Background: More than 90% of children with congenital heart defects now survive into adulthood; just a few decades ago, survival was rare, particularly among patients with complex defects. The new population of adults with congenital heart disease presents a special challenge to physicians from all of the involved specialties.

Method: Selective literature review.

Results and conclusion: A complete cure of the congenital heart defect in childhood is exceptional, and most adult patients continue to suffer from residual problems and sequelae. Further surgery or catheter interventions may be needed. Potential late complications include arrhythmias, heart failure, pulmonary hypertension, endocarditis, and thromboembolic events. The management of these patients during pregnancy or non-cardiac surgery remains a challenge. If this evolving patient population is to receive the best possible care, the adequate provision of specialized medical services is a necessary, but not sufficient, condition: patients and their referring physicians will also need to be aware that these services are available, and then actually make use of them. Moreover, optimal communication among all of the involved physicians is essential.

Cite this as:

Thanks to advances in pediatric cardiology and pediatric heart surgery in the past couple of decades, a new group of patients has emerged—adults with congenital heart defects. This continually growing population presents a particular challenge for cardiologists (e1). The estimated total number of adults with congenital heart defects in Germany is in excess of 250 000 (20% to 50% of these patients are estimated to have complex anomalies) (1, e2). Recommendations for the organization of healthcare services (2) and the qualification of doctors whose remit it is to care for adults with congenital heart defects were published by an interdisciplinary task force (3). Information is available, among others, from the academy for continuing medical education at the German Association for Pediatric Cardiologists (Deutsche Gesellschaft für Pädiatrische Kardiologie, Akademie für Fort-und Weiterbildung, Düsseldorf [theisen@dgpk.org]). Further information on certified cardiologists/pediatric cardiologists is available on the internet (4). Most adult patients with congenital heart defects are primarily being treated by their general practitioners and specialists in general internal medicine/cardiology but they should be referred at regular intervals—depending on the complexity of the underlying condition—to a center for adults with congenital heart disease or a cardiologist specializing in this subject. Accordingly, all doctors from all specialties involved in looking after such patients need to have basic knowledge of congenital heart defects.

This article provides an overview of the most important congenital heart defects, especially of common complications and therapeutic options in patients with congenital heart disease.

The aim is to enable readers to recognize predictable problems in adult congenital heart disease patients and prevent these, if possible. Since hardly any randomized controlled studies exist in this clinical area, the evidence level is comparatively low. The recommendations in this article are based mostly on the results of non-randomized prospective and retrospective studies and on expert opinion.

Method
The article is based on national and international guidelines (5, 6, e3). The authors also searched PubMed (www.ncbi.nlm.nih.gov) and selected articles on the basis of their own subjective assessment of their clinical relevance. The authors also used relevant textbooks and their own personal literature archives.
Overview of common heart defects in adult patients with congenital heart disease

Figures 1–4 provide an overview of the anatomy of common congenital heart defects. Further information is summarized in a supplementary table (available at: http://www.aerzteblatt.de/v4/plus/down.asp?id=7663). This table also contains information about the clinical presentation and common problems in the long term as well as therapeutic considerations for the most important congenital heart defects.

In addition to a merely anatomical description, congenital heart defects should be categorized according to their complexity and the presence of cyanosis (Box 1). This is of fundamental importance for planning follow-up care and in gauging what problems and complications might be expected (6). Except in rare conditions (early correction of uncomplicated patent ductus arteriosus and uncomplicated secundum atrial septal defects, which may be considered “cured,”) patients with congenital heart defects usually require lifelong care in order to secure an optimal outcome. According to current guidelines, patients with congenital heart defects of low complexity should be seen at a center for adults with congenital heart disease at least once, in order to decide on their further management. Patients with congenital heart defects of moderate and high complexity should be referred to a center and be assessed regularly (6).

Cardiac problems during follow-up

Interventions and repeat interventions

Although most patients undergo surgery in childhood, further surgical or interventional procedures are often required in the long term. The decision about the optimal timing of the intervention is a particular challenge in adults with congenital heart disease. In some cases, interventions are necessary before patients develop any symptoms, so as to prevent irreversible damage to the cardiac muscle and pulmonary vascular disease. On the other hand, procedures where reinterventions are anticipated (for example, implantation of a homograft or a biological valve replacement) should not be performed at an unnecessarily early stage.

