The natural and unnatural history of the systemic right ventricle in adult survivors

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Objective: The study objective was to evaluate long-term trends in morbidity and mortality in a national cohort of adult patients with a systemic right ventricle due to the atrial switch for transposition of the great arteries or congenitally corrected transposition of the great arteries.

Methods: We performed a retrospective cohort study from a baseline of 18 years, including life table and Kaplan–Meier analysis for probability of death/transplant, arrhythmia, surgical or percutaneous intervention, and permanent pacemaker insertion.

Results: A total of 97 adults with transposition of the great arteries–atrial switch (Mustard procedure in 80/Senning procedure in 17) and 32 adults with congenitally corrected transposition of the great arteries survived. The median ages at latest follow-up were 29 and 34 years, respectively. At 40 years of follow-up, freedom from death or transplant was 0.90 for those with transposition of the great arteries–atrial switch and 0.84 for those with congenitally corrected transposition of the great arteries ($P = .833$). Freedom from arrhythmia at 40 years of follow-up was 0.51 for those with transposition of the great arteries–atrial switch and 0.93 for those with congenitally corrected transposition of the great arteries ($P = .007$). Freedom from intervention at 40 years of follow-up was 0.33 for those with transposition of the great arteries–atrial switch after initial repair and 0.53 for those with congenitally corrected transposition of the great arteries ($P = .938$). Freedom from pacemaker insertion at 40 years of follow-up was 0.77 for those with transposition of the great arteries–atrial switch and 0.62 for those with congenitally corrected transposition of the great arteries ($P = .161$).

Conclusions: Those patients who survive to adulthood with a systemic right ventricle experience low mortality and good functional status up to 40 years of age. However, there is a substantial burden of atrial tachyarrhythmia, and this occurs significantly earlier in those with transposition of the great arteries–atrial switch. Management of atrial tachyarrhythmia, along with systemic right ventricular dysfunction and systemic atrioventricular valve regurgitation, is likely to be the major challenge for this group of patients over the next decade. (J Thorac Cardiovasc Surg 2013;145:1493-503)

The morphologic right ventricle (RV) supports the systemic circulation in those individuals with congenitally corrected transposition of the great arteries (ccTGA) and those who have survived atrial switch surgery (the Mustard or Senning procedure) for complete transposition of the great arteries (TGA). Although atrial switch is no longer the gold standard for management of TGA, having been superseded by the arterial switch operation in the late 1980s, many adult patients with TGA will have undergone the older procedure.

Patients with either diagnosis are at risk of a host of complications associated with the systemic RV (SRV), including premature heart failure, regurgitation through the systemic atrioventricular (AV) valve, AV nodal block and a need for permanent pacing, tachyarrhythmia, and sudden cardiac death.

The long-term history of patients with an SRV still needs to be clearly defined. Most of the cohort studies of patients after atrial switch focus on establishing which of the Mustard and Senning procedures was the best procedure to use, and the majority provide follow-up to only 20 years. Patients with ccTGA are the subject of only a handful of cohort studies. Our aim was to establish the natural history of the SRV in both conditions and provide a benchmark to which other treatment approaches for the SRV (eg, placing the morphologic left ventricle [LV] in the systemic circulation) can be compared.

We focused on the group of patients who have survived to at least the age of 18 years, because although this group of patients is of great interest to clinicians engaged in provision of an adult congenital heart disease service, to our...
Abbreviations and Acronyms

AV = atrioventricular
cCTGA = congenitally corrected transposition of the great arteries
LV = left ventricle
MRI = magnetic resonance imaging
PA = pulmonary artery
RV = right ventricle
SRV = systemic right ventricle
TGA = transposition of the great arteries
TR = tricuspid regurgitation
VSD = ventricular septal defect

Knowledge there are no studies that assess the prognosis for these survivors.

MATERIALS AND METHODS

We performed a retrospective cohort study, with approval from the regional research ethics committee. We identified 204 patients with TGA-atrial switch or cCTGA (excluding those who had undergone single-ventricle palliation) who had attended the national pediatric or adult congenital cardiac services for Scotland. From this group of patients, we focused on the subset of adult patients (aged at least 18 years) who had an SRV. This resulted in a study population of 129 individuals. Dropout was due to patient death or transplant before the age of 18 years (31 patients), conversion to a systemic LV before the age of 18 years (5 patients), loss of follow-up (16 patients), or age less than 18 years at the time of data collection (23 patients).

Data were collected and anonymized for each individual from our electronic database and hospital case notes between January 2011 and 2012. Patient-specific timelines allowed calculation of survival curves for mortality, atrial arrhythmia, surgical or catheter intervention, and pacemaker insertion. Follow-up was from the age of 18 years. Patients were censored at the time of their last clinic review, date of death, date of cardiac transplant, or date of surgery to restore the morphologic LV to the systemic circulation (a Senning–Rastelli or arterial switch operation).

During long-term follow-up, patients underwent regular clinical review that included establishment of New York Heart Association status and ascertainment of occurrence of cardiovascular events, such as arrhythmia, hospital admission, and intervention.

