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Managing adult Fontan patients: where do we stand?

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ABSTRACT The Fontan operation is performed as a palliative procedure to improve survival in infants born with a functionally univentricular circulation. The success of the operation is demonstrated by a growing adult Fontan population that exists with this unique physiology. Late follow-up has demonstrated expected and unexpected sequelae, and has shown multisystem effects of this circulation. This review discusses the challenges of managing the late complications in terms of understanding this unique physiology and the innovative therapeutic interventions that are being investigated. The challenge remains to maintain quality of life for adult survivors, as well as extending life expectancy.

Introduction

A univentricular circulation describes a congenital heart defect where a biventricular repair is not possible, typically associated with hypoplasia of one of the ventricles and/or atrioventricular valves, and often with other associated features (figure 1). It is rare, with a prevalence of ∼1:10 000 live births [1]. Without intervention in early life, survival to adult life is extremely rare and death in childhood is typical [2, 3]. In 1971, Francis Fontan and his colleagues described a palliative procedure that now bears his name [4], the principles of which have remained unchanged, although the procedure has evolved and is now performed as a standardised surgical or hybrid surgical procedure in developed healthcare economies across the world. Survival to adult life is now expected, but comes with many challenges for the patient, their family and also for their clinicians [5]. In this paper we describe the unique Fontan physiology and the late effects and management strategies employed to maintain health in this complex condition.

The univentricular heart

Figure 1 shows three examples of univentricular hearts, in which there is dominance of a ventricle and associated abnormalities. In each case a biventricular circulation is not achievable, and without intervention chronic cyanosis, effort intolerance and volume loading of the circulation ensues, leading to significant early morbidity and mortality. Fontan described the concept of bypassing the subpulmonary ventricle by routing the systemic venous return directly into the pulmonary circulation, thus committing the single ventricle to the systemic circulation, abolishing cyanosis, reducing the volume load on the ventricle and improving effort tolerance; the original “atriopulmonary connection” (APC) operation (figure 2a). The disadvantage of this arrangement was that the atrial mass became a sump, leading to
energy losses in the circulation and increasing the work of the ventricle, and atrial dilatation predisposed the patient to complex atrial arrhythmias associated with significant morbidity and mortality. Modifications were thus made to optimise the flow characteristics of the Fontan circuit by means of an intra-atrial wall, creating a channel within the atrial mass by which the inferior vena cava flow could be streamed directly to the pulmonary arteries; this is called a lateral tunnel Fontan (figure 2b) [6]. Further modifications were made to remove all atrial tissue from the circulation, believing that this would further improve the flow dynamics and reduce the risk of atrial arrhythmias. This was achieved by means of an extracardiac conduit connecting the inferior vena cava to the pulmonary arteries (figure 2c) [7]. These latter two operations are total cavopulmonary connection (TCPC) forms of Fontan and are the methods employed currently in surgical centres. Initially, adult survivors were mainly APC Fontan patients, but increasing numbers of both forms of TCPC Fontan patients now survive to adult life.

FIGURE 1 Anatomical variations on a univentricular circulation. a) Double inlet left ventricle (LV) with ventriculoarterial discordance and no pulmonary stenosis; b) tricuspid atresia with hypoplastic right ventricle (RV), ventriculoarterial concordance and hypoplastic pulmonary artery (PA); and c) hypoplastic left heart syndrome with atroventricular and ventriculoarterial concordance, hypoplastic left ventricle, aorta and aortic arch. Many classification systems have been proposed for univentricular hearts. The physician treating adult congenital heart disease needs an understanding of atrial arrangement, atrioventricular valve morphology and the morphology of the dominant ventricle, in addition to the surgical procedures that the patient has undergone, as all of these features will contribute to problems that may be faced in adult life. RPA: right pulmonary artery; LPA: left pulmonary artery; SVC: superior vena cava; RA: right atrium; LA: left atrium; PV: pulmonary vein; IVC: inferior vena cava; TV: tricuspid valve; MV: mitral valve.

