Fontan hemodynamics from 100 patient-specific cardiac magnetic resonance studies: A computational fluid dynamics analysis

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Objectives: This study sought to quantify average hemodynamic metrics of the Fontan connection as reference for future investigations, compare connection types (intra-atrial vs extracardiac), and identify functional correlations using computational fluid dynamics in a large patient-specific cohort. Fontan hemodynamics, particularly power losses, are hypothesized to vary considerably among patients with a single ventricle and adversely affect systemic hemodynamics and ventricular function if suboptimal.

Methods: Fontan connection models were created from cardiac magnetic resonance scans for 100 patients. Phase velocity cardiac magnetic resonance in the aorta, vena cavae, and pulmonary arteries was used to prescribe patient-specific time-averaged flow boundary conditions for computational fluid dynamics with a customized, validated solver. Comparison with 4-dimensional cardiac magnetic resonance velocity data from selected patients was used to provide additional verification of simulations. Indexed Fontan power loss, connection resistance, and hepatic flow distribution were quantified and correlated with systemic patient characteristics.

Results: Indexed power loss varied by 2 orders of magnitude, whereas, on average, Fontan resistance was 15% to 20% of published values of pulmonary vascular resistance in single ventricles. A significant inverse relationship was observed between indexed power loss and both systemic venous flow and cardiac index. Comparison by connection type showed no differences between intra-atrial and extracardiac connections. Instead, the least efficient connections revealed adverse consequences from localized Fontan pathway stenosis.

Conclusions: Fontan power loss varies from patient to patient, and elevated levels are correlated with lower systemic flow and cardiac index. Fontan connection type does not influence hemodynamic efficiency, but an undersized or stenosed Fontan pathway or pulmonary arteries can be highly dissipative. (J Thorac Cardiovasc Surg 2014;148:1481-9)
We hypothesize that TCPC power loss varies considerably among connection types, such as extracardiac and intra-atrial Fontans (eg, connection geometry), and high losses may adversely affect systemic hemodynamics and ventricular function.

**MATERIALS AND METHODS**

This analysis was approved by the institutional review boards of the Children’s Hospital of Philadelphia and Georgia Tech. Informed consent was obtained.

**Patients**

A total of 114 consecutive patients with a completed Fontan connection who underwent CMR between 2002 and 2012 were identified. Patients were excluded on the basis of severe CMR artifacts, lack of sufficient phase contrast data, or diagnosis of Ebstein’s anomaly, reducing the number of investigated patients to 100. Demographic information is provided in Table 1.

**Anatomy and Velocity Reconstruction**

A transverse stack of static CMR images served as the basis of TCPC anatomic reconstruction. In-plane resolution was 1.18 mm² on average (range, 0.85-1.88 mm), and average through-plane resolution was 4 mm (range, 3-5 mm). The images were interpolated, segmented in MATLAB (The Mathworks, Natick, Mass), and reconstructed into a 3-dimensional surface in Geomagic Studio (Geomagic, Inc, Research Triangle Park, NC). The Nakata Index was calculated using the mean pulmonary artery radii from the 3-dimensional reconstructions.

Single-slice phase contrast magnetic resonance imaging (PC MRI) was acquired in the ascending aorta, superior vena cava (SVC), inferior vena cava (IVC)/Fontan baffle, proximal left pulmonary artery (LPA) and right pulmonary artery (RPA), and persistent left SVC or aygous vein, when appropriate. These images were segmented semiautomatically to calculate vessel specific blood flow rates. Fenestration flow was estimated from the difference between systemic venous flow (Qs) and pulmonary arterial flows.

**Computational Fluid Dynamics**

Simulations were divided among several investigators and performed using a previously validated solver based on the immersed boundary method, in which the governing equations were solved in their complete unsteady formulation. The patient-specific models were fitted with a surface mesh in Gambit (ANSYS Inc, Lebanon, NH) and registered within a fixed Cartesian grid. Grid spacing was generally set at 2% of the IVC inlet diameter to achieve mesh independence, producing a minimum of 371k nodes in the computational domain.