Tachyarrhythmia was defined as any supraventricular or ventricular tachyarrhythmia captured on ambulatory or 12-lead electrocardiograph recording that was clinically significant (ie, caused symptoms or required treatment). We excluded cases in which tachyarrhythmia occurred exclusively within 30 days of cardiac surgery. Reintervention was defined as any surgical or percutaneous cardiac procedure occurring after baseline. Pacemaker insertion was defined as any permanent epicardial or endocardial pacemaker, and included those with cardiac resynchronization therapy.

Regular imaging with transthoracic echocardiography and, when no contraindications existed, cardiac magnetic resonance imaging (MRI) was performed (Table 1). This was used to establish the degree of tricuspid regurgitation (TR) and SRV dysfunction. A total of 50 patients underwent formal cardiopulmonary exercise testing, and peak oxygen uptake (measured in mL·kg⁻¹·min⁻¹) was recorded if the test was maximal (Table 1). Since 2010, cardiac MRI and cardiopulmonary exercise test have been performed routinely on both groups of patients regardless of functional status, or presumed need for surgery to clarify anatomy and provide more accurate assessment of ventricular function, and we believe the values obtained from these data provide an accurate cross-sectional assessment of the cohort at the time of data collection.

Data were described as frequencies, means and standard error, and medians and interquartile range. Means were compared with the independent samples t test, and medians were compared with the independent samples median test. Chi-square test using linear-by-linear association was used to assess the relationship between severity of TR and severity of SRV impairment.

Cumulative probability of survival was estimated using the Kaplan–Meier and life table method, and differences between groups were evaluated via the log-rank test. We performed analysis for survival from death or transplant, atrial tachyarrhythmia, surgical intervention after baseline, and pacemaker therapy.

Cox proportional hazards models were used to identify predictors for death or cardiac transplant. All analyses were performed using PASW Statistics v18.0.3 (IBM corporation, New York, NY).

RESULTS

We first identified the 204 patients in the pediatric population (Figure 2). Patients were censored at the time of their last review, when they died, when they received a cardiac transplant, or when they underwent surgery that restored the morphologic LV to the systemic circulation and the RV to the pulmonary circulation (ie, a double switch or Senning–Rastelli). Although a clear survival advantage up to 10 years was experienced by those with TGA-atrial switch, mortality between the 2 cohorts was remarkably similar after this time (log-rank P = .657). Of 133 patients with TGA-atrial switch (89 male, 44 female), 88 had ongoing follow-up, 23 died, 19 were lost to follow-up, 2 underwent cardiac transplantation, and 1 underwent a subsequent arterial switch procedure. Of 71 patients with cCTGA (46 male, 25 female), 49 had ongoing follow-up, 14 died, 3 were lost to follow-up, 1 underwent cardiac transplantation, 1 underwent an atrial and arterial double switch operation, and 3 underwent Senning–Rastelli repair.

The red line indicates the period from 18 years of follow-up and the major area of interest for this study. There were 97 adult survivors of atrial switch surgery for TGA and 32 adult survivors of cCTGA. The characteristics are outlined in Table 1.

Most of the patients undergoing atrial switch had undergone a Mustard repair, with only 17% undergoing the Senning procedure. More coexistent lesions were present in those patients with a diagnosis of cCTGA. By the end of the follow-up period, the majority of patients had good functional status, normal or only mild impairment of SRV function, and no or mild TR, with only 2 patients with cCTGA proceeding to tricuspid valve replacement and only 3 patients with severe regurgitation.