FIGURE 2 Fontan circulation. a) Modified atrioventricular connection Fontan in which the systemic venous atrium is directly connected to the pulmonary arteries; b) inferior vena cava flow streamed directly to the pulmonary arteries in a right atrium lateral tunnel Fontan; and c) extracardiac conduit connecting the inferior vena cava to the pulmonary arteries. The latter two operations are forms of total cavopulmonary connection (TCPC) Fontan, and show a fenestration between the total cavopulmonary connection pathway and the right atrium, a common surgical modification performed to relieve pressure in the pathway. RPA: right pulmonary artery; LPA: left pulmonary artery; SVC: superior vena cava; RA: right atrium; LA: left atrium; IVC: inferior vena cava; RV: right ventricle; LV: left ventricle.
The physiology of the Fontan circulation
Fundamentally, the Fontan circulation differs from the normal parallel pulmonary and systemic circulations by existing as a single circulation in which the pulmonary and systemic circulations run in series. This is powered by the single ventricle and is dependent on a low pulmonary vascular resistance (PVR) to maintain ventricular preload and systemic venous hypertension, driving the transpulmonary blood flow. Figure 3 represents the particular challenges faced at each stage of the Fontan circulation and the current strategies employed to deal with them.

Effects of systemic venous hypertension
Systemic venous hypertension is necessary to drive blood flow through the pulmonary vasculature, with systemic venous pressure typically ~5 mmHg higher than the pulmonary venous atrial pressure. Pressures in the hepatic veins are higher than normal, with abnormal liver haemodynamics predisposing to late liver complications [8]. Elevated thoracic duct pressures are thought to contribute to prolonged chylous effusions in the early post-Fontan period [9], and later to the development of plastic bronchitis [10, 11] and protein-losing enteropathy (PLE) [12, 13].

Hepatic effects
There is growing recognition of the deleterious effects of systemic venous hypertension on hepatic function and the development of fibrosis, cirrhosis and, worryingly, hepatocellular carcinoma [14–16]. Patients under follow-up routinely undergo liver surveillance including ultrasonographic assessment of fibrosis in order to identify early hepatocellular carcinoma amenable to resection [17], although the optimal type and timing of screening for liver complications remains uncertain. Subclinical hepatic disease can be challenging for patients undergoing cardiac transplantation for Fontan failure and careful multidisciplinary assessment is required [18].

Protein-losing enteropathy
This complication of the Fontan circulation has challenged physicians for many years [12, 19], diagnosed by a low serum albumin level and the presence of high faecal α1-antitrypsin levels [20]. It is a characteristic feature of Fontan failure, leading to peripheral oedema, ascites, pleural effusions, malabsorption and loss of immunoglobulins. There is no reliable therapy, and treatment strategies involve ensuring that the Fontan circuit is free of narrowing, and intervening if necessary. Fontan fenestration has been performed with the acceptance that cyanosis will follow the procedure [21]; latterly, manipulation of the thoracic duct has been proposed [22]. PLE may successfully resolve following cardiac transplantation [23]. Medical therapies include oral budesonide [24, 25], unfractionated heparin [26], spironolactone [27, 28], loperamide [29] and targeted pulmonary vasodilators [30], but none are reliably effective.

FIGURE 3 Schematic of challenges and management strategies in the Fontan circulation. LPA: left pulmonary artery; PA: pulmonary artery; AVV: atrioventricular valve; ACE: angiotensin-converting enzyme; TCPC: total cavopulmonary connection.
Atrial stretch and arrhythmias

Atrial dilatation is commonly seen in the APC Fontan patient. Dilatation and the presence of surgical scars within the atrial mass predispose these patients to atrial arrhythmias, predominantly of the atypical atrial flutter type [31–33]. Scarring within the atrial mass provides the substrate for macro re-entrant tachycardia circuits to form, and the occurrence of this intra-atrial re-entrant tachycardia is a medical emergency in the Fontan patient, in whom it is poorly tolerated. A possible explanation for this is that the atrial arrhythmia increases the pressure in the pulmonary venous atrium leading to a reduction in the pressure gradient down which the pulmonary blood flows. Reduction in pulmonary blood flow further depletes the preload in the single ventricle and leads to a fall in cardiac output. It is a frequent cause of emergency admission in the adult congenital heart disease (ACHD) population, despite Fontan patients being only a small portion of this population. Delay in cardioversion in susceptible patients can lead to death due to circulatory failure.

