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# **ORIGINAL ARTICLE**

# Myocardial Perfusion in Hypoplastic Left Heart Syndrome

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**BACKGROUND:** The status of the systemic right ventricular coronary microcirculation in hypoplastic left heart syndrome (HLHS) is largely unknown. It is presumed that the systemic right ventricle's coronary microcirculation exhibits unique pathophysiological characteristics of HLHS in Fontan circulation. The present study sought to quantify myocardial blood flow by cardiac magnetic resonance imaging and evaluate the determinants of microvascular coronary dysfunction and myocardial ischemia in HLHS.

**METHODS**: One hundred nineteen HLHS patients (median age, 4.80 years) and 34 healthy volunteers (median age, 5.50 years) underwent follow-up cardiac magnetic resonance imaging  $\approx$ 1.8 years after total cavopulmonary connection. Right ventricle volumes and function, myocardial perfusion, diffuse fibrosis, and late gadolinium enhancement were assessed in 4 anatomic HLHS subtypes. Myocardial blood flow (MBF) was quantified at rest and during adenosine-induced hyperemia. Coronary conductance was estimated from MBF at rest and catheter-based measurements of mean aortic pressure (n=99).

**RESULTS**: Hyperemic MBF in the systemic ventricle was lower in HLHS compared with controls ( $1.89\pm0.57$  versus  $2.70\pm0.84$  mL/g per min; P<0.001), while MBF at rest normalized by the rate-pressure product, was similar ( $1.25\pm0.36$  versus  $1.19\pm0.33$ ; P=0.446). Independent risk factors for a reduced hyperemic MBF were an HLHS subtype with mitral stenosis and aortic atresia (P=0.017), late gadolinium enhancement (P=0.042), right ventricular diastolic dysfunction (P=0.005), and increasing age at total cavopulmonary connection (P=0.022). The coronary conductance correlated negatively with systemic blood oxygen saturation (r, -0.29; P=0.02). The frequency of late gadolinium enhancement increased with age at total cavopulmonary connection (P=0.014).

**CONCLUSIONS:** The coronary microcirculation of the systemic ventricle in young HLHS patients shows significant differences compared with controls. These hypothesis-generating findings on HLHS-specific risk factors for microvascular dysfunction suggest a potential benefit from early relief of frank cyanosis by total cavopulmonary connection.

Key Words: heart defects, congenital 
hypoplastic left heart syndrome 
Fontan circulation 
magnetic resonance imaging
myocardial perfusion imaging

The 3-stage surgical palliation of hypoplastic left heart syndrome (HLHS) is considered one of the major achievements of congenital heart surgery.<sup>1,2</sup> The Fontan circulation after complete surgical palliation presents

new challenges for understanding potential pathophysiological characteristics that increase the risk of forthcoming right ventricular (RV) failure.<sup>1,3,4</sup> Among those, the status of the coronary microcirculation in HLHS has been relatively

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### **CLINICAL PERSPECTIVE**

The status of the coronary microcirculation in the systemic right ventricles (RVs) of patients with hypoplastic left heart syndrome (HLHS) is largely unknown. Coronary dysfunction and myocardial ischemia are important factors in the etiology of heart disease with and without preserved ejection fraction. This prospective cardiac magnetic resonance study aimed at identifying the determinants and risk factors of microvascular coronary dysfunction and myocardial ischemia in HLHS. Volumes and function of the systemic ventricle, myocardial perfusion at rest, and during adenosine-induced hyperemia, diffuse myocardial fibrosis, and late gadolinium enhancement were assessed in 4 different anatomic subtypes of HLHS. Cardiac magnetic resonance imaging in 119 HLHS patients and 34 healthy controls revealed that the coronary microcirculation of the systemic ventricle in young HLHS patients is markedly impaired compared with controls. The coronary conductance at rest correlated negatively with systemic blood oxygen saturation, suggesting a form of compensatory coronary vasodilation, and the frequency of late gadolinium enhancement increased with age at total cavopulmonary connection. Diastolic dysfunction, the presence of the mitral atresia/ aortic stenosis subtype, and an older age at total cavopulmonary connection appear to be risk factors for RV microcirculatory dysfunction. In summary, this study provides new insights into factors unique to HLHS patients that affect the RV microcirculation and tissue structure and may generate novel hypothesis for future long-term studies. Such studies are required to investigate the role of the RV microcirculation leading to RV dysfunction in HLHS patients over time.

