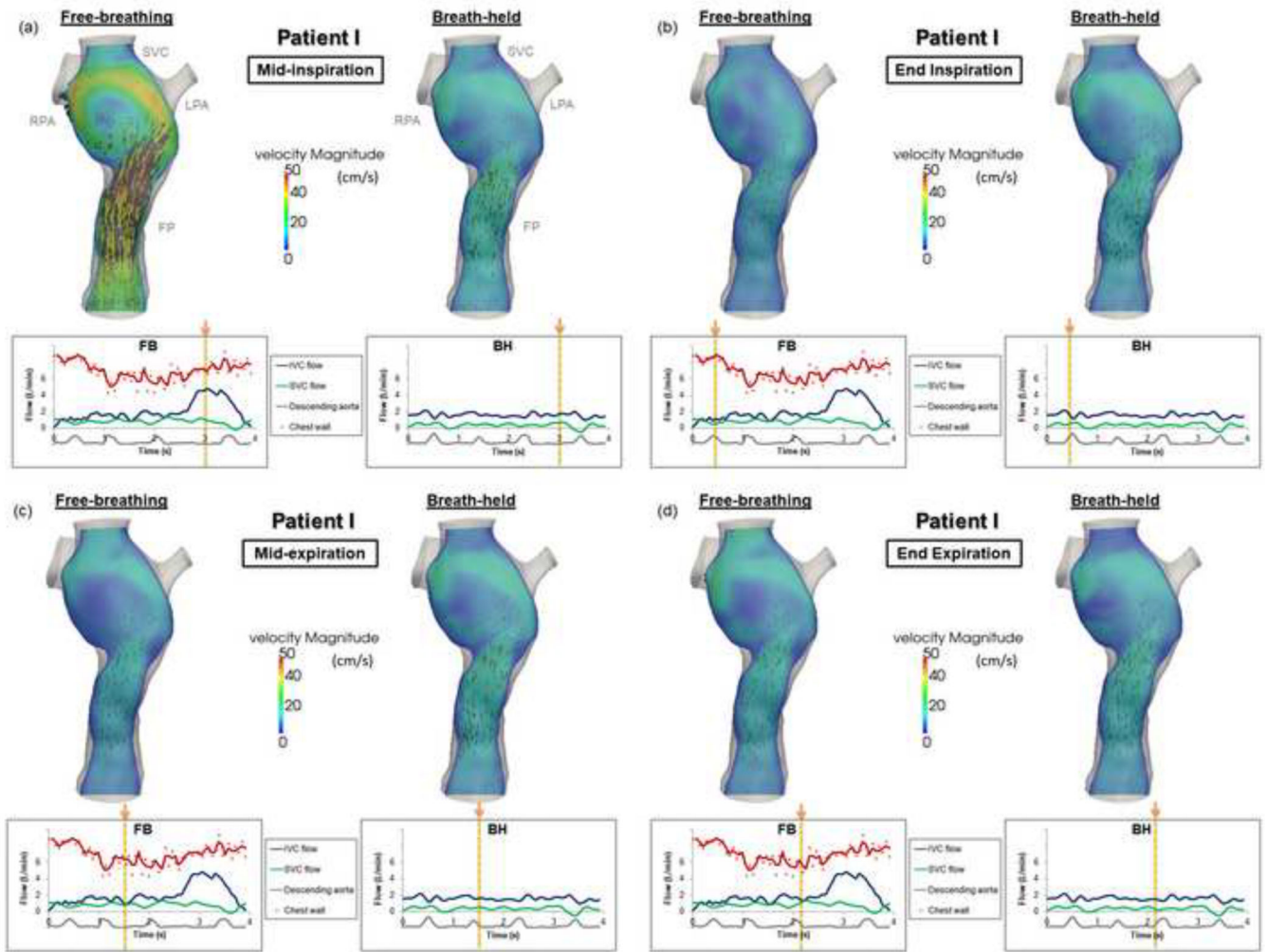


Figure 3. TCPC flow fields under free-breathing and breath-held conditions for Patient I throughout different phases of the respiratory cycle: (a) mid-inspiration, (b) end inspiration, (c) mid-expiration, (d) end expiration. (Velocity magnitude unit: cm/s).

