Pregnancy in a woman with congenitally corrected transposition of the great arteries

A 28-YEAR-OLD WOMAN with a history of complex congenital heart disease presents to an urgent care center with nausea and emesis, which have been worsening for 3 days. She also reports 2 weeks of constipation, with bowel movements occurring approximately every 3 days. Her most recent medical contact was 3 weeks ago for management of sinusitis that had failed to improve with azithromycin therapy. One week before the current presentation, she was prescribed trimethoprim-sulfamethoxazole for this issue. She says she has had no recent fever, lightheadedness, shortness of breath, or chest pain.

Her medical history is remarkable for congenitally corrected transposition of the great arteries (CC-TGA) with a ventricular septal defect and subpulmonic stenosis. At 5 years of age, she underwent initial surgical repair that included subpulmonic resection to relieve right ventricular outflow obstruction and closure of the ventricular septal defect to prevent chronic left-to-right shunting, a situation that could eventually lead to pulmonary vascular remodeling and pulmonary hypertension (Eisenmenger syndrome).

At age 24, she underwent implantation of a mechanical tricuspid valve to prevent the development of systemic ventricular dysfunction related to chronic, severe systemic atrioventricular valve regurgitation. Her postoperative course was complicated by complete heart block, requiring placement of a dual-chamber permanent pacemaker. Transthoracic echocardiography performed 2 months after the valve replacement demonstrated normal systemic ventricular function and no significant prosthetic valve stenosis or regurgitation. She has not undergone repeat cardiac imaging since that time.

FIRST STEP IN EVALUATION

1. Based on the patient’s presentation, what is the most appropriate next step?

☐ Computed tomography of the abdomen
☐ Right heart catheterization
☐ Urine or serum pregnancy test
☐ Abdominal radiography

A pregnancy test is essential before a woman of childbearing age undergoes any diagnostic testing involving radiation exposure, especially if she has symptoms that could possibly be due to pregnancy. While all the above options may help with diagnosis, computed tomography, right heart catheterization, and abdominal radiography require exposure to radiation. The most appropriate and cost-effective first test to evaluate this patient’s symptoms is a urine or serum pregnancy test.

CASE CONTINUED: A POSITIVE PREGNANCY TEST

The patient says she has had no abdominal cramping, bloating, or vaginal bleeding since her last menstrual period, which was about 7 weeks before this presentation.

Her pulse is 68 beats per minute, blood pressure 112/70 mm Hg, oral temperature 36.8 °C (98.2 °F), and respiratory rate 16 breaths per minute.
She is alert, well-nourished, and in no acute distress. Her mucous membranes are moist. Her abdomen is without tenderness, rebound pain, guarding, or palpable masses. Auscultation of the heart reveals a crisp mechanical S1 and a grade 2 of 6 systolic ejection murmur at the left lower sternal border.

A rapid serum quantitative human chorionic gonadotropin test is positive for pregnancy.

**HEMODYNAMIC CHANGES OF PREGNANCY**

Which of the following is a normal hemodynamic adaptation of pregnancy?

- □ Increased cardiac output
- □ Reduced blood volume
- □ Decreased heart rate
- □ Increased systemic vascular resistance

Cardiac output increases during pregnancy, as the maternal cardiovascular system must undergo dynamic adaptations to meet increased metabolic demands and to support fetal growth and development through the uteroplacental circulation. Because nitric oxide is upregulated by circulating gestational hormones and the placental bed has low resistance, systemic and pulmonary vascular resistance decreases by about 30% early in pregnancy, resulting in decreased mean arterial blood pressure.1 Beginning early in the second trimester, maternal circulating plasma volume increases by 30% to 50%, peaking at the beginning of the third trimester and resulting in relative anemia and reduced blood viscosity. Heart rate also increases during that time due to increased sympathetic tone.2,3 Also related to sympathetic tone and increased preload, stroke volume increases by an average of 25%. These physiologic changes cause cardiac output to increase steadily throughout pregnancy, peaking in the third trimester at about 50% above prepregnancy levels.4

Labor and delivery bring additional maternal hemodynamic changes that can challenge the cardiovascular system. Pain and anxiety lead to increased mean arterial blood pressure and oxygen consumption. With each uterine contraction, circulating blood volume further increases by up to 500 mL. These rapid changes result in a further rise in maternal cardiac output to about 80% above prepregnancy levels, in some women reaching more than 10 L/min.5 Some of these hemodynamic shifts can be mitigated with the use of adequate analgesia.6

After delivery, with the uterus no longer compressing the inferior vena cava, venous return increases. This, in combination with intrapartum blood shifts related to uterine contractions and a sudden increase in systemic vascular resistance following delivery of the placenta, leads to relative elevation of intracardiac filling pressures.6 Spontaneous diuresis occurs in most women over the first 24 to 48 hours after delivery. Maternal hemodynamics typically require 2 to 6 weeks to return to baseline after vaginal delivery, and usually longer after cesarean delivery.