### Nonstandard Abbreviations and Acronyms

| AA   | aortic atresia                  |
|------|---------------------------------|
| AS   | aortic stenosis                 |
| CMR  | cardiac magnetic resonance      |
| HLHS | hypoplastic left heart syndrome |
| LGE  | late gadolinium enhancement     |
| LV   | left ventricle                  |
| MA   | mitral atresia                  |
| MBF  | myocardial blood flow           |
| MS   | mitral stenosis                 |
| RPP  | rate-pressure product           |
| RV   | right ventricle                 |
| ТСРС | total cavopulmonary connection  |
| VCC  | ventriculocoronary connection   |

unexplored, despite the fact that the inevitable extensive vascular surgery during the Norwood operation often leads to sympathetic denervation of the heart.<sup>5-7</sup> Thus, the systemic RV's coronary microcirculation in children with HLHS is assumed to differ significantly compared with normal left ventricles (LVs) in non-HLHS subjects.<sup>8-11</sup> Moreover, myocardial perfusion may vary among HLHS anatomic subtypes, in light of previous studies that assigned a higher risk for stage 1 and interstage mortality to the subtype with mitral stenosis (MS) and aortic atresia (AA) compared with the other subtypes.<sup>2,12,13</sup>

Positron emission tomography with radioactive flow tracers has been considered the gold standard to quantify myocardial blood flow (MBF).<sup>14–18</sup> However, due to radiation exposure, nuclear medicine methods are generally less preferred in children. Over the last decade, cardiac magnetic resonance (CMR) imaging has become a major alternative for quantitative determination of myocardial perfusion at rest and during maximal hyperemia.<sup>19</sup>

This study aims to quantify the global and regional coronary microcirculation in the systemic RV of HLHS patients compared with control subjects with healthy hearts and on that basis identify HLHS-specific risk factors for microvascular dysfunction.

## **METHODS**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

### **Study Subjects**

This prospective CMR cohort consists of 119 HLHS patients from the University of Schleswig-Holstein/Campus Kiel Medical Center (median age, 4.8; 67.23% women). The patients' cohort includes all 4 major anatomic subtypes of MS/aortic stenosis (AS), MS/AA, mitral atresia (MA)/AS, and MA/AA as previously described by our center (Figure 1).20 All HLHS patients underwent a comprehensive CMR study after the final palliative step with completion of the total cavopulmonary connection (TCPC) using an intraatrial lateral tunnel. The same lead surgeon (J.S.) and his team at the University of Schleswig-Holstein/Campus Kiel Medical Center operated on all the studied HLHS patients. Patients with moderate or severe tricuspid or aortic valve insufficiency or with a hemodynamically significant gradient (>1.6 m/s by Doppler echocardiography) in the RV outflow tract or aorta, as well as patients with pacemakers were excluded from the study. Most patients (86%) were sedated with propofol and midazolam during the CMR scan. Heart rate, respiratory motion, oxygen saturation, and noninvasive blood pressure were monitored during CMR studies. The study was performed according to the ethical standards of the 1964 Declaration of Helsinki and its later amendments and was approved by our hospital's institutional review board (No. D526/16). Informed written consent was obtained from the patients or their legal guardians.

The control group was recruited among the subjects who underwent a clinically indicated CMR study with administration of gadolinium-based contrast. More specifically, 18 subjects (median age, 5.1 years; 44.4% women) who were ruled out





for any cardiac abnormality were selected as control subjects at the University of Schleswig-Holstein/Campus Kiel Medical Center. To increase the diversity of control subjects, we included the perfusion studies of an additional 16 healthy subjects (<14 years; median age, 7.1 years) from a previously published study<sup>21</sup> from the Oregon Health and Science University, as performed by coauthors E.M., M.S., M.J.-H., and their colleagues using the same type/model of 3T magnetic resonance scanner. All 34 control subjects from Kiel and Oregon underwent the same first-pass magnetic resonance perfusion protocol, which has been originally established by the same investigator (M.J.-H.) at both institutions.<sup>21</sup>

### **CMR** Techniques

CMR studies were performed on a 3.0T MRI scanner (Achieva 3.0T; Philips Medical Systems, the Netherlands), using a phasedarray coil for cardiac imaging, or for small children, a phased-array coil for extremities (SENSE Cardiac coil, SENSE Flex-L coil; Philips Medical Systems). Gradient echo cine CMR (Figure 2A) with retrospective ECG gating was performed to evaluate RV volumes, mass and systolic function, and to define the anatomic subtypes, applying the following scan parameters: field of view, ≤280×224 mm; voxel size, 1.6×1.5×5 mm<sup>3</sup>; repetition time/ echo time, 3.6/1.8 ms; 25 cardiac phases; nonbreath-hold; number of repetitions, 2; and total scan duration, 3 to 6 minutes. Myocardial perfusion images (Figure 2B) were acquired with an ECG-gated, multislice, saturation-recovery-prepared, single-shot gradient echo sequence with the following parameters: repetition time/echo time/flip angle, 1.2/2.6 ms/20°; 192×160 acquisition matrix; voxel size,  $\approx 1.8 \times 2.0$ ; parallel imaging  $\times 2$  acceleration with sensitivity encoding (SENSE); slice, 10 mm; 2 slices/RR; 2 short-axis slices; and 60 dynamics. A low-dosage gadolinium contrast bolus (0.03 mmol/kg of Magnevist; Bayer, Germany) was administered through an antecubital vein with a power injector (MEDRAD Spectris, Bayer, Germany) at a rate of 4 mL/s. Maximal hyperemia was induced by intravenous adenosine with a stepwise dosage escalation from 50 µg/kg per minute, through 100 µg/kg per minute, to 140 µg/kg per minute over ≈9 minutes (3 minutes per stage), before the start of the stress perfusion scan. The adenosine infusion was turned off once the contrast bolus appeared in the LV on the reconstructed perfusion images (≈2 seconds latency time).<sup>22</sup> Rest perfusion images were acquired first, and hyperemic flows were measured with at least 20 minutes of delay after the rest scan.