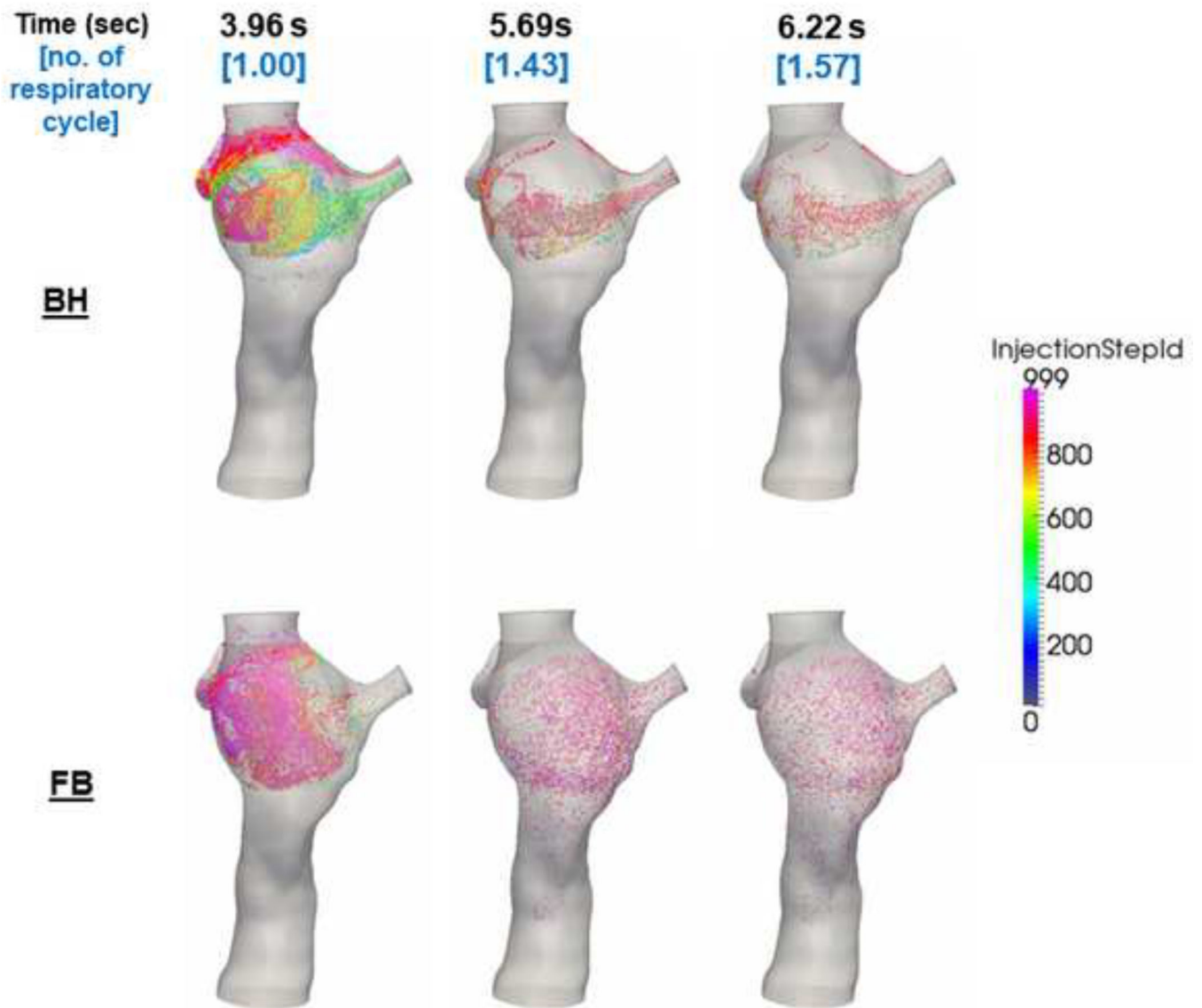


Figure 4. Screenshots of the particle tracking videos of Patient I with particles seeded at the SVC under FB and BH conditions. Higher Injection Step ID corresponds to particles released later in time.