Because the primary CFD end point was a representative, time-averaged measure of power loss, simulations assumed time-averaged boundary conditions based on the PC MRI data for each vessel. At the inlets, a flat velocity profile was imposed. The presence of a fenestration was ignored. At the outlets, flow boundary conditions were imposed on the basis of the ratio of measured vessel flow to total pulmonary arterial flow. In a few cases when a stenosis was present in 1 branch pulmonary artery, the measured flow rate in the stenotic artery was retained for the simulation while flow through the contralateral branch pulmonary artery was assumed to be equal to the total caval return less the flow in the stenotic artery. Because the sum of the measured inflows usually exceeds the sum of the measured outflows (due in part to fenestration flow), imposing the measured pulmonary artery ratio would typically overestimate the measured arterial flows in the simulation. It is known that connection energetics can be dominated by localized pulmonary artery stenoses, so the practice described earlier prevented the artificial increase in imposed flow across the stenotic region. When the right upper lobe branch artery was retained in the computational domain, the measured RPA flow split (taken proximal to the branch) was divided between the lower and upper branches on the basis of the ratio of their respective areas.

The CFD outputs include the pulmonary distribution of IVC flow (assumed to be uniformly carrying hepatic flow distribution [HFD]), calculated via the flux distribution of IVC streamlines, and power loss according to the following:

\[
\text{PowerLoss} = \sum_{\text{inlets}} \int_{A} \left( p + \frac{1}{2} \rho v^2 \right) v \cdot dA - \sum_{\text{outlets}} \int_{A} \left( p + \frac{1}{2} \rho v^2 \right) v \cdot dA
\]

where \( p \) is the static pressure, \( \rho \) is density, \( A \) is vessel area, and \( v \) is the velocity. Indexed power loss (iPL) was as follows:

\[
\text{iPL} = \frac{\rho Q_s}{\text{BSA}^2}
\]

where \( Q_s \) is the systemic venous flow [L \cdot s \(^{-1}\)], and \( \text{BSA} \) is the body surface area.

**Comparison of Time-Averaged and Pulsatile Boundary Conditions**

To quantify the effect of the selected (time-averaged) boundary conditions on power loss, a subset (n = 35) of cases also were simulated using time-varying boundary conditions. For these simulations, the cardiac-gated PC MRI flow rates were directly imposed on the inlet vessels, whereas the outlet boundary conditions were based on the measured time-varying outflow distribution to enforce continuity of mass. Pulsatile power loss values from these simulations were compared with the paired results of the time-averaged simulations using a Bland–Altman analysis and the coefficient of variation (ratio of averaged pairwise standard deviations to the group mean).

**Four-Dimensional Phase Contrast Magnetic Resonance Imaging**

To further justify the modeling assumptions, simulation results and 4-dimensional PC MRI data interpolated with a divergence-free basis function were quantitatively and qualitatively compared for 6 patients: 3 with extracardiac and 3 with intra-atrial connections. Flow distribution from the 4-dimensional PC MRI data was quantified using the built-in temporal interpolation and particle tracking functionalities of ParaView (paraview.org) and compared with CFD data for the same patients.

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**Abbreviations and Acronyms**

- BSA = body surface area
- CFD = computational fluid dynamics
- CMR = cardiac magnetic resonance
- HFD = hepatic flow distribution
- iPL = indexed power loss
- IVC = inferior vena cava
- LPA = left pulmonary artery
- PC MRI = phase contrast magnetic resonance imaging
- Qs = systemic venous flow
- RPA = right pulmonary artery
- SVC = superior vena cava
- TCPC = total cavopulmonary connection
- WU = Wood units
Statistics

Statistical analyses were performed using Minitab (Minitab Inc, State College, Pa). Continuous variables were tested for normality using the Anderson–Darling test. The Student t test and Mann–Whitney test were used for comparisons between groups, as appropriate, and Pearson’s correlation was used to identify relationships between variables. The analysis presented is exploratory in nature, and thus no adjustments were made for multiple comparisons.