To reach a total contrast dosage of 0.1 mmol/kg, a top off bolus was administered after the perfusion studies. Approximately 15 minutes later, late gadolinium enhancement



A, The red line on the long-axis view (4-chamber view) from a cine acquisition denotes the orientation of short-axis image slices. B, For perfusion image analysis, the right ventricular (RV) wall in short-axis view was divided into 4 anatomic segments (1, septal/ posterior; 2, inferior; 3, anterior; 4, superior). This is a typical first-pass perfusion image acquired during hyperemia in an HLHS patient with rudimentary left ventricular (LV) cavum.



Myocardial Perfusion in HLHS

(LGE) images were acquired with an inversion recovery–prepared 3-dimensional gradient echo sequence (field of view, 300×178×80 mm; voxel size, 1.17×1.27×10 mm; repetition time/echo time, 3.7/1.83 ms; flip angle, 15°; TI adjusted to null normal myocardium) with respiratory navigator for detection of diaphragmatic motion. The presence and location of LGE was qualitatively identified within the myocardium by windowing images to null the signal in normal myocardium. Myocardial and blood T1s were quantified with a cine Look-Locker technique in one short-axis slice in the mid-RV as described previously.<sup>23,24</sup>

# Hemodynamic Measurements During Cardiac Catheterization

Cardiac catheterization was routinely performed under deep sedation on the day after the CMR scan. The full hemodynamic assessment included measurements of the mean aortic pressure and the RV end-diastolic pressure. Cine angiograms, including a selective injection into the native aortic root were acquired in standard projections and used to identify ventriculocoronary connections (VCCs) as previously described by our group.<sup>25</sup>

RV diastolic function was defined as an invasive end-diastolic pressure >10 mmHg concordant with previous studies on diastolic dysfunction in Fontan patients.<sup>26,27</sup>

### **Image Analysis and Calculations**

All CMR studies were analyzed on a workstation with dedicated software (ViewForum release 6.3; Philips Medical Systems). Anatomic HLHS subtypes were determined from axial and short-axis magnetic resonance cine image stacks and echocardiography. RV end-diastolic and end-systolic volumes were determined by planimetry of all short-axis images. Stroke volume was calculated by defining the RV volume at end diastole and then subtracting the RV volume at end systole. RV ejection fraction was calculated by dividing the stroke volume by the RV end-diastolic volume. LGE in the RV was quantified using the full-width, half-maximum criterion and expressed in grams or percentage of RV mass.

The perfusion images were segmented along the endo- and epicardial borders, and the RV myocardium was subdivided into 4 segments of equal angular extent from the RV centroid point (Figure 2B). The region of interest was drawn in the center of the RV to sample the arterial input of gadolinium contrast. This arterial input was corrected for saturation effects, using a model-based calibration. MBFs were estimated by modelindependent deconvolution of the myocardial signal-intensity curves with the saturation-corrected arterial input measured, as described previously.<sup>28</sup> Examples of signal-intensity curves for an HLHS patient and a volunteer are shown in Figure I in the Data Supplement. Unless stated otherwise, all results for MBF at rest refer to the MBF values normalized by the patient's rate-pressure product (RPP; heart rate in beats per minute multiplied by systolic pressure in mm Hg/10<sup>4</sup>). RPP was used here as a surrogate measure for the resting cardiac workload. In keeping with previous studies of MBF with positron emission tomography<sup>29,30</sup> and magnetic resonance imaging,<sup>31</sup> the MBF at rest was normalized by the RPP. RPP was calculated as heart rate, multiplied by systolic blood pressure and divided by 10000, to result in MBF values around 1.0, not dissimilar

to uncorrected MBF. Hyperemic MBF was not normalized by RPP, since the hyperemic response is caused by vasodilation with adenosine, which at best has a weak association with the cardiac workload. Coronary conductance was estimated from resting MBF in mL/min per gram, divided by the mean aortic pressure (mmHg)<sup>32</sup> measured during catheterization within a day of the CMR study.

Look-Locker images were segmented along the RV (HLHS) or LV (controls) endo- and epicardial borders, which were further divided into 6 standard segments. T1 was determined for each segment with a nonlinear least-squares algorithm and correction for radiofrequency pulse effects. The myocardial partition coefficient for gadolinium was estimated as the rate of change of myocardial change of R1, as a function of the R1 change in the blood pool. The extracellular volume fraction for each segment was calculated as the product of the partition coefficient with (1–hematocrit fraction), as described in previous studies.<sup>33</sup>