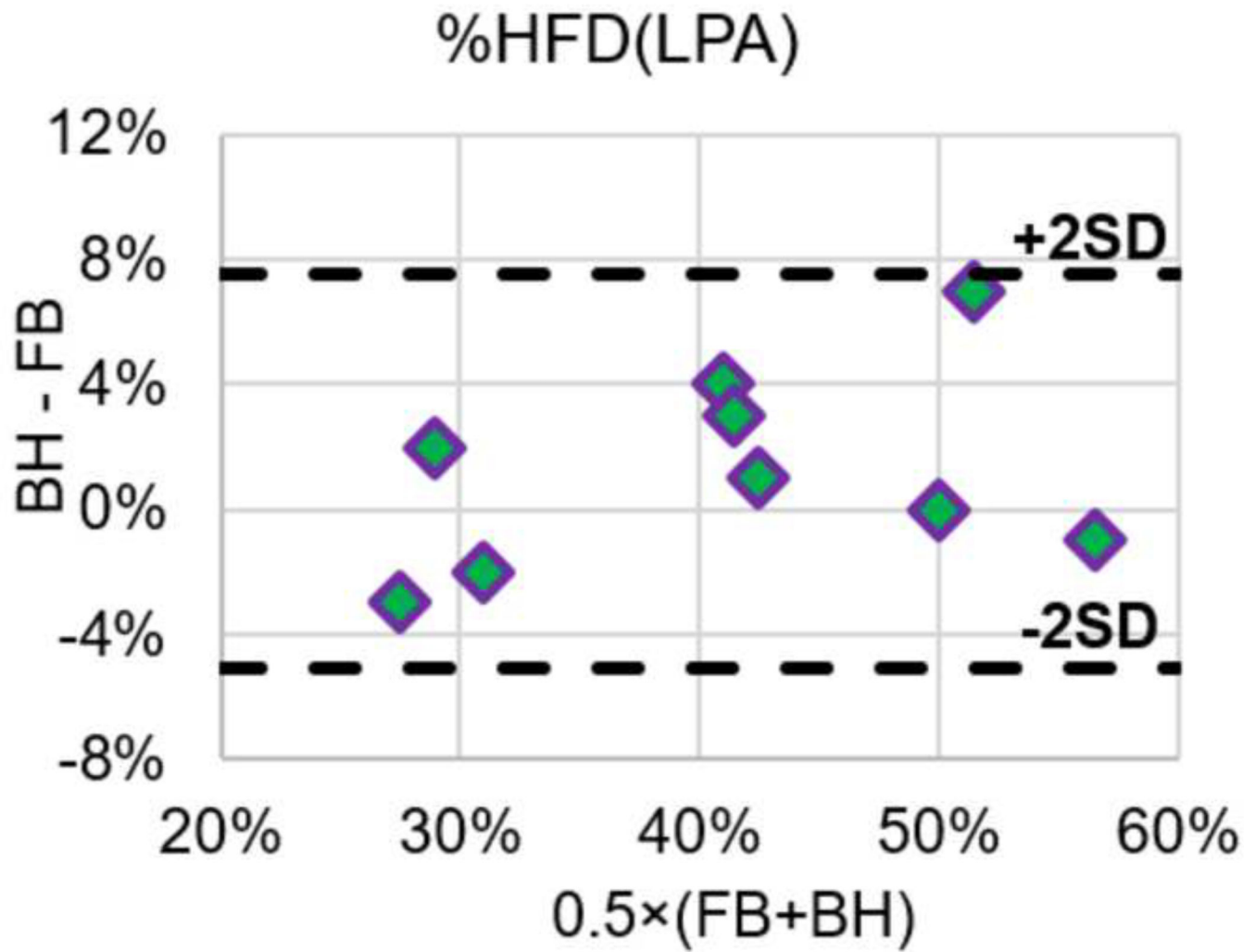