Occasionally, congenital heart defects are detected only in adulthood. Apart from valvular disease (especially bicuspid valve), the most common abnormalities are atrial septal defects, occasionally Ebstein’s anomaly, and, more rarely, complex defects such as congenitally corrected transposition of the great arteries without associated abnormalities. If cardiac anatomy is unclear or ventricular dilatation and dysfunction occur (e.g., right ventricular dysfunction in the setting of atrial septal defects or partial anomalous pulmonary venous connection [PAPVC]; left ventricular dysfunction, e.g., in ductus arteriosus), physicians should consider an undetected congenital heart defect, even in adults, and seek further diagnostic evaluation.

A particular challenge is posed by patients with complex congenital heart defects in whom surgery was contraindicated for historical reasons or in whom only
alleviating procedures have been performed, and in whom it needs to be considered whether surgical correction is still possible in adult life.

Arrhythmias

Arrhythmias are a common late complication in adults with congenital heart defects, owing to the underlying cardiac disorder, hemodynamic problems, and surgical corrections resulting in scarring (7, e4). They range from bradycardiac arrhythmias to atrial tachycardias to life threatening ventricular tachycardias or atrial fibrillation.

In patients with symptoms such as palpitations, tachycardia, dizziness, and syncope, but also signs of heart failure, arrhythmias should be considered. We will now present some examples of cardiac abnormalities that predispose to arrhythmias.

Tetralogy of Fallot (ToF)—Patients with corrected ToF are predisposed to atrial and ventricular arrhythmias and are at increased risk for sudden cardiac death (about 2.5% per 10 years) (6). The substrate for these arrhythmias is represented by myocardial fibrosis (e5) and re-entry circuits around ventriculotomy and atriotomy scars or the VSD (ventricular septal defect) patch (e6). Identified risk markers for sudden cardiac death include a wide QRS complex >180 ms (8), severe pulmonary insufficiency, right and left ventricular dysfunction, extensive myocardial fibrosis, late corrective surgery, and palliative shunts that have been in place for a long time. The importance of regular Holter-ECGs is the subject of controversial debate (e7). In selected symptomatic patients, electrophysiological studies can yield prognostic information, and, in ventricular re-entry circuits ablation treatment may be an option (9). Implantation of an implantable cardioverter defibrillator (ICD) is widely accepted in the context of secondary prophylaxis (“survived sudden cardiac death”), but criteria for primary preventive implantation are currently not well defined (e8).

Patients with transposition of the great arteries (TGA) after atrial switch surgery—Most of the patients with TGA develop relevant cardiac arrhythmias during their lifetime. These include bradycardiac and tachycardiac arrhythmias. As a result of the procedure and owing to scarring in the atrial region, more than 10% of patients with TGA develop early postoperative bradycardiac arrhythmias after atrial switch operation, which require pacemaker insertion (e9). It has been reported that 20 years postoperatively, only 40% of patients are in sinus rhythm (e10). Symptoms of arrhythmia, heart failure, and documented arrhythmias in the patient’s medical history have been identified as a predictor for sudden cardiac death, whereas findings from resting ECG and Holter-ECG monitoring do not seem to carry prognostic information (10). Little is currently known about the importance of electrophysiological examination for prognostic assessment.

Patients with functional single ventricle after Fontan repair—Recurrent supraventricular arrhythmias are common in patients after Fontan repair, especially patients who have undergone the classic Fontan procedure (connection between right atrium and pulmonary artery). In such patients, intra-atrial

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macro-re-entry circles with a long cycle-length are common, which are hemodynamically poorly tolerated. Patients with recurrent supraventricular tachycardias should be referred to an experienced center for electrophysiological evaluation (11). This applies in particular to patients after a “classic” Fontan repair, whose prognosis is limited (5 year survival 85%) (e11, e12).

Heart failure
In the long term, the development of heart failure is common in patients with congenital heart disease, especially those with complex abnormalities and systemic right ventricle. It has been described that heart failure develops in 22% of patients with congenitally corrected transposition of the large vessels, 32% of TGA patients after atrial switch surgery, and up to 40% of patients after Fontan repair (12). Potentially correctable causes (such as hemodynamically relevant shunts, valve insufficiency and stenosis, and other intracardial or vascular obstructions, arrhythmias, pulmonary arterial hypertension) should be identified and treated accordingly. Angiotensin converting enzyme (ACE) inhibitors and beta blockers are used empirically, but evidence supporting the prognostic benefit of these medications is lacking in adult patients with congenital heart defects (13). Transplantation is the treatment of last resort in patients with end-stage heart failure, but this may be technically demanding and has an increased risk compared with other groups of patients (e13).