Many strategies have been employed to reduce the arrhythmia burden, including oral antiarrhythmics such as sotalol and amiodarone. However, the side-effect profile of amiodarone in the young adult makes this less than ideal [34]. A catheter-based ablation strategy has been employed with success in many cases [35–38], but the recurrence rate remains high [38], in part due to the multiple potential substrates within the heart. It is likely that atrial arrhythmias are inevitable in the APC Fontan population [39]. Much has been published about the favourable outcomes in TCPC Fontan, owing to less atrial stretch [40, 41], and with this in mind many centres have performed conversion operations creating a TCPC Fontan and performing cryoablation surgery to remove the substrate for intra-atrial re-entrant tachycardia and improve Fontan flow characteristics [42–46]. However, the conversion operation comes with a significant morbidity and mortality risk [47], and should be considered only in selected cases, having been assessed by an experienced team [48]. Furthermore, TCPC Fontan operations have not completely eliminated the problem of atrial arrhythmia, and pose particular challenges for the interventional electrophysiologist. The use of oral anticoagulants is recommended in all patients who have had an arrhythmia.

Cyanosis and collateral formation

The maintenance of adequate preload for the single ventricle is a challenge, and one of a number of “autoregulatory” phenomena seen in the Fontan population is the development of veno-venous collateral vessels that pass from the systemic veins to the pulmonary venous circulation [49]. The benefit is to maintain preload and cardiac output, but this comes at a cost of often profound cyanosis that worsens on exertion. Many centres have employed a catheter-based strategy of embolisation of these abnormal vessels [50, 51]. Whether this impacts on symptoms or survival has not been determined [52], and a further reduction in saturations post-procedure suggests that recurrence rates may be high [51].

Effects of the lack of a subpulmonary ventricle

The fundamental difference between the Fontan and a normal biventricular circulation is the lack of a subpulmonary ventricle. In all its iterations, the Fontan circulation lacks a synchronised ejection from a subpulmonary ventricle. In all its iterations, the Fontan circulation lacks a synchronised ejection from a subpulmonary ventricle, relying upon changes in intrathoracic pressure during breathing to draw blood through the lungs, and systemic venous hypertension to provide the “header” pressure in the circulation. We have already discussed the vulnerability of this circulation to the effects of atrial arrhythmia, but let us consider its impact on cardiac output and the ability to exercise.

Gewillig et al. [53] addressed this elegantly in a discussion paper on what controls the cardiac output in the Fontan circulation. They explored the areas of contractility, heart rate, afterload, preload and transpulmonary blood flow, before developing a conceptual model of the ageing Fontan circulation (figure 4a–d). Key to the maintenance of cardiac output is the preservation of adequate preload, which depends on a low PVR, with even modest changes in PVR leading to a significant fall in cardiac output and exercise capacity. This hypothesis is difficult to test in routine practice, as standard measurements of PVR by cardiac catheter are technically challenging in the Fontan circulation, but there is evidence of the importance of PVR in the failing Fontan.

Diastolic dysfunction is often quoted as a late effect of the Fontan circulation [54–61], but it remains poorly understood and is likely to be multifactorial. Moreover, measurement of diastolic function using echocardiography may well be inaccurate in a circulation that is, by its nature, preload-depleted. Similarly, chronotropic incompetence may occur due to scar-related sinus node dysfunction [62]. However, atrial pacing does not necessarily augment cardiac output [63], which suggests that the Fontan circulation functions optimally in a narrow range of heart rates. Whether diastolic dysfunction and chronotropic incompetence are likely long-term outcomes of the Fontan circulation or merely a reflection of the physiology remains to be determined.
Other organ system involvement
With increasing length of follow-up, other organ systems appear to be vulnerable to abnormalities. Neurocognitive and developmental issues are of concern [64], and there appears to be an abnormally high incidence of some psychiatric issues, such as anxiety [65]. Bones [66], endocrine systems [67, 68] and lymphatics [69] are also known to be affected in a substantial number of patients with Fontan physiology; the pathogenesis, diagnosis and optimal treatments for these issues are areas of current research activity. Overt renal failure is not routine post-Fontan, but lower-than-normal glomerular filtration rates are found in ∼10% of Fontan patients during long-term follow-up [70].