RESULTS

Computational Fluid Dynamics and Cardiac Magnetic Resonance Comparison

Figures 1 and 2 show instantaneous velocity streamlines for the time-averaged CFD and representative time points from the phasic CMR results for 3 patients with extracardiac connection and 3 patients with intra-atrial connection, respectively. In patient A, both modalities show that the caval flow collision favors SVC flow to the RPA, whereas IVC flow was evenly split. For patient B, the SVC flow streams entirely to the LPA with noticeable flow acceleration through the subaortic section of the pulmonary artery. A distinct flow collision/stagnation point (Figure 1, arrow) is seen in the center of the connection for patient C with IVC flow favoring the LPA.

In Figure 2, the intra-atrial flows are marked by complex mixing and recirculation patterns. Some dynamic attributes are seen from the CMR (eg, the transient translation of the central vortex in patient D) that are not entirely captured by the time-averaged CFD results; however, the general characteristics and overall velocity magnitudes are still conserved. For example, the SVC flow is constrained along the lateral right wall for all 3 patients (with large vortices forming in patients D and F) and primarily exits the RPA.

Comparison of the HFD results between modalities is provided in Figures 1 and 2. Differences ranged from 3% to 16%, with an average absolute difference of 11.8% and a coefficient of variation of 14%.

Time-Averaged Versus Pulsatile Boundary Conditions

Figure 3 shows the Bland–Altman mean versus difference plot of time-averaged versus pulsatile boundary condition power losses. The 95% limits of agreement are denoted by the “2SD” lines. Pulsatile power loss was generally higher than time-averaged for a given patient, as indicated by the negative average difference. Despite this systematic

TABLE 1. Patient demographic details

<table>
<thead>
<tr>
<th>Details</th>
<th>Values</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (y)</td>
<td>12.0 ± 6.8</td>
<td></td>
</tr>
<tr>
<td>BSA (m²)</td>
<td>1.23 ± 0.46</td>
<td></td>
</tr>
<tr>
<td>Nakata Index (mm²/m²)</td>
<td>219 ± 89</td>
<td></td>
</tr>
<tr>
<td>Gender (male/female)</td>
<td>55/45</td>
<td></td>
</tr>
<tr>
<td>Connection type</td>
<td>64/33/3</td>
<td></td>
</tr>
<tr>
<td>HLHS vs non-HLHS</td>
<td>31/69</td>
<td></td>
</tr>
<tr>
<td>Bilateral SVC connections</td>
<td>15</td>
<td></td>
</tr>
</tbody>
</table>

Data are reported as mean ± standard deviation. BSA, Body surface area; HLHS, hypoplastic left heart syndrome; SVC, superior vena cava.

FIGURE 1. Three-dimensional velocity fields from time-averaged computational simulations (left) and representative phases from the CMR acquisitions (right) for 3 patients with an extracardiac connection. Comparison of hepatic distribution results (as %LPA) shown for each case. Arrow for patient C highlights a conserved region of flow stagnation at the caval flow collision site. CFD, Computational fluid dynamics; HFD, hepatic flow distribution; LPA, left pulmonary artery; MRI, magnetic resonance imaging; Pt., patient.
bias, the variation in these differences was small compared with the mean power loss values, with a coefficient of variation of 11%.

Cohort Results

Flow data derived from the through-plane PC MRI acquisitions and normalized to body surface area (BSA) are summarized in Table 2. On average, fenestration flow was less than 10% of Qs.

Hemodynamic findings from the CFD analysis are presented in Table 3. Mean HFD was 44% (to the LPA), and it correlated with the average pulmonary flow distribution of 45% to the LPA ($r = 0.36, P < .001$; Figure 4, A). Additional correlations included BSA with both resistance and iPL ($r = 0.28$ and $0.37$, respectively; $P < .05$; Figure 4, B) and age with both cardiac index and iPL ($r = -0.37$ and 0.26, respectively; $P < .05$). After correcting for the presence of flow in the power loss indexing scheme with partial correlation, the natural logarithm of iPL was significantly correlated with both $Q_s$ ($r = -0.31, P = .001$; Figure 4, C).