Pulmonary hypertension and Eisenmenger’s syndrome
Pulmonary arterial hypertension (PAH) is a common complication in adults with congenital heart defects, especially with uncorrected shunt lesions and unprotected pulmonary circulation or after a late repair. Patients with PAH after shunt closure pathophysiologically resemble patients with idiopathic PAH, but severe PAH in patients who have not had surgery usually results in shunt reversal with Eisenmenger’s syndrome (a combination of PAH and cyanosis). Endothelin receptor antagonists, phosphodiesterase inhibitors, and prostacyclines are effective drugs for the treatment of PAH (14, e14); however, far fewer data exist for adults with congenital heart defects than for those with idiopathic PAH.

Infectious endocarditis
In principle, all congenital heart defects are associated with an increased lifetime risk for infectious endocarditis (IE). However, since it has to be assumed that most cases of IE are not caused by medical or dental interventions, and since the assumption that administration of an antibiotic can prevent IE has never been conclusively proved in humans, the current guidelines of the international societies recommend IE prophylaxis only in high-risk patients and patients who have previously had IE (15). This high-risk group comprises patients with complex, usually cyanotic conditions, patients with implanted prosthetic valves, and patients with residual defects in immediate proximity to...
implanted patches or conduits. Furthermore, prophylaxis against endocarditis for 6 months is recommended after cardiac surgery with implantation of patch or conduit material and after catheter intervention using implants.

By type of intervention, prophylaxis is restricted to dental procedures that may be accompanied by injury to the mucosa, gingiva, or apical dental region, as well as bronchoscopy with incision of the mucosa or biopsy, as well as tonsillectomies and adenotomies. We cannot emphasize enough that patients need to be reminded to observe good oral hygiene and visit their dentist regularly.

It is of particular importance in adults with congenital heart disease to consider endocarditis early in case of suspicious symptoms (raised temperature, night sweats, embolism, etc) and initiate the necessary diagnostic steps early on—especially echocardiography and blood cultures, before an antibiotic is given (e15, e16).

**Common non-cardiac problems in the context of aftercare for adults with congenital heart defects**

**Contraception and pregnancy**

Cardiovascular disorders are currently the most common cause of maternal mortality during pregnancy (16). An individual risk assessment in any adult woman with a congenital heart defect should be done early on, in order to prevent unwanted high-risk pregnancies.

The physiological changes and their possible implications for different congenital heart defects have just been discussed in detail in a review article in Deutsches Ärzteblatt International (16). Because of the increase in cardiac volume and circulating blood volume, stenotic abnormalities and increased pulmonary vascular resistance are particularly problematic. Cyanotic abnormalities also present a particular problem, because of the drop in peripheral vascular resistance and the increase in right-to-left shunts. The combination of severe pulmonary arterial hypertension and cyanosis (as in Eisenmenger’s syndrome) is associated with particularly high maternal mortality of up to 50% (17); the deaths often occur during the puerperium. Generally, symptoms of heart failure (NYHA >2), reduced left ventricular function, previous cardiovascular problems during pregnancy, and severe pulmonary valve insufficiency—especially combined with reduced right ventricular function—are all associated with increased risk for pregnant women (18–20). Patients with Marfan syndrome and a dilated aortic root (>4 cm) have an increased risk for dissection (5). Treating patients with mechanical prosthetic valves is problematic because of the need for adequate anticoagulation and the increased risk for fetal abnormalities owing to treatment with vitamin K antagonists (for example, Marcumar [phenprocoumon]). Alternative strategies for anticoagulation are not easily controlled with standard coagulation tests; they are associated with an increased risk of thrombosis, barely evidence based, and thus not without problems. The management of patients with a

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**BOX 1**

**Complexity of common heart defects**

- **Heart defects of low complexity**
  - Uncorrected heart defects
    - Isolated small atrial septal defects (ASD II)
    - Isolated restrictive ventricular septal defects (VSD)
    - Mild pulmonary stenosis (gradient <30 mm Hg)
    - Restrictive persistent ductus arteriosus
    - Isolated congenital aortic stenosis
    - Isolated congenital mitral valve disorder