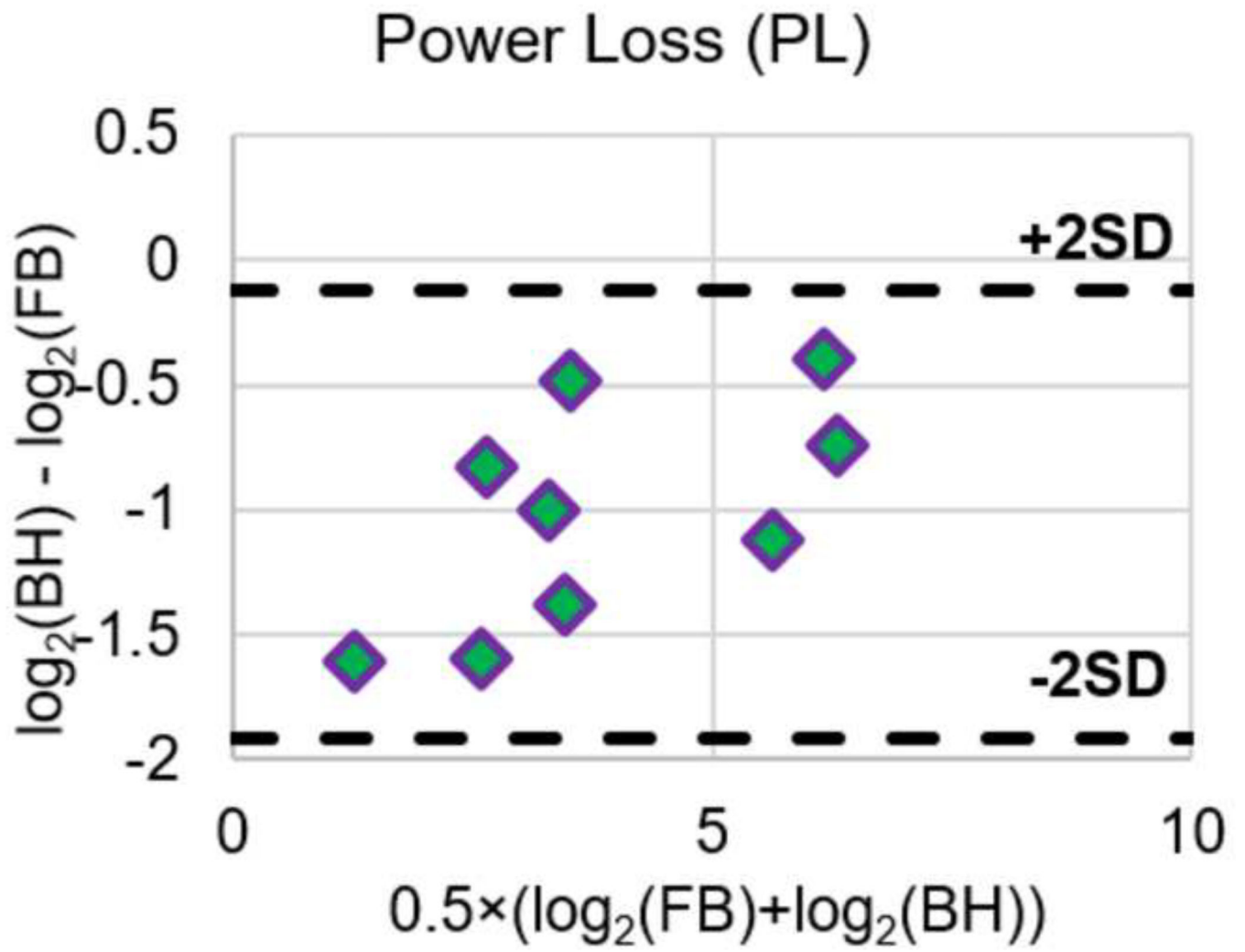