### **Statistical Analysis**

Statistical analysis was performed using the R software (version 4.0.2, 2019; R Foundation for Statistical Computing, Vienna, Austria).<sup>34</sup> Data are reported as mean±SD or if not normally distributed, as median and interquartile range. Means between two groups were compared with the Student t test or the Wilcoxon-Mann-Whitney test as nonparametric analog. Fisher exact test was used to test associations between categorical variables. ANOVA was used to compare multiple groups, and P values obtained with multiple post hoc testing were adjusted with the Holm method. The strength of correlations between variables was assessed with Spearman rank correlation. A multivariate linear regression model was constructed for hyperemic MBF and included anatomic subtypes, the presence of a rudimentary LV, age at the time of TCPC, presence of LGE or percentage of LGE, RV diastolic dysfunction, oxygen saturation, age at the time of CMR scan, and sex as predictors. The presence of LGE was analyzed with a multivariate logistic regression model. Predictors considered for the logistic regression model were age at TCPC, age at the time of CMR scan, HLHS subtype, and hyperemic MBF. The final set of predictors for linear and logistic regression models was determined by stepwise forward and backward selection of predictors using the Akaike Information Criterion (stepAIC function in R package MASS) to arrive in each case at the most parsimonious model. Variance inflation factors were determined to guard against any potential predictor collinearities, for example, age at TCPC and age at the time of CMR scan, using a variance inflation factor threshold of 2.5. Statistical tests were 2 tailed, and P values of <0.05 were considered statistically significant.

## RESULTS

### **Patient Characteristics**

HLHS patients underwent CMR at a median of 1.8 years (interquartile range, 2.7; range, 0.8–16 years) after TCPC. Six HLHS patients with tricuspid regurgitation (n=3) or aortic regurgitation (n=3) were excluded from further analysis. The HLHS patients were comparable in age,

weight, body height, and body surface area to controls, as shown in Table 1. Heart rate, systolic blood pressure, and the RPP were significantly lower in HLHS (P<0.01). A Blalock-Taussig shunt in the context of the Norwood operation was initially established in 115 (96.6%) HLHS patients, and 4 (3.4%) received a (nonvalved) RV-PA conduit (Sano-Shunt). Additional patient characteristics are summarized in Table 1.

The distribution of HLHS anatomic subtypes (Figure 1) was comparable to previous studies.<sup>2,13,35</sup> A rudimentary LV with a visible LV cavum on short-axis CMR images was present in the subtypes MA/AS (6%), MS/AA (24%), and MS/AS (29%), see example in Figure 2. In the subtypes MA/AA (42% of cases), no visible LV cavum was present.

### **RV Volumes and Function**

Table 2 summarizes the CMR results for the systemic ventricle in controls and HLHS patients. In comparison to healthy control subjects, HLHS patients had higher indexed systemic ventricle volumes at end diastole and end systole, a higher indexed wall mass, and a lower ejection fraction but a similar cardiac index. The RV cardiac index did not vary significantly (ANOVA, P=0.128) among HLHS anatomy subtypes. RV volume or function parameters did not significantly differ among the HLHS subtypes. RV ejection fraction correlated negatively with age at TCPC (r, -0.244; P=0.01).

### Late Gadolinium Enhancement

LGE imaging was available for 103 (87%) HLHS patients. LGE imaging was performed at the end of the exam and had to be skipped in patients who awoke early from sedation. LGE was not reported for cases where part of the RV wall was affected by artifacts from metallic implants. Myocardial LGE was detected in 17 (14%) of the HLHS patients. In 8 (7%) patients, LGE was present in the RV, with a median of 2.8 g (6.9 % of RV mass; interguartile range, 6.7%) of scar tissue. The age at TCPC and hyperemic MBF were identified by stepwise selection as best predictors in a logistic regression model for the presence of LGE. Based on this logistical regression model, LGE was more likely with increasing age at TCPC (odds ratio, 2.1:1 for 1-year increase; P=0.019; Figure II in the Data Supplement), while the probability of LGE trended lower with increasing hyperemic MBF (odds ratio, 0.32:1 for 1 mL/min per g increase of hMBF; P=0.066).

### **Myocardial Perfusion**

MBF at rest was estimated in 108 (91%) and hyperemic MBF in 101 (85%) of HLHS patients. Myocardial perfusion could not be quantified in cases with severe