- **Heart defects after correction**
  - After closure of ASD (except for atrioventricular defect)
  - After uncomplicated closure of VSD
  - After closure of persistent ductus arteriosus

- **Heart defects of moderate complexity**
  - Atrioventricular septal defects (AVSD) / ASD I
  - Sinus venosus defects
  - Aortic isthmus stenosis
  - Ebstein’s anomaly
  - Tetralogy of Fallot
  - Ventricular septal defects with associated anomalies
  - Moderate and higher grade pulmonary stenosis or regurgitation
  - Anomalous pulmonary venous connection
  - Persistent ductus arteriosus (uncorrected)

- **Heart defects of high complexity**
  - Cyanotic heart defects (independent of anatomy)
  - Complex heart defects with functionally univentricular anatomy
  - Patients after Fontan repair
  - Transposition of the great arteries (TGA)
  - Any form of discordant connection between atria, ventricles, and great arteries
  - Patients with implanted conduits

ASD, atrial septal defect; AVSD, atrioventricular septal defect; TGA, transposition of the great arteries; VSD, ventricular septal defect. *1 Adapted from Warnes et al (6)
Cyanosis and erythrocytosis

Patients with uncorrected complex cardiac defects and patients after palliative procedures (for example, Glenn surgery) may present with cyanosis. Chronic cyanosis is characterized by notably reduced exercise capacity, frequent infections (for example, endocarditis and cerebral abscesses), and multiple end organ damage in the sense of a multiorgan disorder (e17). Typical complications in addition to the infections include hemorrhage, thromboembolism, arrhythmias, impaired renal function, gallstones, and joint problems. A comprehensive explanation of the problems would exceed the scope of this article.

Erythrocytosis develops secondary to chronic hypoxia, is associated with an increased oxygen carrying capacity of the blood, and improves tissue oxygenation; however, it requires sufficient availability of iron. Preventing iron deficiency is of particular importance (e18). Phlebotomy should be limited to patients with confirmed symptoms of hyperviscosity (headache, dizziness, impaired vision, paresthesias, muscle aches, etc.) after iron deficiency has been excluded; the hematocrit is usually in excess of 65% in such cases (5). However, even hematocrit values above 70% are commonly tolerated without any symptoms. Repeated unnecessary phlebotomy will then result in iron deficiency, reduced exercise capacity, and increased risk of stroke (e17, e19). Dehydration needs to be avoided in such patients; if it does develop it requires immediate treatment.

Cyanotic patients, especially those with Eisenmenger’s syndrome, are sometimes placed in grave danger by simple surgical procedures, but also by other medical interventions, if their particular characteristics are not considered accordingly. Any medication or surgical treatment, including semi-invasive and invasive examinations should be coordinated with a center for adults with congenital heart disease.

Non-cardiac surgical interventions

Many adult patients with congenital heart defects require general surgery during their lifetime. Because of the range of such procedures, and also depending on their urgency, medical centers in primary as well as secondary care are often confronted with this problem. The risk of a procedure may vary depending on the underlying defect and associated problems. The treating surgeon and anaesthetist should consider the influence of anesthesia, mechanical ventilation, possible required vasoactive substances, and blood volume shifts on the underlying cardiac physiology and consult a cardiologist with experience in adult patients with congenital heart disease. High-risk patients (Fontan repair, severe PAH, cyanosis, complex heart defects especially in the presence of heart failure, and others) should be treated in specialized centers.

Psychosocial aspects, choice of job, and sports

The quality of life of adult patients with congenital heart disease is impaired by:

- Reduced exercise capacity (21)
- Recurring arrhythmias (22, 23)
- Cosmetic impairments (e20)
- Repeatedly required catheter interventions or surgical procedures.