Figure 5.
Bland-Altman comparison of %HFD(LPA) under FB and BH conditions



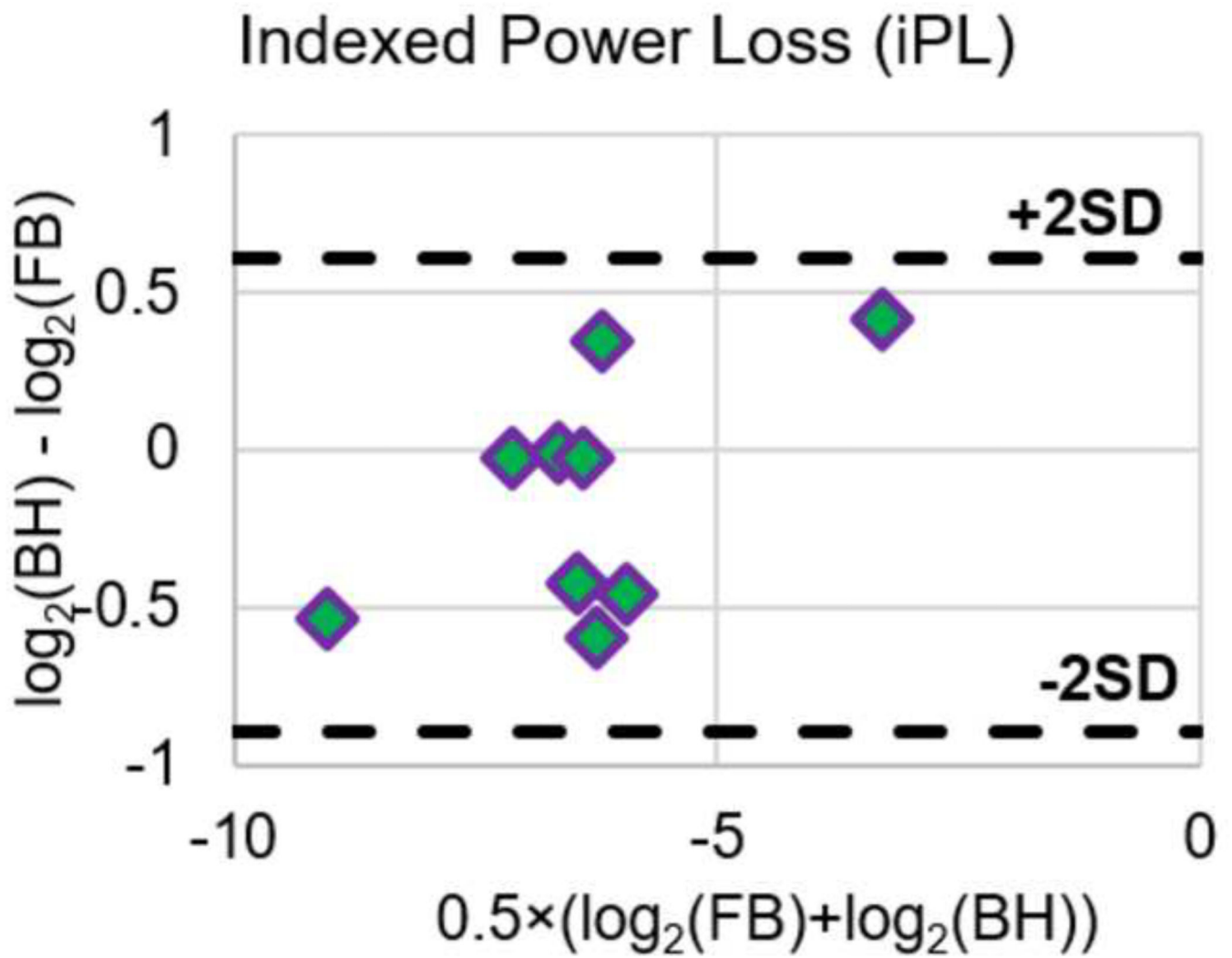


Figure 6.
Bland-Altman plots of PL and iPL under FB and BH conditions

Table 1.