Pulmonary vascular remodelling
There is morphological and haemodynamic evidence of pulmonary vascular remodelling and increased PVR. In lung biopsies taken at the time of bidirectional Glenn procedure, patients who subsequently struggled following Fontan completion were demonstrated to have muscularised distal pulmonary arteries; this was a common finding in patients undergoing Fontan completion, despite normal pre-procedural pulmonary haemodynamics [71, 72]. Furthermore, elevated PVR has been demonstrated post-transplantation in patients who have undergone cardiac transplantation for a failing Fontan circulation [73]. Modest elevations in indexed PVR have been found in healthy Fontan patients when studied in detail, with acute lowering of the indexed PVR with inhaled nitric oxide [74]. Therefore, the evidence that pulmonary vascular remodelling exists in these patients is strong, as is evidence that it impacts on the health of these patients. That it can be reduced acutely under experimental conditions is promising, but there is a lack of evidence of a persisting effect beyond the acute reduction, and of an effect that can be achieved with oral pulmonary vasodilators.

Pulmonary vasodilator therapy in the Fontan circulation
Phosphodiesterase type-5 inhibitors
Phosphodiesterase type-5 inhibitors have been studied both acutely and in case series, with promising preliminary results [75]. In the acute setting, sildenafil has been shown to improve the peak oxygen uptake (V'O₂) in adult Fontan patients [76]. In unselected Fontan populations, sildenafil improves some exercise parameters [75, 77], but in selected failing Fontan patients, sildenafil has been reported to improve symptoms and features of Fontan failure [30, 78–82]. This effect appears dependent on reduction of PVR [83] rather than direct effects on ventricular function [84]. It would appear that “targeted” pulmonary phosphodiesterase type-5 inhibition may be of benefit to the Fontan patient, although adequately powered clinical trials are still lacking in this regard.
**Endothelin receptor antagonists**

The finding of early pulmonary vascular remodelling in the failing Fontan [71] makes the use of endothelin receptor antagonists (ERAs) an attractive option. In early animal studies, ERAs have been demonstrated to reverse established pulmonary vascular remodelling, in addition to pulmonary vasodilation [85, 86]. Bosentan therapy has been used successfully in many forms of pulmonary arterial hypertension [87]; ambrisentan [88] and macitentan [89] have also shown promise. Increasingly, the focus has been on the combination of therapies targeting the endothelin, cyclic GMP and cyclic AMP pathways [90]. The use of ERAs in the Fontan circulation appears to have an acceptable safety profile [91] and may be effective [92, 93]. The TEMPO study [92] evaluated medium-term bosentan therapy (14 weeks) in a large cohort of adolescent and adult Fontan patients in Scandinavia. Patients were not selected on functional status, and were randomised 1:1 to placebo or bosentan. There was a modest improvement in peak VO₂ and an increase in exercise duration with bosentan therapy, and ~25% of the treatment arm improved functional class. The majority of the study population were in functional class I with no perceived limitations. Whether targeting high-risk failing Fontan patients with ERAs, or even combination therapy, is an effective strategy remains to be established and requires larger collaborative randomised clinical trials.

**Prostanoids**

Two studies have evaluated the use of inhaled iloprost in the management of Fontan patients. Inhaled iloprost was well tolerated and led to symptomatic improvement and improvement of peak VO₂ in limited patients [94, 95]. The use of prostanoids is not widespread and requires further clinical trial data to recommend its routine use.