We assessed the relationship between SRV impairment and severity of TR. The 2 patients who underwent tricuspid valve replacement were excluded from the analysis. The chi-square value for linear-by-linear association between SRV impairment and severity of TR was 10.064.
<table>
<thead>
<tr>
<th>TGA-atrial switch</th>
<th>ccTGA</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>N</strong></td>
<td>97</td>
<td>32</td>
</tr>
<tr>
<td><strong>Male</strong></td>
<td>64</td>
<td>20</td>
</tr>
<tr>
<td><strong>Female</strong></td>
<td>33</td>
<td>12</td>
</tr>
<tr>
<td><strong>Age (y)</strong></td>
<td>Median (IQR) 29 (25-32)</td>
<td>33 (23-46)</td>
</tr>
<tr>
<td>Follow-up from 18 y</td>
<td>Median (IQR) 10.8 (6.5-13.9)</td>
<td>13.2 (5.1-28.2)</td>
</tr>
<tr>
<td><strong>Atrial switch operation (%)</strong></td>
<td>Mustard 80 (83)</td>
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</tr>
<tr>
<td></td>
<td>Senning 17 (17)</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Coexistent congenital lesions</strong></td>
<td>Dextrocardia 0</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>VSD 10</td>
<td>20</td>
</tr>
<tr>
<td></td>
<td>Pulmonary stenosis 13</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>Pulmonary atresia 0</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Ebstein 0</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Coarctation 3</td>
<td>0</td>
</tr>
<tr>
<td><strong>Outcome</strong></td>
<td>Ongoing follow-up 86</td>
<td>28</td>
</tr>
<tr>
<td></td>
<td>Lost to follow-up 5</td>
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<tr>
<td></td>
<td>Died 5</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Transplant 1</td>
<td>1</td>
</tr>
<tr>
<td><strong>Cause of death</strong></td>
<td>Heart failure 1</td>
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</tr>
<tr>
<td></td>
<td>Sudden 3</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Sepsis 1</td>
<td>0</td>
</tr>
<tr>
<td><strong>Tachyarrhythmia</strong></td>
<td>No 72</td>
<td>26</td>
</tr>
<tr>
<td></td>
<td>Yes 21</td>
<td>4</td>
</tr>
<tr>
<td><strong>Up-to-date imaging (%)</strong></td>
<td>Echocardiography 73 (75)</td>
<td>24 (73)</td>
</tr>
<tr>
<td></td>
<td>MRI 22 (22)</td>
<td>5 (16)</td>
</tr>
<tr>
<td><strong>SRV impairment</strong></td>
<td>Nil 26</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>Mild 32</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>Moderate 18</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>Severe 6</td>
<td>1</td>
</tr>
<tr>
<td><strong>Cardiac MRI</strong> (mean ± SE)</td>
<td>SRV EDV 111 ± 8.8</td>
<td>127 ± 18.3</td>
</tr>
<tr>
<td></td>
<td>SRV EF % 54 ± 2.3</td>
<td>61 ± 2.6</td>
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<tr>
<td></td>
<td>LVEDV 79 ± 6.0</td>
<td>92 ± 11.0</td>
</tr>
<tr>
<td></td>
<td>LVEF % 60 ± 2.6</td>
<td>66 ± 3.3</td>
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<tr>
<td><strong>TR</strong></td>
<td>Nil 42</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Mild 40</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>Moderate 13</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>Severe 0</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Replaced 0</td>
<td>2</td>
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</table>

(Continued)
(P = .002), implying a statistically significant correlation between the 2 variables.

More detailed assessment of ventricular function and functional capacity was performed by means of cardiac MRI and cardiopulmonary exercise testing, because this is a routine form of evaluation for all eligible patients at this center with an SRV, regardless of clinical status. From cardiac MRI in 27 patients, the mean ejection fraction of the SRV was lower in those with TGA-atrial switch than in those with ccTGA (54% vs 61%), although this was not significant. There was no clear relationship between time from baseline and ventricular volumes or ejection fraction. Mean peak oxygen uptake for 50 patients in both groups was 25 mL/kg/min. There was no statistically significant difference in peak oxygen uptake, heart rate reserve, and minute ventilation to carbon dioxide output ratio between the 2 cohorts.

Intervention was common. Baffle problems accounted for most of this in the atrial switch group, with 26 patients (26.8% of the adult TGA-atrial switch cohort) requiring further intervention. Of this group of patients, 24 (of a possible 80) had undergone a Mustard repair, compared with 2 patients undergoing a Senning (of a possible 17). The majority of intervention was performed percutaneously, and baffle obstruction was a more frequent indication for intervention than a leak. In the majority of cases, the initial intervention was successful at treating the baffle leak or stenosis, with only 6 patients (Mustard in 5, Senning in 1) requiring more than 1 repeat intervention. In 2 patients (Senning repair), partially successful percutaneous intervention for a baffle leak was followed by definitive surgical treatment 2 years later. One patient (who underwent a Mustard repair) required 2 surgical baffle repairs, albeit 14 years apart, for obstruction of the pulmonary venous pathway. Two patients each required 2 attempts at catheter

**FIGURE 1.** Coexistent lesions and initial surgery in 32 patients with ccTGA. *ccTGA*, Congenitally corrected transposition of the great arteries; *VSD*, ventricular septal defect; *LVOTO*, left ventricular outflow tract obstruction; *LV*, left ventricle; *PA*, pulmonary artery; *TVR*, tricuspid valve replacement.

**FIGURE 2.** Long-term survival in 133 patients after atrial switch surgery for TGA and 71 patients with ccTGA. The red line indicates the period from 18 years of follow-up, and the major area of interest for this study. *ccTGA*, Congenitally corrected transposition of the great arteries.
intervention, and the remaining 2 patients required percutaneous intervention after initial surgical revision of the baffles.

A total of 13 of 97 patients with TGA-atrial switch underwent subsequent pulmonary artery (PA) banding for treatment of SRV dysfunction, 1 of whom later died suddenly 3 years post-banding (presumed arrhythmic in cause). Up-to-date imaging was not available for 1 other patient who underwent this procedure. For the remaining 11 patients, the median follow-up period was 5.8 years. The mean gradient through the PA band was 3.4 m/s. SRV function was normal or mildly impaired in 9 patients, moderately impaired in 1 patient, and severely impaired in 1 patient. There was no or mild systemic TR in 6 of 11 patients, with 4 patients having moderate regurgitation and 1 patient having severe regurgitation. None of the patients with PA banding were ever deemed suitable for a subsequent double-switch.

Other indications for reintervention in the TGA-atrial switch group were enlargement of the subpulmonary LV outflow tract, repair of aortic coarctation, and cardiac transplant. No TGA-atrial switch cases have yet been referred for tricuspid valve replacement.