Demographic details of the patients analyzed and summary of CMR acquisition parameters

Patient characteristics	Range
Age (years)	17–28
Body surface area (m ²)	1.64–1.92
Gender (Male/Female)	6/3
IVC connection type (IA/EC) *	8/1
Ventricular morphology (LV/RV/MV) †	3/5/1
<u>Transverse static steady state free precession CMR</u>	
Number of slices	45 – 70
Matrix (pixel)	256 X168
Spatial resolution (mm)	1.17 – 1.48
Slice thickness (mm)	3 – 4
Echo time (ms)	1.21 – 1.27
<u>Real time PC-MRI at FP and SVC</u>	
Encoding velocity (cm/s)	60 – 150
Number of images	199 – 511
Temporal resolution (ms)	32 – 77
<u>Conventional PC-MRI at LPA and RPA</u>	
Encoding velocity (cm/s)	60 – 100
Number of images	24 – 30

* IA – Intra-atrial, EC- Extra-cardiac.

† LV = left ventricle, RV = right ventricle, MV = mixed ventricle

Table 2.

Hepatic flow distribution (HFD) under FB and BH conditions

Patient	Age (years)	Body Surface Area (m ²)	Fontan type*	%PFD(LPA)	%HFD(LPA)	
					FB	BH
A	21	1.64	IA	46%	42%	43%
B	19	1.74	IA	27%	39%	43%
C	28	1.64	IA	25%	29%	26%
D	26	1.82	IA	38%	32%	30%
E	27	1.92	IA	53%	50%	50%
F	17.2	1.83	IA	52%	57%	56%
G	22	1.92	IA	49%	40%	43%
H	17	1.8	EC	41%	28%	30%
I	19	1.77	IA	31%	48%	55%
Average ± Standard Deviation		21.8±4.2	1.79±0.10	40±11%	40±10%	42±11%

* IA – Intra-atrial, EC- Extra-cardiac.

Table 3.

TCPC power loss PL under FB and BH conditions

Patient	Age (years)	Body Surface Area (m ²)	Fontan type *	Pulsatile PL (mW)		Non-pulsatile PL (mW)	
				FB	BH	FB	BH
A	21	1.64	IA	8.43	4.75	5.31	4.62
B	19	1.74	IA	13.63	9.77	11.84	9.05
C	28	1.64	IA	72.61	33.37	57.75	32.38
D	26	1.82	IA	10.43	3.45	6.28	3.1
E	27	1.92	IA	14.01	6.99	10.58	6.02
F	17.2	1.83	IA	102.8	61.24	100.79	60.5
G	22	1.92	IA	17.75	6.8	11.42	6.29
H	17	1.8	EC	81.86	62.11	63	60.2
I	19	1.77	IA	4.25	1.39	2.9	1.33
Average ± Standard Deviation	21.8±4.2	1.79±0.10		36.19±38.15	21.10±24.86	29.99±35.05	20.39±24.45

* IA – Intra-atrial, EC- Extra-cardiac.

Table 4.

TCPC power loss iPL under FB and BH conditions

Patient	Age (years)	Body Surface Area (m ²)	Fontan type *	Pulsatile iPL		Non-pulsatile iPL	
				FB	BH	FB	BH
A	21	1.64	IA	0.0163	0.0108	0.0103	0.0105
B	19	1.74	IA	0.0073	0.0071	0.0063	0.0066
C	28	1.64	IA	0.0885	0.1181	0.0704	0.1146
D	26	1.82	IA	0.0023	0.0016	0.0014	0.0014
E	27	1.92	IA	0.0133	0.0099	0.0101	0.0085
F	17.2	1.83	IA	0.0121	0.0154	0.0119	0.0152
G	22	1.92	IA	0.0193	0.0141	0.0124	0.0130
H	17	1.8	EC	0.0101	0.0101	0.0078	0.0098
I	19	1.77	IA	0.0121	0.0119	0.0083	0.0114
Average ± Standard Deviation	21.8±4.2	1.79±0.10		0.0201±0.0261	0.0212±0.0362	0.0154±0.0209	0.0221±0.0352

* IA – Intra-atrial, EC- Extra-cardiac.