There is an emerging evidence base for the use of targeted pulmonary vasodilator therapy in the failing Fontan patient. What remains to be determined is which patients will benefit most, which pulmonary vasodilator is best suited for use in the Fontan circulation and whether combination therapy using currently available targeted pulmonary vasodilators has greater efficacy in Fontan patients than monotherapy. Challenges remain regarding how best to establish a treatment effect and how best to identify patients who may benefit from this therapeutic strategy. While it can be difficult to demonstrate an increase in maximal performance, there may be merit in developing end-points of submaximal exercise performance, such as endurance time [96] or oxygen uptake efficiency slope [97, 98]. How to categorise functional class in patients who do not perceive themselves to be limited is a complex issue. Adaptations to lifestyle in early life is common in the congenital heart disease population and quality of life from a personal perspective is often good [99]. The development of a congenital heart disease specific quality of life score [100] allows investigators to differentiate between physical and psychological aspects of quality of life, and this yields interesting results. Whilst Fontan patients and families report a good quality of life in general [99], when this is examined further, there is a substantial reduction in the physical quality of life while psychosocial quality of life remains good [101, 102]. It is therefore important to assess functional class objectively by means of a structured survey, in order to measure any treatment effect on physical symptoms.

**Follow-up of adult Fontan patients**

The process of transition to adult care is by its nature flexible, reflecting the individual circumstances of the patient, their family and their clinical status. Ideally, transition should start no later than 14 years of age and should start in the paediatric setting, building up to transfer to adult care between 16 and 18 years.

Empowerment of the Fontan patient is aided by a detailed discussion of their diagnosis and how it may impact on various aspects of their adult life. This should include a discussion regarding pregnancy and contraception for all female patients, and all should be counselled against the use of recreational drugs and cigarette smoking. Although alcohol is permitted, patients should not binge drink, as this can precipitate arrhythmias. Whether alcohol consumption impacts on the late hepatic function is not established. Body piercing and tattoos are discussed, as is the need for regular dental follow-up. Advice should be given regarding career options, for example, currently the UK armed forces do not accept recruits with any history of congenital heart disease. Discussions also include, where appropriate, sporting and other leisure activities, including whether patients can scuba dive.

The initial adult assessment should include routine cardiovascular examination and the following routine examinations: 12-lead ECG, transthoracic echocardiogram, 24-h ECG monitoring, cardiopulmonary exercise and cardiac magnetic resonance (CMR) imaging in those in whom it is possible. This provides a baseline haemodynamic assessment with which to compare future results.

It is good practice to provide patients with a copy of their 12-lead ECG for them to carry in case they are ever reviewed in an emergency department with palpitations; it is good practice for patients to have copies of all of their healthcare records. Patients should have contact details for the unit, the hospital and clinical nurse specialists. Follow-up is every 12 months, or at short notice if a situation arises.
Standard medical management
The nonpulsatile and often turbulent blood flow through the Fontan predisposes the patient to thromboembolism [103]. Subclinical microthrombi in the Fontan pathway could lead to an increase in the PVR, with large occlusive thrombus being a known cause of death [104, 105]. Persistent right-to-left shunts are frequent due to Fontan fenestrations and veno-venous collaterals, with an associated risk of paradoxical embolus. For these reasons many centres routinely anticoagulate their Fontan population, but this remains a subject of debate. There is no evidence in clinical studies of superiority of vitamin K antagonists over antiplatelet strategies [106–108]; however, patients prescribed neither of these do encounter frequent thrombotic complications [109].

Many Fontan patients are treated with angiotensin-converting enzyme inhibitors, despite there being minimal evidence for their use [110, 111], and their use remains controversial [112]. β-adrenergic blockers and amiodarone are the main therapeutic strategies for secondary prevention of arrhythmias.

Quantitative assessments of performance demonstrate severe limitation in all but the best Fontan patients (rare asymptomatic patients with normal exercise test parameters), but many patients would regard themselves as free of significant effort intolerance. Many have suggested that exercise as a therapeutic strategy could benefit the Fontan patient, due to improved oxygen extraction at the periphery and the additional benefit of the venous muscle pump during exercise. Exercise has been shown to be effective in increasing the peak oxygen consumption of Fontan patients [113–115].

Imaging modalities used in adult Fontan patients
Transthoracic echocardiography (TTE) remains the most widely available imaging modality in adult cardiology. However, the interpretation of TTE images of the univentricular circulation requires experience and training. Additional training in cardiac morphology of operated and unoperated hearts is of great use.