In the ccTGA group, most intervention was directed at correction of underlying anatomic lesions, such as a ventricular septal defect (VSD) (Figure 1), although 1 patient with preexisting left ventricular outflow tract obstruction and a VSD required PA banding for a failing SRV, and 1 patient required tricuspid valve replacement. Both patients were doing well at their most recent follow-up. Although a number of pediatric patients have ultimately undergone a double switch or Senning–Rastelli, none of the adult survivors have.

Cumulative survival from each adverse outcome for the 129 patients who reached the age of 18 years is outlined in detail in Figure 3. The dropoff at baseline (age 18 years) reflects the number of adverse events that had occurred in individuals before the age of 18 years.

There was no significant difference in survival between patients with TGA-atrial switch and patients with ccTGA (log-rank test \( P = .833 \)). Cumulative survival (including cardiac transplantation) was 0.99 ± 0.01, 0.90 ± 0.04, and 0.90 ± 0.04 at 20, 30, and 40 years post-atrial switch, respectively, and 0.97 ± 0.03, 0.91 ± 0.06, and 0.84 ± 0.09 at 20, 30, and 40 years post-ccTGA, respectively. Although there seemed to be an abrupt decrease in survival to approximately 0.40 at more than 45 years from baseline for the TGA-atrial switch cohort, only 2 patients had been followed for this length of time; therefore, this value was not thought to be sufficiently representative of the whole cohort. There was no significant difference in survival from 18 years of follow-up between Mustard and Senning cases (log-rank test \( P = .625 \)), although after 27 years of follow-up graphic trends favored those undergoing the Senning. There was no significant difference between simple and complex TGA-atrial switch (\( P = .562 \)). There was no significant difference in survival between simple and complex ccTGA after 18 years of follow-up (log-rank test \( P = .662 \)). There was no significant difference in survival between the 14 patients in both groups who underwent PA banding and those who did not (\( P = .659 \)). A univariate analysis with a Cox proportional hazards model was performed to establish risk factors for death or transplant (Table 2). Severe SRV impairment at baseline was most strongly associated with risk of death or transplant, albeit with wide confidence intervals.

Patients with ccTGA demonstrate low arrhythmia burden with cumulative survival from tachyarrhythmia of 1.00, 1.00, and 0.93 ± 0.07 at 20, 30, and 40 years, respectively. This was significantly better than in those who received atrial switch, who experienced survival of 0.91 ± 0.03, 0.73 ± 0.06, and 0.54 ± 0.14 at 20, 30, and 40 years, respectively (log-rank test \( P = 0.007 \)). All documented tachyarrhythmias were supraventricular: There were no instances of documented ventricular tachycardia causing symptoms or requiring treatment (although the sudden deaths that occurred may have been due to this).

Freedom from intervention at 20, 30, and 40 years of follow-up was 0.74 ± 0.05, 0.57 ± 0.06, and 0.33 ± 0.08 for TGA-atrial switch, respectively, and 0.57 ± 0.09, 0.53 ± 0.10, and 0.53 ± 0.10 for ccTGA, respectively (log-rank test \( P = .938 \)).

There was no significant difference between the 2 groups for cumulative survival from pacemaker insertion (log-rank test \( P = .161 \)), although graphic trends favor the TGA-atrial switch group. Freedom from pacemaker insertion at 20, 30, and 40 years of follow-up was 0.87 ± 0.04, 0.82 ± 0.04, and 0.77 ± 0.06 for TGA-atrial switch, respectively, and 0.79 ± 0.08, 0.69 ± 0.09, and 0.62 ± 0.11 for ccTGA, respectively.

DISCUSSION

Mortality

Overall mortality to 40 years is low in both cohorts, with the cumulative probability of survival estimated at 90% survival after atrial switch and 84% after ccTGA. Of the 7 deaths after 18 years of follow-up, 4 were sudden and 2 were secondary to heart failure (1 patient with ccTGA died after referral for cardiac transplant assessment). This is similar to other studies. Sudden death and heart failure accounted for the majority of late deaths in 2 large retrospective studies of a combined total of approximately 700 atrial switch cases\(^1\)\(^-\)\(^2\) and in a large retrospective study of survivors of surgery for ccTGA.\(^5\) Therefore, strategies to identify those at risk of sudden cardiac death and to manage systemic RV dysfunction will be crucial to improve outcomes in these patients.

Direct comparison between our cohort of adult survivors and the larger cohort studies of SRV is difficult because most studies include early postoperative and pediatric
deaths. We therefore included our pediatric data, and with
this cumulative mortality was broadly similar (Table 3). It
remains to be seen whether the adaptation of an early ag-
gressive strategy for patients with ccTGA, by which the
morphologic LV is connected to the systemic circulation,
results in improved survival into adulthood. However,
Brawn4 reports survival of 90% at up to 9 years in a cohort
of 56 patients with ccTGA who underwent both types of
double switch (atrial and arterial switch, and Senning–Ras-
telli), with an early mortality of 5.6%.4 Ly and colleagues5
reported 100% survival at 60 months in 20 patients with
complex ccTGA who underwent the Senning–Rastelli pro-
procedure. Shin’oka and colleagues6 reported 74.5% survival
at 15 years in 15 patients undergoing atrial-arterial switch
and 80% survival at 16 years in 69 patients undergoing
the Senning–Rastelli procedure. Poorer survival was re-
ported in the Senning–Rastelli group compared with the
atrial-arterial switch group in another study with data up
to 10 years of 65 patients undergoing anatomic repair.7
However, this is in contrast to a 2006 meta-analysis of

FIGURE 3. Mortality and morbidity in 129 adult survivors of ccTGA and atrial switch surgery for TGA. ccTGA, Congenitally corrected transposition of the
great arteries.