# Table 1. Clinical Characteristics and CMR-Computed Measures in HLHS Patients and Controls

|                                    | Control (n=34)   | HLHS (n=119)     | P value |  |
|------------------------------------|------------------|------------------|---------|--|
| Female sex                         | 18 (52.94)       | 81 (67.23)       | <0.001  |  |
| Age at CMR, y                      | 5.50 (3.42-8.40) | 4.80 (4.00-6.55) | 0.797   |  |
| Height, cm                         | 115.27±27.73     | 111.57±17.56     | 0.472   |  |
| Weight, cm                         | 24.94±13.29      | 20.75±10.23      | 0.101   |  |
| BSA, m²                            | 0.88±0.34        | 0.79±0.24        | 0.184   |  |
| $O_{_2}$ saturation, %             | 98.43±1.27       | 90.52±4.68       | 0.008   |  |
| Hematocrit, %                      | 41.33±1.75       | 42.36±3.74       | 0.378   |  |
| Heart rate, bpm                    | 89.15±17.78      | 75.94±15.93      | <0.001  |  |
| BP sys, mmHg                       | 92.00±15.89      | 82.96±12.19      | 0.005   |  |
| BP dias, mmHg                      | 45.52±14.07      | 43.35±8.98       | 0.460   |  |
| MAP, mm Hg                         | 60.95±13.36      | 56.42±9.30       | 0.080   |  |
| RPP, bpm×mmHg/10 <sup>4</sup>      | 0.82±0.21        | 0.64±0.16        | < 0.001 |  |
| Medication usage in HLH            | S patients       |                  |         |  |
| β-Blocker                          | 7 (5.8)          |                  |         |  |
| Aspirin                            | 82 (68.3)        |                  |         |  |
| Diuretics                          | 7 (5.8)          |                  |         |  |
| ACE inhibitor                      | 19 (15.8)        |                  |         |  |
| Marcumar                           | 16 (13.3)        |                  |         |  |
| HLHS-specific characteristics      |                  |                  |         |  |
| HLHS anatomy subtype               | es               |                  |         |  |
| MA/AA                              | 50 (42.0)        |                  |         |  |
| MA/AS                              | 7 (5.9)          |                  |         |  |
| MS/AA                              | 28 (23.5)        |                  |         |  |
| MS/AS                              | 34 (28.6)        |                  |         |  |
| Age at TCPC, y (media              | 2.59 (2.21-2.96) |                  |         |  |
| Total bypass time, min             | 394±106          |                  |         |  |
| Ischemia time, min                 | 69±51            |                  |         |  |
| Fenestration                       | 90 (75.6)        |                  |         |  |
| Fenestration closure               |                  |                  |         |  |
| Spontaneous                        | 6 (5.0)          |                  |         |  |
| Interventional                     | 23 (19.3)        |                  |         |  |
| Native ascending aorta             |                  |                  |         |  |
| Diameter, mm (echocard             | 3.4±1.3          |                  |         |  |
| Diameter, mm/m <sup>2</sup> (echoc | 16.1±6.5         |                  |         |  |
| Diameter, mm (date of              | 8.5±3.2          |                  |         |  |
| Diameter, mm/m <sup>2</sup> (date  | 10.8±4.2         |                  |         |  |
| Cardiac catheterization            |                  |                  |         |  |
| Mean aortic pressure, r            | 65±11            |                  |         |  |
| RV-EDP mean, mm Hg                 | 5.63±2.33        |                  |         |  |
| RV-EDP max, mm Hg                  | 8.65±2.84        |                  |         |  |
| Transpulmonary gradient            | 5.21±1.21        |                  |         |  |
| Mean pressure lateral to           | 12.87±2.19       |                  |         |  |

Continuous and normally distributed variables are summarized as mean±SD and otherwise as median (interquartile range). Counts are shown as number (%). *P*values are from the Wilcoxon test and Fisher exact test (sex). AA indicates aortic atresia; ACE, angiotensin-converting-enzyme; AP, atrial pressure; AS, aortic stenosis; BP dias, diastolic blood pressure; BP sys, systolic blood pressure; BSA, body surface area; CMR, cardiac magnetic resonance imaging; EDP, end-diastolic pressure; HLHS, hypoplastic left heart syndrome; IOR, interquartile range; MA, mitral atresia; MAP, mean arterial pressure; MS, mitral stenosis; RPP, rate-pressure product; RV, right ventricle; SVC, superior vena cava; and TCPC, total cavopulmonary connection.

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CMR parameter Control (n=34) HLHS (n=119) 75.97±18.26 53.24±13.19 EDV, mL/m<sup>2</sup> < 0.001 ESV, mL/m<sup>2</sup> 17.57±6.80 35.15±13.24 < 0.001 Wall mass, g/m<sup>2</sup> 49.75±13.24 67.28±19.71 < 0.001 ECV 0.26±0.02 0.34±0.08 0.001 Ejection fraction, % 67.62±6.40 54.37±8.83 <0.001 Cardiac index [mL/(min×m<sup>2</sup>)] 3.03±0.75 3.28±1.01 0.232 Rest MBF, mL/min per g 0.92±0.16 0.75±0.17 < 0.001 Rest MBF RPP-norm, mL/ 1.19±0.33 1.25±0.36 0.446 min per g Hyperemic MBF, mL/min 2.70±0.84 1.89±0.57 < 0.001 per g 118.31±23.71 93.46±15.38 0.001 Heart rate (hyperemia), bpm Perfusion reserve  $3.00 \pm 1.06$ 2.59±0.78 0.045 1.95±0.85 1.63±0.53 Perfusion reserve (rest MBF 0.165 normalized by RPP)

 Table 2.
 Comparison of CMR-Measured Indices in Patients

 With HLHS and Controls (LV)

Values for ventricular volumes and function pertain to the systemic ventricle in controls (LV) and HLHS patients (RV). *P* values are from the Wilcoxon or Fisher exact test. CMR indicates cardiac magnetic resonance; ECV, extracellular volume fraction; EDV, end-diastolic volume; ESV, end-systolic volume; HLHS, hypoplastic left heart syndrome; LV, left ventricle; MBF, myocardial blood flow; norm, normalized; RPP, rate-pressure product; and RV, right ventricle.