### Table 2. Average flow data (L/min/m²) by vessel from phase contrast magnetic resonance imaging

<table>
<thead>
<tr>
<th>Vessel</th>
<th>Mean ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>IVC</td>
<td>1.72 ± 0.55</td>
</tr>
<tr>
<td>SVC</td>
<td>1.00 ± 0.53</td>
</tr>
<tr>
<td>LSVC*</td>
<td>0.64 ± 0.35</td>
</tr>
<tr>
<td>Azygos vein*</td>
<td>0.92 ± 0.37</td>
</tr>
<tr>
<td>LPA</td>
<td>1.21 ± 0.63</td>
</tr>
<tr>
<td>RPA</td>
<td>1.38 ± 0.50</td>
</tr>
<tr>
<td>Fenestration†</td>
<td>0.30 ± 0.54</td>
</tr>
</tbody>
</table>

Data are reported as mean ± standard deviation. IVC, Inferior vena cava; LPA, left pulmonary artery; LSVC, left superior vena cava; RPA, right pulmonary artery; SVC, superior vena cava. *Not applicable to all patients. †Not measured for all patients.

**FIGURE 2.** Three-dimensional velocity fields between time-averaged computational simulations (left) and representative phases from the CMR acquisitions (right) for 3 patients with an intra-atrial connection. Comparison of hepatic distribution results (as %LPA) shown for each case. CFD, Computational fluid dynamics; HFD, hepatic flow distribution; LPA, left pulmonary artery; MRI, magnetic resonance imaging; Pt., patient.

**FIGURE 3.** Bland–Altman mean versus difference comparison of power losses derived from simulations using pulsatile and time-averaged boundary conditions. The 95% confidence intervals are small compared with average power losses.
and cardiac index \( r = -0.21, P = .04 \). Finally, the Nakata Index had a power law fit to both iPL \( r = -0.52, P < .001 \) and connection resistance \( r = -0.43, P < .001 \).

### Intra-Atrial Versus Extracardiac Connections

In this comparison (neglecting 3 atriopulmonary connections), patients with intra-atrial connection were significantly older and had larger BSA, whereas there was a strong trend \( (P = .07) \) toward a higher cardiac index in patients with extracardiac connection (Table 4). There were no statistical differences in resistance or iPL between groups \( (P = .48 \) and \( .24, \) respectively). The pulmonary flow distribution to the LPA was significantly higher in extracardiac connections, but there was a trend toward lower HFD to the LPA \( (P = .08) \) in those connections.

### Highest Versus Lowest Power Loss Connections

Table 5 compares the characteristics and hemodynamics of the 5 highest and 5 lowest power loss (iPL) connections. The highest loss cases, although being older and having a higher BSA, all had energetic measures at least 1 order of magnitude larger (ie, more dissipative) than the lowest loss ones. \( Q_s \) and cardiac indices also were notably lower. Finally, the composition of the highest loss group with respect to connection type is notable because 3 of the 5 intra-atrial connections were baffle conduits (ie, not intra-atrial lateral tunnel connections).

Figure 5 shows 3-dimensional velocity streamlines for the 5 highest power loss connections. In all cases, appreciable narrowing of the Fontan pathway to varying degrees created local acceleration, pressure decrease, and recirculation within the connections. Figure 6 shows a similar 3-dimensional velocity streamline analysis of the lowest power loss connections, which had lower maximal velocities and more ordered flow patterns than in Figure 5.