TTE gives immediate assessment of ventricular function, outflow tract obstruction and atrioventricular valve structure and function. It rarely gives clear images of the Fontan pathway, and is not sufficient to exclude obstruction in the pathway or pulmonary venous pathway obstruction. CMR imaging is now routinely used in the assessment of the Fontan pathway flow, to exclude obstruction and to assess pulmonary venous return [116–119]. In addition, CMR imaging gives very accurate assessment of ventricular function and assessment of branch pulmonary arteries, and, importantly, can exclude re-coarctation or narrowing in the Damus pathway. Transoesophageal echocardiography, in experienced hands, is a useful adjunct to TTE. The pulmonary veins can be readily identified and thrombus within the Fontan circuit can be seen clearly, in addition to obtaining quantitative assessment of the atrioventricular valves.

Cardiac catheterisation can provide both functional and pressure data and indirect estimates of cardiac output by measurement of the mixed venous saturation; however, this is used less in the adult population, where routine assessment largely relies on CMR imaging. Catheter-based interventions are not common in the adult population, but may be required to alleviate obstruction in the Fontan pathway [120], for managing profound cyanosis [50, 121] or for stenting of branch pulmonary artery stenosis or venous pathway obstruction [120]. Emergency management of haemoptysis is usually performed in the catheter lab, where identification and embolisation of aorto-bronchial fistulae may be necessary [122].

Assessment of exercise performance
The assessment of exercise performance in ACHD patients is best achieved using cardiopulmonary exercise testing, which gives accurate, reproducible and quantifiable data on cardiac and respiratory performance. This allows for tracking of performance and assessment of unexpected deterioration. Most Fontan patients have undergone two or three open procedures and may have undergone a thoracotomy. As with other complex lesions, this leads to a restrictive lung defect in many Fontan patients. This finding, in addition to the restrictions in augmentation of the cardiac output on exercise, contribute to a reduction in the measured peak V′O2, with average Fontan patients achieving a peak V′O2~60–70% that of age-/sex-/size-matched controls [123].

Peak V′O2 has been used as an end-point in a number of clinical studies of the Fontan population [92, 93, 124–126]. This has merit in that results are easily quantifiable and reproducible even in the Fontan population [126]. Additional prognostically relevant parameters, such as anaerobic threshold, minute ventilation/carbon dioxide (CO2) slope and changes in peak end tidal CO2 are easily quantifiable and could be used to identify patients who may benefit from certain treatment strategies. The haemodynamic response to exercise in Fontan patients is illustrated in figure 5 [127].

Pregnancy and contraception
A number of centres have recently reported the outcomes of pregnancies in women with a Fontan circulation [128–132], and all of these support earlier published series [133–135]. Both fetal and maternal
outcomes are generally (but not uniformly) good. Although many pregnancies are carried to term, prematurity is not uncommon and the average gestational age ranges between 31 and 33 weeks [128, 130]. There remains a high risk of fetal loss in the first two trimesters, with miscarriage rates as high as 50% being typical. Careful pre-pregnancy assessment and counselling is warranted, delivered through dedicated pre-conception counselling. Such clinics inform of the risks of pregnancy and prepare women for the necessary detailed antenatal care that will follow conception. Following counselling, many women choose to defer pregnancy [136], and despite this very high level of specialised care, viable pregnancies occur in the minority of conceptions [136]. Specific measures during antenatal care include management of anticoagulation, with many centres converting warfarin to low molecular weight heparin for the duration of the pregnancy, with the need for weekly anti-Xa monitoring. Angiotensin-converting enzyme inhibition is omitted for the duration of the pregnancy. Delivery planning involves a full multidisciplinary team including an obstetrician, anaesthetist and cardiologist, with a preference for a normal vaginal delivery at term with a short second stage.

The choice of contraception for Fontan patients is relatively straightforward: contraception containing oestrogen is not recommended as this may increase the risk of venous thromboembolism. A number of methods of progestogen-only contraception exist in the form of oral contraception, depot and implantable subdermal contraception. The progestogen intrauterine coil is effective and safe for those who have undergone a pregnancy. Although barrier methods are cardiac-safe they are not as effective as other methods, and those more effective forms of contraception should be employed.