**EFFECTS OF TRANPOSITION OF THE GREAT ARTERIES**

Which of the following statements about TGA is correct?

- □ In infants with dextro-transposition, a shunt is essential for survival

![Figure 1. (Left) Congenitally corrected transposition of the great arteries (CC-TGA). The right ventricle pumps oxygenated blood in the systemic circulation, while the left ventricle pumps deoxygenated blood in the pulmonary circulation. No intervention may be needed. (Right) Dextro- or D-TGA. Pulmonary and systemic blood flows are separate, leading to rapid clinical deterioration in the first day of life unless a communication between the circulations exists or is created.](https://www.ccjm.org)
A patient with CC-TGA may develop heart failure due to dysfunction of the systemic right ventricle.

A patient with CC-TGA may develop regurgitation of the systemic atrioventricular (morphologic tricuspid) valve.

Heart block can occur in up to a third of patients with CC-TGA.

All of the above are correct.

In dextro-transposition, two separate circulations

In TGA, the aorta arises from the right ventricle and the pulmonary artery from the left ventricle (Figure 1). In complete TGA (dextro- or D-TGA), the normal relationship between the atria and the ventricles is preserved: the right atrium empties into the right ventricle through the tricuspid valve, and the left atrium empties into the left ventricle through the mitral valve. Thus, the pulmonary circulation is separate from the systemic circulation. As a result, infants develop cyanosis and rapidly clinically deteriorate within the first day of life unless a communication exists through either an atrial septal defect, ventricular septal defect, patent ductus arteriosus, or atrial septostomy.

In CC-TGA, normal oxygenation

In CC-TGA, also known as levo- or L-TGA, the right atrium empties into the left ventricle through the mitral valve and the left ventricle in turn empties into the pulmonary artery, while the left atrium empties into the right ventricle through the tricuspid valve, and the right ventricle in turn empties into the aorta.

The condition is described as “congenitally corrected” because, unlike D-TGA, the pulmonary and systemic circulations exist in series, and hence, oxygenated blood does reach the systemic circulation. In fact, the only functional difference between CC-TGA and normal anatomy is that the right ventricle is responsible for pumping oxygenated blood into the systemic circulation while the left ventricle pumps deoxygenated blood into the pulmonary circulation. Hence, patients with isolated CC-TGA do not commonly require surgical or transcatheter intervention.

Cardiac output peaks in the third trimester at about 50% above prepregnancy levels.

Figure 2. Electrocardiogram demonstrating complete heart block with a junctional escape rhythm in a patient with CC-TGA. The typical findings in patients with CC-TGA found here include Q waves in the inferior leads (II, III, and aVF) and right limb lead (aVR), and the absence of Q waves in V5 and V6.

From Yoon et al, reference 9.
CC-TGA: Associated anomalies, conduction abnormalities

CC-TGA is associated with other congenital cardiac anomalies in up to 90% of patients. The most common include deformities of the tricuspid valve, ventricular septal defects, and subpulmonic ventricular outflow obstruction. CC-TGA can also be associated with abnormalities of the cardiac conduction system.

The major arrhythmogenic complication of CC-TGA is conduction system disease. Due to the presence of atrioventricular discordance, the atrioventricular node is unable to connect normally to the bundle of His. A second anteriorly positioned atrioventricular node, therefore, typically develops and gives rise to an elongated His bundle that is vulnerable to fibrosis over time. The risk of complete heart block is around 2% per year, with a cumulative lifetime incidence of 30%. Patients with CC-TGA should undergo a surveillance electrocardiogram at least yearly to screen for development of conduction system disease (Figure 2).