### DISCUSSION

**Fontan Power Loss and the Single Ventricle**

TCPC design is one of the factors amenable to interventional/surgical manipulation in the care of patients with

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**TABLE 3.** Cohort hemodynamic results

<table>
<thead>
<tr>
<th></th>
<th>Cardiac index (L/min/m²)</th>
<th>Pulmonary flow distribution</th>
<th>Resistance (WU)</th>
<th>iPL</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>Qs (L/min/m²)</td>
<td>HFD (%LPA)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>3.47</td>
<td>2.89</td>
<td>45</td>
<td>0.23</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>1.11</td>
<td>12</td>
<td>0.17</td>
</tr>
<tr>
<td></td>
<td>Median</td>
<td>3.43</td>
<td>43</td>
<td>0.19</td>
</tr>
<tr>
<td></td>
<td>25th percentile</td>
<td>2.86</td>
<td>37</td>
<td>0.10</td>
</tr>
<tr>
<td></td>
<td>75th percentile</td>
<td>3.85</td>
<td>52</td>
<td>0.28</td>
</tr>
</tbody>
</table>

HFD, Hepatic flow distribution; iPL, indexed power loss; LPA, left pulmonary artery; Qs, systemic venous flow; SD, standard deviation; WU, Wood units.
single ventricles. These data demonstrate that patient-to-patient variation led to a 2 orders of magnitude range in iPL, so elucidating its relationship to patient functional status and outcomes is of significant clinical relevance.

The mean TCPC resistance value from this cohort was 0.23 WU, compared with 1.3 to 1.8 WU for normal pulmonary vascular resistances in patients undergoing the Fontan.24,25 So these data indicate that the TCPC adds 15% to 20% on average to post-hepatic resistance. The significant inverse relationships of iPL with Qs and cardiac index suggest these losses have a relevant effect on cardiac function, particularly when elevated. These results are similar to the linear correlation between TCPC resistance and cardiac index reported previously.25

A physiology-based model of ventricular filling (ie, diastolic function) and cardiac output shows that resistance elements downstream of vascular capacitance have a drastic effect on limiting the ability of the ventricle to fill and thus provide adequate output.26 In the Fontan physiology/circulation, the TCPC and pulmonary vasculature are 2 such critical resistive elements. Thus, a possible mechanism for the present findings is that elevated power loss resulting from suboptimal connection geometry contributes to restricted preload and preload reserve27,28 of the single ventricle, which may limit efficient long-term performance.

### Extracardiac Versus Intra-Atrial Connections

Extracardiac connections had a strong trend (P = .08) toward lower HFD to the LPA compared with intra-atrial connections (38% vs 47%, respectively), despite significantly higher total LPA flow percentage in extracardiac connections (51% vs 41%). A recent study found that HFD correlated with the total pulmonary distribution in intra-atrial connections because of caval mixing within the connection. Conversely, HFD in extracardiac connections have been reported to correlate only with caval offset.29 These observations agree well with the present findings. Although HFD did correlate with the total flow distribution for extracardiacs in this analysis, the significant influence of caval offset was still apparent because more than 60% of IVC flow perfused the RPA (the predominant direction of caval offsetting for such connections) despite a nearly balanced total distribution (51% LPA). Most important, neither result suggests an inherent risk factor for unilateral hepatic distribution and pulmonary arteriovenous malformations.30

The more relevant comparison is in regard to the comparative resistance of the 2 approaches because the superiority of one approach over the other has been the subject of debate for a number of years.31-34 In this series, there were no differences in power loss between the connection types.

The question then remains as to what distinguishes between hemodynamically favorable and unfavorable connections. The importance of pulmonary artery size has been well documented,7,20 and the large difference in the average Nakata indices between the most and least efficient groups provides strong support for its prominence. In addition, the effect of local constriction/undersizing of the Fontan pathway also was apparent from the present data. The worst performing connections were characterized by the presence of high-velocity Fontan flow immediately proximal to the cavopulmonary connection that created large separated flow regions and vortices within the junction (Figure 5). The combination of multiple suboptimal geometric features (eg, stenosis, sharp bends, sudden expansions) in some cases further compounded these inefficiencies. As a result, the iPL in those cases was 3 times greater than the cohort mean. Furthermore, these hemodynamics are likely to be significantly exacerbated under higher cardiac output conditions, and could thus contribute to poor exercise tolerance. In 2 of the cases (L2, L5), lateral tunnel stenosis was present and contributed to these adverse hemodynamics, whereas a small intra-atrial conduit was a problem for 3 patients (L1, L3, L4). On the basis of the present cohort, such adverse geometric characteristics are rare but seem to warrant attention (eg, via stenting the lateral tunnel pathway) when observed clinically.