Late effects of the Fontan circulation
We have already discussed the deleterious effects of systemic venous hypertension. The development of arrhythmias could be thought to be an inevitable part of the Fontan patient’s progress [39]. Late mortality occurs, often related to the development of PLE, ventricular dysfunction, thrombotic complications and increasingly life-threatening complications of chronic liver disease [137, 138]. The goal of our management of the adult patient is to preserve symptom-free survival for as long as possible.

This requires a detailed and routine assessment, looking for evidence of deterioration in key areas. Regular review of imaging and exercise performance, monitoring for evidence of liver fibrosis and failure and development of hepatocellular carcinoma are now routine practice in large Fontan centres. This requires the integration of ACHD cardiologists, interventional congenital cardiologists, imaging cardiologists, hepatologists, respiratory physicians, obstetricians and fetal medicine teams, cardiac anaesthetists and intensivists, primary care physicians, clinical nurse specialists, congenital cardiac surgeons and the patients and their families into a multidisciplinary team, dedicated to the identification and management of complications of the Fontan circulation.

Fontan failure may be heralded by a number of symptoms and signs including effort intolerance, increasing arrhythmias, increasing cyanosis, ascites and peripheral oedema and thrombotic complications. The management strategy is one of identification of modifiable conditions and intervention where necessary. Full imaging of the Fontan pathway, ventricle, the outflow tract aorta and arch using TTE and CMR imaging is a first step. In the event of an obstruction being identified, catheter-based interventions may be possible, as previously discussed. Surgical intervention is effective in certain cases. Evaluation of atrioventricular valve regurgitation can be difficult and transoesophageal echocardiography is often
necessary to understand the mechanism. Although surgery for atrioventricular valve regurgitation can be undertaken, careful consideration of the cases in which this can be attempted needs to be made, as “redo” Fontan surgery is accompanied by high mortality and uncertain outcomes.

The lack of effective management strategies has led to increasing consideration of heart transplantation for the failing Fontan, which in itself comes with a number of challenges. The organ donor shortage in many developed healthcare economies has led to increasingly harsher listing criteria, such that only the most urgent cases are likely to be transplanted. In addition, having multiple previous operations increases the likelihood of the sensitisation of the recipient to the donor organ, further reducing the donor pool [139]. Prolonged donor ischaemic times are more likely, given the need for complex explantation of the recipient’s heart. The outcomes of Fontan transplants are less good than non-Fontan ACHD transplants, but despite higher early mortality, long-term survival is better in the ACHD population than the non-ACHD population.

Patients with preserved ventricular function and reduced albumin levels do less well than those whose indication for transplant is primarily pump failure [140]. The presence of liver disease presents an additional challenge and requires a collaborative approach with hepatologists in the pretransplant assessment and the early postoperative phase [18]. It remains the case that the majority of failing Fontan patients do not receive a heart transplant [141].

The reality of lack of organ availability and uncertainty over suitability for transplantation has led to the concept of mechanical support for the failing Fontan. While specific Fontan mechanical support strategies are being considered [142–148], centres have inserted ventricular assist devices in a number of failing paediatric Fontan patients, providing a successful bridge to heart transplantation [149–152]. Whether a total artificial heart strategy could be a long-term solution in this population remains to be established.

Conclusion

The Fontan circulation is in effect an alternative physiology that has challenged the assumptions of physicians. Terms such as chronotropic incompetence and diastolic dysfunction are in effect descriptions of the autoregulatory effects of the “normal” Fontan circulation; this challenges physicians to understand this novel physiology and to develop novel management strategies when they encounter what appears to be inevitable deterioration in adult life. Collaborative clinical trials are required in large populations of symptomatic Fontan patients to determine the best diagnostic and treatment strategies. Longitudinal data collection will define the natural history of the condition and will allow us to inform patients and families. Presently, our adult Fontan population are pioneers for their paediatric Fontan counterparts. We hope that by investigating our failing adult Fontan patients we can increase the life expectancy and quality of life of future generations of Fontan patients.

References