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124 patients undergoing anatomic repair for ccTGA in which the Senning–Rastelli procedure conferred a survival benefit at least for in-hospital mortality; however, there were no data regarding longer-term outcomes.8 Although these results are clearly encouraging, the prognosis to adulthood and beyond remains to be established.

**Morbidity**

**Tachyarrhythmia.** Atrial tachyarrhythmia was common in patients with an SRV. However, it occurred significantly earlier in survivors of atrial switch surgery \( P = .007 \). This may be due to the extensive atrial scar tissue and potential for macro-reentry circuits. The relatively high prevalence of arrhythmia in both cohorts at 40 years is cause for concern, because its management is extremely challenging even for experienced electrophysiologic physicians.

**Systemic right ventricle impairment and tricuspid regurgitation.** At the time this study was performed, the majority of patients in both cohorts exhibited insignificant or mild SRV impairment and TR. However, severe SRV impairment was a significant risk factor for death, and management is controversial. We confirmed a clear correlation between severity of SRV impairment and severity of TR, in keeping with the published literature.9,10 Cardiac MRI data suggested that overall mean SRV function for the group was good in ccTGA cases, with an SRV ejection fraction of 61%, and only mildly impaired in atrial switch cases, with an SRV ejection fraction of 54% \( (P = .190) \). Of note, mean SRV end-diastolic volume was slightly higher in the ccTGA cases; however, the difference was not statistically significant.

Approaches to management of SRV failure are controversial. Early intervention for a regurgitant systemic tricuspid valve has been advocated in view of its correlation with SRV failure. However, results are poor in patients who have preexisting SRV impairment; therefore, this is a prophylactic strategy rather than an option for patients who already have a failing SRV. Furthermore, long-term prognostic data are lacking, and there are no large-scale case-control studies.10-12 Generally, tricuspid valve replacement rather than repair is advocated if the RV is planned to remain in the systemic circulation.10,13

The atrial-arterial double switch or Senning–Rastelli procedures, if performed early enough in a center with sufficient expertise, eliminate the problem of SRV dysfunction and TR. However, results are unfavorable in adult patients.4,14,15 Furthermore, there is a risk of a range of new complications, including dysfunction of the newly systemic LV and insufficiency through the neoaoartic valve (in the case of atrial-arterial double switch).4

The process of training the LV through the placement of a PA band is used as a treatment in its own right, because higher LV pressures avoid septal dyskinesis exacerbating SRV impairment, and in theory helps to prevent tricuspid annular dilatation, thus lessening the severity of TR. In the small subcohort of our group of patients undergoing PA banding, most maintained normal or only mildly impaired SRV function. However, there was no overall

### TABLE 2. Hazard ratios for death or cardiac transplant in adult survivors with a systemic right ventricle

<table>
<thead>
<tr>
<th>Variable</th>
<th>HR</th>
<th>P value</th>
<th>95% CI</th>
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</thead>
<tbody>
<tr>
<td>Male</td>
<td>2.286</td>
<td>.303</td>
<td>0.474-11.033</td>
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<tr>
<td>Initial VSD</td>
<td>2.370</td>
<td>.208</td>
<td>0.619-9.081</td>
</tr>
<tr>
<td>Initial BT shunt</td>
<td>1.537</td>
<td>.692</td>
<td>0.183-12.89</td>
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<tr>
<td>Tachyarrhythmia</td>
<td>0.046</td>
<td>.717</td>
<td>0-∞</td>
</tr>
<tr>
<td>Pacemaker</td>
<td>2.349</td>
<td>.290</td>
<td>0.483-11.417</td>
</tr>
<tr>
<td>Severe TR (or valve replacement)</td>
<td>3.064</td>
<td>.305</td>
<td>0.360-26.072</td>
</tr>
<tr>
<td>Severe SRV impairment</td>
<td>12.092</td>
<td>.002</td>
<td>2.424-60.309</td>
</tr>
<tr>
<td>PA band</td>
<td>1.606</td>
<td>.662</td>
<td>0.191-13.484</td>
</tr>
</tbody>
</table>

**HR**, Hazard ratio; **CI**, confidence interval; **VSD**, ventricular septal defect; **BT**, Blalock–Taussig; **TR**, tricuspid regurgitation; **SRV**, systemic right ventricle.