image artifacts due to metallic implants, foldover artifacts not apparent before contrast enhancement, or because the patient refused contrast or sedation wear off and the patient became restless. At the segmental level, MBF at rest was quantified in 86% (744 of

864) of all segments and hyperemic blood flow in 84% (680 of 808) of all segments. Segmental could not be quantified due to spillover from the blood pool around the RV outflow tract because the myocardial wall was too thin or due to the presence of image artifacts. MBF of the systemic ventricle at rest, normalized by the RPP, did not significantly differ between HLHS patients and control subjects (rest MBF, 1.25±0.36 versus 1.19±0.33 mL/min per g; P=0.446), but during maximal hyperemia, HLHS patients had MBFs significantly lower than controls (hyperemic MBF, 1.89±0.57 versus 2.70±0.84 mL/min per g; P<0.001), as shown in Figure 3. Myocardial perfusion reserve, calculated as hyperemic MBF, divided by rest MBF was also lower in HLHS patients compared with controls (2.59±0.78) versus 3.00±1.06; P=0.045), but once rest MBF was normalized by the RPP, the difference of the reserve was insignificant (P=0.165; Table 2).

Hyperemic MBF varied among anatomic subtypes (P=0.058 for ANOVA test) and was significantly lower by 0.33 mL/min per gram in the MS/AA group compared with the MA/AA group (adjusted P=0.017; Figure 4) and with respect to all other anatomic subtypes pooled together (P=0.019 for *t* test). VCCs, not uncommon in the MS/AA subtypes,<sup>13,25</sup> were found in 8 (29%) HLHS patients with the MS/AA subtype. The presence of VCCs had no significant effect on myocardial perfusion at rest or hyperemia. Late gadolinium enhancement in HLHS was associated with a reduced rest MBF (RPP normalized; P=0.040) and hyperemic



### Figure 3. Myocardial blood flow in hypoplastic left heart syndrome (HLHS) and controls at rest and hyperemia.

Normalized at rest myocardial blood flow did not differ between HLHS and controls (**left**), since the rate-pressure product was significantly different between the two groups ( $0.82\pm0.21 \text{ vs} 0.64\pm0.16 \text{ bpm mm} \text{Hg}/10^4$ ; *P*<0.001). Myocardial blood flow at rest without normalization by the rate-pressure product (not shown) was significantly lower in the hypoplastic left heart syndrome group due to the lower cardiac workload. Hyperemic myocardial blood flow was significantly lower in patients with HLHS compared with healthy controls. *P* values are from unpaired Student *t* tests.



**Figure 4. Hyperemic myocardial blood flow (MBF) in anatomic subtypes of hypoplastic left heart syndrome (HLHS).** Hyperemic MBF was the lowest in the anatomic subtype of HLHS with mitral stenosis (MS) and aortic atresia (AA; adjusted *P*=0.017 compared with mitral atresia [MA]/AA). *P* values are from Student *t* test and adjusted for multiple comparisons. AS indicates aortic stenosis.

MBF (P=0.042; Figure 5A and 5B). Hyperemic MBF was lower in the posterior wall segment when a rudimentary LV cavum was present (P=0.016; Figure III in the Data Supplement).

In HLHS patients, the coronary conductance at rest, calculated as rest MBF, divided by the mean aortic pressure measured during catheterization within a day of the CMR exam, averaged  $0.018\pm0.0031$  mL/min per g/mmHg. The coronary conductance at rest was negatively associated with blood O<sub>2</sub> saturation (r, -0.3; *P*=0.017; Figure 6A). The blood O<sub>2</sub> saturation correlated negatively with the RV cardiac index (r, -0.34; *P*<0.001; Figure 6B) and was negatively associated with blood

hematocrit (r, 0.22; *P*=0.017) and blood hemoglobin (r, 0.28; *P*=0.003).

### **Cardiac Catheterization**

Invasive hemodynamic measurement results are shown in Table 1. Blood pressures (mm Hg) within the lateral tunnel, RV, and aorta were in the similar range/age previously reported for children with HLHS in Fontan circulation from our center and others.<sup>36,37</sup> There were no significant pressure differences among anatomic HLHS subtypes. Of the 99 patients who underwent coronary catheterization, 12



Figure 5. Myocardial blood flow (MBF) at rest and hyperemia in hypoplastic left heart syndrome (HLHS) with or without late gadolinium enhancement (LGE).

**A**, Myocardial LGE, which is relatively infrequent in the HLHS cohort (14%), was associated with a lower MBF at rest (P=0.040). **B**, LGE was also associated with a lower hyperemic MBF (P=0.042). AA indicates aortic atresia; MS, mitral stenosis; and RPP, rate-pressure product.