### Table 1

<table>
<thead>
<tr>
<th>Low risk (risk of cardiac complications or death &lt;1%)</th>
<th>Moderate risk (risk of cardiac complications or death = 1–5%)</th>
<th>High risk (risk of cardiac complications or death &gt;5%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>– Left-right shunt without PAH</td>
<td>– Moderate aortic or mitral valve stenosis at most</td>
<td>– Severe aortic or mitral valve stenosis</td>
</tr>
<tr>
<td>– Status post correction of tetralogy of Fallot without relevant residual defects (no relevant pulmonary insufficiency or right ventricular dysfunction)</td>
<td>– Cyanotic abnormalities without PAH (but high fetal risk)</td>
<td>– Eisenmenger syndrome or PAH (very high pregnancy risk: maternal mortality 30–60%)</td>
</tr>
<tr>
<td>– Status post correction of aortic stenosis without relevant residual defects (residual stenosis, aneurysm)</td>
<td>– Univentricular heart with good ventricular function (particular aftercare required during pregnancy)</td>
<td>– Univentricular heart with reduced ventricular function</td>
</tr>
<tr>
<td>– Bicuspid aortic valve without associated problems</td>
<td>– Marfan syndrome without significant aortic dilatation</td>
<td>– Marfan syndrome with aortic dilatation</td>
</tr>
<tr>
<td>– Status post biological valve replacement (with normal prosthetic and ventricular function)</td>
<td>– Systemic right ventricle (ccTGA, after atrial switch operation) with intact systemic ventricular function</td>
<td>– Systemic right ventricle (ccTGA, after atrial switch operation) with reduced systemic ventricular function</td>
</tr>
<tr>
<td>– Asymptomatic mitral or aortic valve insufficiency with intact left ventricular function</td>
<td>– Uncorrected aortic isthmus stenosis (depending on gradient)</td>
<td></td>
</tr>
<tr>
<td>– Mild to moderate pulmonary stenosis</td>
<td></td>
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</tbody>
</table>

PAH, pulmonary arterial hypertension; ccTGA, congenitally corrected transposition of the great arteries. Modified from (16, e22)
Furthermore, patients often have fears and anxieties regarding possible complications, increasing disability, and reduced life expectancy. These aspects have thus far not received much attention, but they are increasingly gaining importance. Appropriate psychosocial care is of great importance and should be provided at supraregional centers for adult patients with congenital heart disease. In addition, concrete questions regarding participation in sports and choice of job position should be answered. This requires experience and the objective assessment of maximal exercise capacity in the context of spiroergonomic examinations.

The organization of healthcare service structures

The treatment of patients with heart defects of moderate to high complexity should be coordinated by supraregional centers for adult patients with congenital heart disease. In a position paper published in 2001 (e21) it was stipulated that a supraregional center should provide care for a population of 5 to 10 million. For Germany, this would mean that 8 to 16 supraregional centers are needed. Cardiologists at such centers should have particular experience in the treatment of adult patients with congenital heart disease (according to the American model, a minimum of 2 years’ experience with adult patients with congenital heart disease, ideally 5 or more years). Furthermore, an experienced cardiothoracic surgical team should be available, as should be the option of electrophysiological investigations and treatments as well as all other diagnostic methods (cardiac MRI, CT, and invasive diagnostic approaches).

The interdisciplinary task force for adult patients with congenital heart disease that was mentioned earlier has published recommendations for improving the quality of interdisciplinary care for adults with congenital heart defects (2), which mainly follow the North American suggestions. The cooperation of basic care provided by general practitioners, regional services providing maximal care, and supraregional centers are explained and the requirements of the different structures are laid out. Good communication between the general practitioner, the cardiologist in private practice, and the center for adult patients with congenital heart disease is of crucial importance. Close collaboration between pediatric and adult cardiologists is also vital in order to provide a seamless transition for patients from pediatric cardiology services to adult cardiology services.

Conflict of interest statement

The authors declare that no conflict of interest exists.

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REFERENCES


KEY MESSAGES

- A minority of adult patients with congenital heart defects can be regarded as “cured”, and long term complications are common.
- In spite of surgical interventions in childhood, further surgical or other interventional procedures are often required in the longer term.
- Patients with heart defects of moderate to high complexity should be referred to specialized centers for adults with congenital heart disease and should be evaluated regularly in those centers.
- Since cardiovascular disorders are currently the most common cause for maternal mortality during pregnancy, an individual risk assessment should be made in women with congenital heart defects early on, in order to prevent unwanted high-risk pregnancies or ensure optimal care during the pregnancy.
- Good communications between patients’ general practitioners, cardiologists in private practice, and specialist centers, as well as close cooperation between pediatric and adult cardiologists is required in order to ensure optimal treatment for adult patients with congenital heart defects.


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A supplementary table containing further information on congenital heart defects is available at:
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eReferences

REVIEW ARTICLE

Congenital Heart Defects in Adulthood

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The supplementary table containing further information on congenital heart defects will be available here on Wednesday, July 27, 2011.