### TABLE 3. Reported long-term survival for 1282 patients after atrial switch surgery and 488 patients with congenitally corrected transposition of the great arteries

<table>
<thead>
<tr>
<th>Author</th>
<th>Population</th>
<th>Year</th>
<th>N</th>
<th>Cumulative probability of death</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>10 y</td>
</tr>
<tr>
<td>TGA-atrial switch</td>
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</tr>
<tr>
<td>Dobson et al</td>
<td>Mustard and Senning (pediatric data)</td>
<td>2012</td>
<td>133</td>
<td>0.96</td>
</tr>
<tr>
<td>Lange et al3</td>
<td>Mustard only</td>
<td>2006</td>
<td>395</td>
<td></td>
</tr>
<tr>
<td>Moons et al2</td>
<td>Mustard and Senning</td>
<td>2004</td>
<td>283</td>
<td>0.92</td>
</tr>
<tr>
<td>Sarkar et al30</td>
<td>Mustard and Senning</td>
<td>1999</td>
<td>358</td>
<td>0.82*</td>
</tr>
<tr>
<td>Wilson et al31</td>
<td>Mustard only</td>
<td>1998</td>
<td>113</td>
<td>0.90</td>
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<tr>
<td>ccTGA</td>
<td></td>
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</tr>
<tr>
<td>Dobson et al</td>
<td>ccTGA (operated and unoperated)</td>
<td>2012</td>
<td>71</td>
<td>0.87</td>
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<td>Lim et al2</td>
<td>ccTGA (123 physiologic and 44 anatomic repair)</td>
<td>2010</td>
<td>167</td>
<td>0.8</td>
</tr>
<tr>
<td>Hraska et al32</td>
<td>ccTGA (operated and unoperated)</td>
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<td>123</td>
<td>0.72</td>
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<tr>
<td>Yeh et al33</td>
<td>ccTGA (physiologic and 9 anatomic repair)</td>
<td>1999</td>
<td>127</td>
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TGA, Transposition of the great arteries; ccTGA, congenitally corrected transposition of the great arteries. *Mustard. | Senning.
improvement in the degree of TR when pre-PA band data existed. This is confirmed by 2 relatively large studies of 39 and 20 patients, respectively, although some older, smaller trials suggest a modest beneficial effect on TR. The biggest problem with PA banding seems to be the development of dysfunction of the subpulmonary LV.

There is only modest evidence for the efficacy of cardiac resynchronization therapy, and again the main problem seems to be the development of LV dysfunction. Cardiac transplantation is effective; however, there is the problem of limited donor organ availability in many parts of the world and the additional risk conferred by the complexity of abnormal anatomy and previous surgical intervention.

Functional Data

In keeping with the other cohort studies described previously, patients with atrial switch and patients with ccTGA continue to enjoy good functional status. Approximately 98% of patients with atrial switch for whom data were available (85/87) and 93% of patients with ccTGA (26/28) were in New York Heart Association class I or II. Median follow-up was to 29 and 33 years of age, respectively.

Formal cardiopulmonary exercise testing demonstrated reduced maximum oxygen uptake compared with peak predicted, with a mean peak oxygen uptake of 25 mL/kg/min for both atrial switch and ccTGA cases. This is in keeping with previous studies assessing exercise performance in patients with an SRV. We also demonstrated evidence of an abnormally high heart rate reserve and increased minute ventilation to carbon dioxide output ratio, again in keeping with previous work.

Reintervention

There was relatively high attrition in both cohorts, with cumulative probability of freedom from reintervention of 57% in atrial switch cases and 53% in ccTGA cases at 30 years.

In the TGA-atrial switch cohort, the relatively high rate of baffle reintervention may be explained by the fact that most patients had undergone a Mustard repair, and this technique was associated with a higher incidence of baffle obstruction than the Senning repair in an earlier meta-analysis. There was only 1 cardiac transplant. There have been no systemic AV valve replacements. Although 1 arterial switch procedure was performed in a patient who had previously undergone atrial switch, this was performed at the age of 8 years, and therefore before at least 18 years of follow-up had elapsed.

In the ccTGA cohort, physiologic repair of coexistent cardiac lesions accounted for most intervention. Most surgery was performed during childhood, with intervention after 18 years far less common than intervention before 18 years. The only surgery performed after 18 years of age was 1 tricuspid valve replacement and 1 late physiologic repair involving VSD closure and relief of left ventricular outflow tract obstruction via pulmonary valvotomy.

Systemic AV (tricuspid) valve replacement in patients with an SRV is controversial. Some centers advocate early replacement of an incompetent systemic AV valve because of the relationship between SRV impairment and systemic atrioventricular valve regurgitation. A double switch strategy, with a subsequent arterial switch operation, also is controversial.

Pacemaker Therapy

There was no significant difference between the 2 cohorts for survival from pacemaker insertion, and this was a particularly common problem in the ccTGA cohort. Use of cardiac resynchronization therapy was low, presumably reflecting the lack of evidence to support its use. Of note, there were no automatic implantable cardioverter defibrillator insertions. Although there is a high risk of sudden cardiac death, there are no established guidelines for formal risk assessment, and automatic implantable cardioverter defibrillator insertion in patients with an SRV is based on a “common sense” approach extrapolated from studies in other populations.