### TABLE 4. Comparison of intra-atrial and extracardiac connections

<table>
<thead>
<tr>
<th></th>
<th>Intra-atrial (N = 64)</th>
<th>Extracardiac (N = 33)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (y)</td>
<td>13.5 ± 6.5</td>
<td>7.9 ± 4.8</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>BSA (m²)</td>
<td>1.33 ± 0.43</td>
<td>0.96 ± 0.37</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Nakata Index (mm²/m²)</td>
<td>219 ± 78</td>
<td>206 ± 58</td>
<td>.40</td>
</tr>
<tr>
<td>Cardiac index (L/min/m²)</td>
<td>3.30 ± 0.87</td>
<td>3.90 ± 1.41</td>
<td>.07</td>
</tr>
<tr>
<td>Qs (L/min/m²)</td>
<td>2.89 ± 0.89</td>
<td>3.10 ± 0.91</td>
<td>.33</td>
</tr>
<tr>
<td>Pulmonary distribution (L²/LPA)</td>
<td>41 ± 9</td>
<td>51 ± 15</td>
<td>.001</td>
</tr>
<tr>
<td>HFD (%LPA)</td>
<td>46 ± 18</td>
<td>37 ± 26</td>
<td>.08</td>
</tr>
<tr>
<td>Resistance (WU)</td>
<td>0.25 ± 0.19</td>
<td>0.20 ± 0.12</td>
<td>.48</td>
</tr>
<tr>
<td>iPL</td>
<td>0.040 ± 0.031</td>
<td>0.030 ± 0.017</td>
<td>.24</td>
</tr>
</tbody>
</table>

Mean ± standard deviation. BSA, Body surface area; iPL, indexed power loss; LPA, left pulmonary artery; Qs, systemic venous flow; WU, Wood units.

### TABLE 5. Comparison of highest and lowest power loss total cavopulmonary connections

<table>
<thead>
<tr>
<th></th>
<th>5 most efficient</th>
<th>5 least efficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (y)</td>
<td>9.2 ± 4.9</td>
<td>16.6 ± 3.8</td>
</tr>
<tr>
<td>BSA (m²)</td>
<td>0.91 ± 0.39</td>
<td>1.70 ± 0.29</td>
</tr>
<tr>
<td>Nakata Index (mm²/m²)</td>
<td>357 ± 132</td>
<td>180 ± 41</td>
</tr>
<tr>
<td>Connection type (intra-atrial/ extracardiac/atriopulmonary)</td>
<td>4/1/0</td>
<td>5/0/0</td>
</tr>
<tr>
<td>Cardiac index (L/min/m²)</td>
<td>3.89 ± 0.64</td>
<td>3.33 ± 0.66</td>
</tr>
<tr>
<td>Qs (L/min/m²)</td>
<td>3.57 ± 0.85</td>
<td>2.54 ± 0.22</td>
</tr>
<tr>
<td>Resistance (WU)</td>
<td>0.05 ± 0.01</td>
<td>0.66 ± 0.09</td>
</tr>
<tr>
<td>iPL</td>
<td>0.007 ± 0.003</td>
<td>0.117 ± 0.012</td>
</tr>
</tbody>
</table>

Mean ± standard deviation. BSA, Body surface area; iPL, indexed power loss; Qs, systemic venous flow; WU, Wood units.
Four-Dimensional Phase Contrast Magnetic Resonance Imaging and Computational Fluid Dynamics in Fontan Evaluation

This study used both 4-dimensional PC MRI and CFD to characterize Fontan hemodynamics; advancements in both technologies have increased the frequency of their use for such applications, yet few, if any, studies have provided a direct comparison of their results in the TCPC or any other vascular region. The primary findings from this comparison were quantitative similarities in HFD measures and qualitative similarities in the 3-dimensional flow fields between modalities, which support the accuracy of the CFD simulations.