**Figure 6.** Association of coronary conductance and right ventricular (RV) cardiac index with blood oxygen saturation. **A**, Coronary conductance was negatively associated with blood  $O_2$  saturation, suggesting a form of compensatory vasodilation at baseline. Coronary conductance was estimated from myocardial blood flow at rest (mL/min per g), divided by the mean aortic pressure (mmHg) as measured during catheterization within a day of the cardiac magnetic resonance study. **B**, The RV cardiac index decreased with blood  $O_2$ saturation (both *P*<0.001). The continuous lines in the graph were calculated with linear models for the variable on the *y* axis and  $O_2$  saturation (*P*=0.017) as predictor.

(12%) had diastolic dysfunction (RV-end-diastolic pressure, >10 mmHg). Diastolic dysfunction was present in 20% (5 of 25) of HLHS patients in the MS/AA subgroup and in 9% (7 of 74) for the other subgroups (P=0.173 for Fisher exact test).

Diastolic dysfunction was associated with a lower hyperemic MBF ( $1.54\pm0.38$  versus  $1.95\pm0.58$  mL/min per g; *t* test *P*=0.0053).

### Multivariate Prediction of Hyperemic MBF

After stepwise selection of predictors for multivariate regression model for the prediction of hyperemic MBF in HLHS, the MS/AA subtype (-0.329 mL/min per g; P=0.034), age at TCPC (-0.179 per year delay in TCPC; P=0.022), RV diastolic dysfunction (-0.316 mL/min per g; P=0.088), and male gender (P=0.171) remained in the final model. This selection of predictors did not change if LGE presence was replaced by the percentage of LGE in RV. Age at CMR, O<sub>2</sub> saturation, a rudimentary LV cavum, and myocardial LGE were eliminated. The model intercept estimate of 2.1 mL/min per gram corresponds to the hyperemic blood flow of a female HLHS patient with TCPC at 2.7 years, in the MA/AA anatomic subtype and without RV diastolic dysfunction (Figure IV in the Data Supplement).

### **Extracellular Volume Fraction**

T1 mapping pre- and post-contrast were performed in 51 (43%) HLHS patients after a look-locker T1-mapping technique became available. In HLHS patients, the RV ejection fraction trended lower with increasing extracellular volume (P=0.087). There was no significant correlation of extracellular volume fraction with rest or hyperemic MBF, respectively.

## DISCUSSION

This prospective study in a cohort of 119 HLHS patients in Fontan circulation is the first CMR-based investigation focusing on the pathophysiological characteristics that put the RV microcirculation at risk after 3-step surgical palliation. We found significant differences of the microcirculation of the systemic ventricle in HLHS patients compared with control subjects.

### Effects of HLHS Subtypes on MBF

The MS/AA subtype represents a risk factor unique to HLHS patients,213,35 starting with the Norwood procedure, during the interstage phase and beyond.<sup>2,13,35,38</sup> Over 2 decades ago, Sugiyama et al<sup>39</sup> observed in an angiographic and histopathologic study that myocardial fibrosis, calcification, and necrosis occur more frequently in HLHS patients with MS/AA anatomy, compared with other subtypes. They also reported an impaired RV posterior wall motion, concluding that the rudimentary LV (piggyback ventricle) is a functional burden to the heart. Siehr et al<sup>13</sup> noted that severe myocardial dysfunction, mostly due to coronary ischemia, is possibly related to VCCs and renders the MS/AA group more vulnerable to surgical palliation or cardiopulmonary bypass. Our data indicate that myocardial perfusion continues to be impaired in the MS/AA group after switch to Fontan circulation compared with the other anatomic subgroups, but we could not find a significant difference in patients with and without VCC's.

The reduction of hyperemic MBF related to RV diastolic dysfunction in HLHS patients parallels previous findings on the effects of LV diastolic dysfunction on microcirculary function in anatomically normal hearts with angiographically normal coronary arteries.<sup>40</sup> A rudimentary LV has been characterized as representing a functional burden to the RV, manifested by diastolic dysfunction.<sup>12,41</sup> In our study, diastolic dysfunction was about twice as frequent in the MS/AA group (20%) with rudimentary LV compared with the other subgroups (9%).

### LGE in HLHS

LGE presence in HLHS patients was associated with lower resting and hyperemic blood flows. Nevertheless, LGE did not make it into the most parsimonious model for hyperemic MBF after stepwise variable selection (Figure III in the Data Supplement), while age at the time of TCPC was retained in the final model. The presence of LGE was strongly associated with the age at TCPC completion, suggesting that LGE may partially result from later conversion to Fontan circulation (Figure I in the Data Supplement) and be a secondary effect of a later relief from cyanosis. Age at TCPC-the variable that is retained after stepwise selection-may capture multiple adverse effects to the heart besides LGE, for example, the RV end-diastolic volumen indexed by body surface area and RV end-systolic volumen indexed by body surface area increased significantly with age at TCPC, and RV ejection fraction decreased with age at TCPC independent of the age at the time of CMR. Therefore, the lateral tunnel may be advantageous to achieve an early relief of cyanosis. The current study did not show any significant association between a CMR marker of diffuse fibrosis (extracellular volume fraction) and myocardial perfusion. Further longitudinal and comparative studies need to ultimately clarify the effect of type and timing of TCPC on myocardial fibrosis and blood flow.