Study Limitations

As with all retrospective studies, our study is limited by those patients lost to follow-up both before and after the age of 18 years, although these numbers were relatively small. Another group of patients missing from the analysis are those with asymptomatic uncomplicated ccTGA, who have yet to present to medical services or are not recognized as having cardiac anatomy consistent with ccTGA. However, on the basis of our own experience, few patients with ccTGA present after the age of 18 years. Furthermore, all longitudinal cohort studies of ccTGA will have the same problem.

Functional data regarding ventricular function as assessed by cardiac MRI are lacking for a relatively large number of patients, and in many cases this is unavoidable because of the presence of a cardiac rhythm device. Furthermore, our service has yet to fully assess the cohort of patients with SRV by means of a cardiopulmonary exercise test, and the sickest patients are typically too unwell to allow determination of peak exercise capacity by this method.

CONCLUSIONS

Overall mortality is low at up to 40 years of follow-up in adult survivors of atrial switch surgery for TGA or ccTGA. Furthermore, mortality is remarkably similar between the 2 cohorts. This suggests that for the majority of patients, it is actually the SRV itself, rather than the nature of any prior surgery, that determines true long-term prognosis. SRV
dysfunction increases the overall hazard for death or transplant, and most deaths are sudden or due to heart failure. Morbidity is substantial, with a high incidence of tachyarrhythmia, surgical or catheter intervention, and pacemaker insertion. Tachyarrhythmia is a particular problem for TGA-atrial switch cases because it occurs significantly earlier than in ccTGA cases. However, moderate to severe SRV dysfunction, TR, and functional impairment are relatively rare up to 40 years. Risk stratification for sudden cardiac death and the identification of suitable strategies to manage SRV impairment will be key to improving survival in the future. In view of the high burden of SRV impairment, an early aggressive strategy aimed at anatomic repair may be warranted in the patients with ccTGA who require corrective surgery for coexistent lesions. For the asymptomatic patient with uncomplicated ccTGA, there is insufficient adult follow-up data to comment on whether an anatomic repair strategy is better than conservative management.

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References

Discussion
Dr Shunji Sano (Okayama, Japan). There is always debate on whether the morphologic RV supports the systemic circulation for the patient’s entire life or not. The American College of
Cardiology/American Heart Association guidelines show that ventricu-
lar dysfunction seems to be related to systemic AV valve re-
gurgitation. In the absence of associated congenital anomalies, 
primary RV failure is uncommon. Abnormality of the systemic 
AV valve is Ebstein-like malformation in 90% of the patients. In 
your series in the adult group, only 4 patients had Ebstein’s. 
From your data, mortality is low with survival of 90% for 
TGA-atrial switch and 84% for ccTGA. However, cumulative 
mortality at 40 years from birth is 68% for TGA-atrial switch 
and 67% for ccTGA. This means most of the deaths in these pa-
tients occur in the pediatric age, and in the patients who survived 
to adult age, mortality is low with good RV function.

Do you have any data of TR and RV impairment of ccTGA in 
the pediatric age population, and what was the reason of death 
of these patients after TGA-atrial switch and ccTGA in the pediat-
ric age population?

Dr Dobson. The main aim of this study was to look at outcomes 
in the adult patients who were cared for by the adult congenital 
heart disease service. In terms of causes of death in the pediatric 
population, we didn’t assess this in detail. Furthermore, I have lim-
ited data regarding the severity of TR and systemic RV impairment 
as estimated from echocardiography or MRI in patients before the 
age of 18 years.

However, one can make some deductions. A significant number 
of the pediatric deaths occurred in association with the initial re-
pair and early postoperative course, and most of these were in 
the late 1970s or early 1980s. SRV failure also was a significant 
problem: Five patients in the pediatric population subsequently 
went on to have a successful double switch procedure, and 1 pa-
tient required cardiac transplantation after unsuccessful PA band-
ing. The other cause of mortality in the pediatric patients was 
sudden cardiac death. This raises the question of how to prevent 
reports due to SRV failure.

We know that as patients become older it is difficult to train the 
morphologic LV in preparation for an anatomic repair. Therefore, 
patients should be prepared for this at a young age, really as soon 
as it has become evident that they have significant TR or signifi-
cant SRV failure also was a significant problem: Five patients in the pediatric population subsequently 
went on to have a successful double switch procedure, and 1 pa-
tient required cardiac transplantation after unsuccessful PA band-
ing. The other cause of mortality in the pediatric patients was 
sudden cardiac death. This raises the question of how to prevent 
reports due to SRV failure.

The key role of any research in this group is to try and identify 
a strategy that promptly identifies and intervenes in those patients 
at high risk of SRV failure, while at the same time recognizing that 
there are other patients who will do well with a more conservative 
approach.

Dr Sano. Your freedom from arrhythmia at 40 years was 0.51 
for TGA-atrial switch and 0.93 for ccTGA. There is a discrepancy 
between TGA-atrial switch and ccTGA. Most of the arrhythmia 
in those with TGA-atrial switch was atrial tachyarrhythmia, and most 
of your atrial switch procedures were Mustard. Therefore, the rea-
son of the arrhythmia is due to late complication of Mustard rather 
than RV dysfunction. Is the rate of arrhythmia the same in the Sen-
ning and Mustard groups?