CASE CONTINUED: THE PATIENT CONSULTS A SPECIALIST

Once pregnancy is confirmed, the patient is surprised, as she had been regularly using a barrier method of contraception. She is excited, as this is her first pregnancy, but becomes concerned about potential complications that she might experience due to her preexisting heart disease. She promptly schedules an appointment with her adult congenital heart disease specialist to discuss this.

CARDIOVASCULAR RISK IN PREGNANCY

Which of the following conditions pose the highest risk of pregnancy-associated maternal mortality?

- Eisenmenger syndrome
- Mechanical prosthetic valve
- Subaortic atroventricular valve regurgitation with preserved subaortic ventricular function
- Moderate pulmonary valve stenosis

Eisenmenger syndrome poses the highest risk of maternal mortality of the conditions listed.

Hemodynamic changes during pregnancy, delivery, and the postpartum period may be poorly tolerated in women with congenital or acquired heart disease. Cardiovascular conditions are the leading cause of maternal mortality in the United States: 25% to 30% of pregnancy-related deaths between 2011 and 2016 were attributable to preexisting cardiovascular disease.

All women of childbearing age with preexisting heart disease should receive regular counseling regarding their individual risk of adverse pregnancy-related outcomes. The updated 2018 American Heart Association-American College of Cardiology guideline recommends that all women with congenital heart disease receive pregnancy counseling with input from a congenital heart disease specialist to determine maternal cardiac, obstetric, and fetal risks, as well as potential long-term risks to the mother (recommendation class 1 [strong recommendation], evidence level C [based on expert consensus, case studies, or standard of care]).

It is vitally important for physicians to understand the pregnancy risks associated with specific cardiac lesions and disease states. The risk for adverse pregnancy-related events in women with preexisting cardiovascular disease can be estimated based on the type and severity of the underlying cardiac disease, as well as preconception functional status.

Several clinical tools are available to assist in patient-specific preconception risk assessment. The Cardiac Disease in Pregnancy II (CARPREG II) risk score, which was derived from 1,938 pregnancies between 1994 and 2014, is the most contemporary. Ten predictors of adverse pregnancy-related events were identified and correlated to a weighted-point system to assist in counseling patients:

- Previous cardiac events or arrhythmias: 3 points
- Baseline New York Heart Association class III or IV heart failure or cyanosis: 3 points
- Mechanical valve: 3 points
- Ventricular dysfunction: 2 points
- High-risk left-sided valve disease or left ventricular outflow tract obstruction: 2 points
- Pulmonary hypertension: 2 points
- Coronary artery disease: 2 points
- High-risk aortopathy: 2 points
- No prior cardiac intervention: 1 point
- Late pregnancy assessment: 1 point.

Isolated CC-TGA does not commonly require surgical or transcatheter intervention.
Patients with CC-TGA should have an ECG at least yearly to screen for conduction system disease.

The risk of a primary cardiac event is calculated from the total points:

- 0–1 points: 5%
- 2 points: 10%
- 3 points: 15%
- 4 points: 22%
- > 4 points: 41%.

The World Health Organization has also developed a maternal cardiovascular risk classification, which is based on expert opinion (Table 1).\(^\text{13}\)

Echocardiography, cardiac magnetic resonance imaging, and exercise testing may be necessary to better define risk and understand the patient’s anatomy, cardiac physiology, and functional status.

Genetic counseling

Patients with congenital heart disease or other inherited forms of preexisting cardiac disease should be offered genetic counseling. The

### TABLE 1

**World Health Organization classification of maternal cardiovascular risk**

**Class I**

Uncomplicated, small or mild pulmonary stenosis, patent ductus arteriosus, or mitral valve prolapse; successfully repaired simple lesions (atrial or ventricular septal defect, patent ductus arteriosus, anomalous pulmonary venous return)

Cardiovascular risk: No increased risk of maternal mortality, possible mild increase in morbidity

**Class II**

Unrepaired atrial or ventricular septal defect; repaired tetralogy of Fallot; supraventricular tachycardia

Cardiovascular risk: Small increased risk of maternal mortality, moderate increase in morbidity

**Class II–III**

Mild left ventricular dysfunction; native or tissue valvular heart disease not considered World Health Organization class I or IV; Marfan syndrome without aortic dilation; aorta less than 45 mm in association with bicuspid aortic valve; repaired coarctation of the aorta; hypertrophic cardiomyopathy