Furthermore, although there are considerable areas of overlap in the knowledge gained from these modalities, there are important distinctions that should be noted that made imaging-based CFD a unique and important asset in this analysis. The first point is the availability of large datasets: The CFD protocol used requires basic imaging inputs, which allowed for the use of retrospective data to be used in generating a large patient sample. By comparison, 4-dimensional PC MRI is only recently gaining in popularity and is still not a part of routine clinical protocols; a study of this size based solely on 4-dimensional PC MRI analyses is still several years away. Assessing power loss is another strength of CFD because of the different spatial resolution characteristics of the 2 modalities: Fontan 4-dimensional PC MRI studies have reported voxel sizes ranging from 5.8 to 17.5 mm$^3$, whereas the grid size for the solver in our study was on the order of 0.05 mm$^3$. These differences have significant implications for the ability of each method to resolve velocities close to vessel boundaries and accurately evaluate spatial derivatives of the velocity fields, both of which are crucial for hemodynamic power loss evaluation. Finally, there are technical considerations related to the acquisition of CMR and the potential for cycle averaging, noise, finite velocity measurement resolution, and turbulent de-phasing to affect measurement fidelity. CFD has its own limitations, as discussed next, but an accurate computational result clearly provides a superior means of quantitative analysis of the velocity field. Moving forward, both techniques will play critical, complementary roles in further understanding this complex physiology.

Study Limitations

The simulations were all performed with the assumptions of time-averaged flow conditions and rigid vessel walls. Figure 3 indicates that the time-averaged flow assumption introduced a small quantitative bias in the power loss results compared with simulations assuming pulsatile flow boundary conditions and rigid walls (which is generally considered to be the current state-of-the-art). However, on the basis of the constrained limits of agreement and the low...
coefficient of variation, the effect of the time-averaged flow assumption on power loss was small and acceptable for quantification of this time-averaged end point. Moreover, the quantitative consistency and qualitative similarities between the CFD and 4-dimensional PC MRI data provide similar assurance that despite the simplifying assumptions made, the simulated results presented are reasonable representations of the in vivo dynamics. Spatial resolution limitations in the imaging technique can have negative effects on the fidelity of reconstructed vessel sizes, particularly by exaggerating the severity of stenotic PAs. However, the techniques used for image interpolation and segmentation have been rigorously validated against phantoms, so we are confident in the overall accuracy of these models.

Fenestration flow was systematically ignored, which is a limitation of the retrospective data analysis. However, because fenestration flow accounted for less than 10% of systemic flow on average, that assumption was acceptable.

There was a significant difference in age between the extracardiac and intra-atrial groups, which presents a challenge in making direct comparisons. However, the observed association between age and iPL was weak and positive, suggesting that as the patients in the extracardiac group (younger group) age, their average iPL may increase toward the current average of the intra-atrial group from this cohort. Thus, it is unlikely that the finding of no difference between groups is affected by this age discrepancy.

Finally, the approach to the Fontan connection can vary considerably from center to center and surgeon to surgeon. Although this is a single-center experience, the present findings comparing intra-atrial and extracardiac connections may be globally applicable because there are no specific practices at this center that would create a significant distinction with the broader approach to Fontan surgery.

CONCLUSIONS

Congenital heart disease occurs in 1% of the population, and single ventricle defects make up a smaller subset; the ability to acquire and analyze large patient data samples is a unique challenge and significant accomplishment. From this largest computational analysis of the Fontan connection to date, several clinically relevant insights were gained. First, power loss varies widely among patients with Fontan and may vary with age and development. Second, there is an inverse correlation between indexed TCPC power loss and $Q_s$, which supports the hypothesis that TCPC hemodynamics can negatively affect ventricular filling and preload. Third, no energetic differences were observed between intra-atrial and extracardiac Fontan connections, but the presence of undersized pulmonary arteries or Fontan pathway stenosis did have a detrimental effect on power loss.

References