### **Coronary Conductance**

There was a significant association between coronary conductance at rest and blood  $O_{2}$  saturation (Figure 6A) in HLHS patients, suggesting a form of compensatory vasodilation at rest. More specifically, patients with lower oxygen saturation had a higher coronary conductance<sup>32,42</sup> estimated from the ratio of MBF at rest divided by the mean aortic pressure (Figure 6A). HLHS patients with lower oxygen saturation tended to have higher blood hematocrit and hemoglobin values, both of which may contribute to higher blood viscosity with decreasing O<sub>o</sub> saturation.43,44 Maintaining RV myocardial perfusion in the presence of increased blood viscosity requires a compensatory increase of perfusion pressure and coronary conductance. As blood pressures are already lower than normal in many HLHS patients, compensatory coronary vasodilation at baseline appears to be the more likely response to the reduced  $O_{\circ}$  saturation (Figure 6A). Lower O<sub>o</sub> saturations can reflect a higher right-to-left shunt (eg, through an open fenestration, suture dehiscence of the baffle, or veno-venous collaterals). This indicates that blood bypasses the lungs, which is compensated by an increase of the cardiac index and would

be consistent with the observed correlation between  $O_2$  saturation and cardiac index (Figure 6B).

There is a paucity of published studies on myocardial perfusion in children with HLHS. A study by Donnelly et al<sup>8</sup> in 1998 with ammonium positron emission tomography in a subgroup of 5 HLHS patients at a mean interval of 13.2 days after Norwood surgery, but none of them in Fontan circulation, is a notable exception.

Previous nuclear medicine studies in children with healthy hearts have shown that myocardial perfusion under stress increases  $\approx$ 3-fold compared with resting perfusion.<sup>16</sup> The results in our control group approximately correspond to the MBF ranges at rest and hyperemia reported for children<sup>21</sup> and young adults.<sup>14,15,28,30,45</sup>

# Possible Mechanisms for Impaired MBF in HLHS

Myocardial perfusion depends on the capillarization of the ventricular myocardium, which may be reduced in HLHS patients, according to a previous histological investigation by Salih et al.<sup>10</sup> An altered or impaired vasoreactivity due to inadequate cardiac (re-)innervation may be another mechanism that reduces MBF under stress in these patients and possibly also affects coronary vasomotor tone at rest. The Norwood operation involves surgical sectioning of the native aorta resulting in sympathetic denervation and potential inadequate cardiac reinnervation similar to transplanted hearts.<sup>6,46-48</sup> Extensive aortic surgery during arterial switch operation 14,49 or heart transplantation<sup>6,46-48</sup> is also known to impair the coronary vasodilator response. Cardiac efferent sympathetic signals affect MBF during a stress,<sup>50</sup> especially at the arteriolar level.

In HLHS patients with no rudimentary LV, no significant differences in hyperemic MBF between RV wall segments were observed. In contrast, patients with rudimentary LV had lower hyperemic MBF in the posterior segment (Figure II in the Data Supplement), which is the segment shared with the rudimentary LV, if present. A rudimentary LV may induce RV diastolic dysfunction in HLHS patients while the impaired diastolic relaxation is known to reduce the coronary vasodilator response.<sup>12,36,40</sup>

### Limitations

Some study limitations should be considered for interpretation of the results. Unfortunately, catheter-based measurements within a day of CMR study were not feasible for all patients. The MA/AS subtype of HLHS is known to be a rare phenotype, and with only 7 patients in our cohort, we had to exclude them from the analysis that included anatomic subtypes as effect.

Techniques for myocardial T1 mapping and extracellular volume fraction quantification only became available on our scanner during the study. Since some patients had a short sedation, LGE imaging and the T1 mapping usually performed at the end of the CMR protocol were not carried out in all cases. Therefore, only a subgroup of HLHS patients underwent T1 mapping (43%) or LGE imaging (87%).

### Conclusions

This study investigating a cohort of 119 HLHS patients in Fontan circulation is, to the best of our knowledge, the first CMR-based investigation focusing on the pathophysiological characteristics that predispose RV microcirculation to dysfunction after 3-step surgical palliation. Compared with control subjects, CMR-based quantification of MBF in the HLHS patients' systemic ventricle revealed a significant impairment of the coronary autoregulation after complete surgical palliation. Diastolic dysfunction, the presence of the MA/AS subtype, and an older age at TCPC appear to be risk factors for RV microcirculatory dysfunction. Accordingly, the associations reported in the present article can be viewed as correlations that elucidate how known pathological features of HLHS are related to microvascular dysfunction. Microvascular dysfunction is an important factor in heart failure with reduced ejection fraction, and there is also growing evidence of its role in the etiology of heart failure with preserved ejection fraction.51,52

In summary, this study provides new insights into factors unique to HLHS patients that affect the RV microcirculation and tissue structure and may generate novel hypothesis for future long-term studies. Such studies are required to investigate the role of the RV microcirculation leading to RV dysfunction in HLHS patients over time.

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None.

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