Dr Dobson. The key thing is that the Senning group was 
smaller. I don’t have the exact figures at hand, but I will find it 
for you later. If one looks at the absolute values, certainly Mustard 
cases accounted for most of the burden of tachyarrhythmia in the 
TGA-atrial switch cohort. Of the 21 patients in the atrial switch co-
hort with documented tachyarrhythmia, only 2 underwent Senning 
repair; the rest underwent Mustard. However, the smaller number 
of patients underwent Senning, accounting for only one fifth of the 
atrial switch cases, and subsequent analysis demonstrated that 
there was no statistically significant difference in freedom from 
tachyarrhythmia between the 2 groups, with a P value of .892 as 
estimated by the log-rank test: this data set is therefore underpow-
ered in this respect.

However, when one compares freedom from tachyarrhythmia 
between the TGA-atrial switch cohort and the ccTGA cohort 
you are absolutely correct, there were higher rates of arrhythmia 
in TGA-atrial switch cases compared with ccTGA cases. My own 
feeling is that this reflects the extensive atrial scar tissue 
that has occurred as a result of the initial surgery, leading to a pro-
Brim for atrial macro-reentrant tachycardia.

Dr Sano. So that means if you perform a Senning operation, 
most of these arrhythmias will be avoided. Recent reports from 
many centers show a 20-year survival of more than 70% to 80% 
after the double switch, and the Fontan operation is now adopted 
in many cases rather than complex intracardiac repair, with a 10-
to 20-year survival of 70% to 90%, and the reintervention after 
Fontan is low.

We recently reported our results of ccTGA. Although the mean 
follow-up is only 10 years, in a double switch with the Senning ar-
terial switch group, there was no mortality. In the Senning–Rastelli 
group, we measured the length between interventricular septum and 
aorta, and all early and late mortality came from the group with 
a long distance between the interventricular septum and aorta. There 
was no mortality in the patients with the short distance. Therefore, 
we do a Fontan operation in the long distance group and a double 
switch in the short distance group. I hope the long-term survival 
will improve using different techniques in individual patients.

Dr John Foker (Minneapolis, Minn). This is a valuable study, 
because there are a number of these patients. I did 50 atrial baffle 
procedures (modified Shumacker) for simple transposition. They 
all left the hospital, 2 or 3 died later suddenly, but many of them 
are doing well, including 1 woman who recently gave birth to 
a child with TGA.

My question concerns the RV dysfunction. These patients 
underwent operation at a time when myocardial protection was rudi-
mentary at best and was, in essence, clamp and go, with the RV out 
under the hot lights. Do you think the RV dysfunction, when you 
see it, could be due mostly to this poor myocardial protection 
rather than any intrinsic inability of the RV to serve as the systemic 
ventricle? If so, when you looked at these hearts, would you see 
subendocardial fibrosis in those that were failing?

Dr Dobson. One would expect that more primitive myocardial 
protection techniques would result in greater myocyte necrosis in 
the SRV and consequent impairment. However, it is uncertain 
whether this is permanent or there is potential for recovery. There 
are clearly many other factors that could contribute to the develop-
ment of SRV failure, not least because we know that some patients 
with isolated ccTGA who have never proceeded to surgery some-
times develop significant cardiac failure. To answer your question, 
one would need a larger study population. One could then compare 
outcomes between patients who had been repaired in the current
era with all the advantages of modern techniques of myocardial protection and patients who had been repaired 20 or 30 years ago when such technology was less advanced. Multivariate analysis would then allow adjustment for confounding variables and provide insight into the effect of cardiopulmonary bypass in predicting those who will go on to develop late systemic RV impairment.

In answer to the second part of your question, subendocardial fibrosis is something that is currently difficult to assess by MRI. Even with late gadolinium enhancement, you are unlikely to be able to distinguish whether someone definitely has fibrosis. There is some interesting work looking at the use of MRI edema sequences for this purpose. It would be a fascinating study to try and objectively assess the amounts of fibrosis in SRVs and then compare these with the era of surgical repair. But I don’t have the data for that in this study. The other problem is that many of these patients require permanent pacing, which precludes assessment by MRI at least in the current era.

Dr Gerhard Ziemer (Chicago, Ill). I understand that you have to simplify your conclusions in a talk like this, but I have to make 2 remarks.

For atrial repair of TGA, you cannot really put patients with and without VSD into 1 group and give them a single incidence for arrhythmias. The same is true for RV function, because you may have a different cause of RV function impairment, either secondary to TR or preceding it. This relates to VSD closure or no VSD closure. So at least in your article you should discuss this.

My second point relates to you just comparing arrhythmia incidence for ccTGA with arrhythmias in normal transposition. But you only can compare them for their types of arrhythmias: Patients post-atrial switch for TGA mainly have atrial arrhythmias and patients post-ccTGA have sudden AV block with and without surgery. So there is really a different mechanism of late failure, but I understand the simplification in your presentation.

Dr Dobson. Thank you.

Dr Craig Smith (New York, NY). This study is a sobering reminder that when we are dealing with a population with such a long life expectancy, the duration of follow-up required to answer any of these questions is long, and I look forward to the 40-year follow-up.