Cardiovascular risk: As in class II or III, depending on individual

**Class III**

Mechanical valve; systemic right ventricle with good or mildly reduced function; Fontan circulation without complication; unrepaired cyanotic heart disease; other complex congenital heart disease; aortic dilation 40 to 45 mm in Marfan syndrome; aortic dilation 45 to 50 mm in bicuspid aortic valve disease; previous peripartum cardiomyopathy with complete recovery of ventricular function

Cardiovascular risk: Significantly increased risk of maternal mortality or severe morbidity. Expert counseling required. If pregnancy continues, intensive specialist cardiac and obstetric monitoring needed throughout pregnancy, childbirth, and for 6 weeks afterwards

**Class IV**

Pulmonary arterial hypertension from any cause; severe systemic ventricular dysfunction (ejection fraction < 30%, New York Heart Association functional class III-IV); systemic right ventricle with moderate or severely reduced function; Fontan circulation with any complication; severe mitral stenosis; severe symptomatic aortic stenosis; Marfan syndrome with aortic dilation more than 45 mm; aortic dilation more than 50 mm in bicuspid aortic valve disease; native severe coarctation of the aorta; previous peripartum cardiomyopathy without complete recovery of ventricular function

Cardiovascular risk: Pregnancy is contraindicated because of extremely high risk of maternal mortality or severe morbidity. If pregnancy occurs, discuss termination. If pregnancy continues, care as for class III

Information from reference 13.
prevalence of congenital heart disease is 0.8% in the general population; risk increases up to 10-fold if congenital heart disease is also present in a first-degree relative. Rates of transmission are higher when the mother is affected compared with the father. In patients with autosomal dominant conditions, such as Marfan syndrome, there is a 50% chance of transmission to the fetus.

**COMPLICATIONS IN PREGNANCY IN WOMEN WITH CC-TGA**

What is the most common pregnancy-associated cardiovascular complication in women with CC-TGA?

- Symptomatic heart failure
- Worsening systemic atrioventricular valve regurgitation
- Arrhythmia
- Endocarditis

With medical management, most women with CC-TGA reach childbearing age and are healthy enough to consider pregnancy. Patients and their physicians should understand their specific risks so that they can carefully consider the appropriateness and timing of pregnancy.

Although helpful, global risk assessment tools such as CARPREG II developed to predict pregnancy-associated maternal cardiovascular risk typically do not account for the presence of a systemic right ventricle. In the maternal World Health Organization classification system, patients with a systemic right ventricle with normal or mildly reduced function are categorized as class III, consistent with significantly increased risk of maternal mortality or severe morbidity, and patients with moderately to severely reduced systemic right ventricular dysfunction should be advised against pregnancy (class IV).

**Arrhythmia risk poses challenges**

The most common pregnancy-associated cardiovascular complication seen with CC-TGA is supraventricular arrhythmia. It presents a management challenge, as few antiarrhythmic medications have been studied for safety and efficacy in pregnancy. Beta-blockers, especially metoprolol succinate and labetalol, are acceptable in pregnancy. However, they should be used with caution in patients with CC-TGA, who have an increased predisposition for heart block, which these drugs can exacerbate. In a minority of cases, supraventricular arrhythmias can lead to maternal hypoxia, which can be associated with premature delivery and low birth-weight infants.

**Risk of heart failure depends on baseline function**

Systemic right ventricular dysfunction and resulting congestive heart failure symptoms are common by the third decade of life and increase in frequency as patients age. The World Health Organization classification of risk in patients with a systemic right ventricle is derived from the idea that the systemic right ventricle is not designed to support systemic blood flow, predisposing patients to systemic ventricular dysfunction and other cardiovascular complications of pregnancy. Although pregnancy-related changes in cardiovascular hemodynamics may provoke new-onset or worsening systemic ventricular dysfunction, most women with CC-TGA who have normal systemic ventricular function before conception do not develop symptomatic heart failure during pregnancy or postpartum.

**CASE CONTINUED: ANTICOAGULATION IN PREGNANCY**

The patient engages in detailed discussions with a specialist in adult congenital heart disease and the high-risk obstetrics team about her potential risks during pregnancy. She next asks about additional considerations regarding her mechanical systemic atrioventricular valve. She currently takes warfarin 7.5 mg daily to prevent mechanical valve thrombosis. Her international normalized ratio (INR) is within the therapeutic range.

How should anticoagulation be immediately handled in this patient?

- Continue warfarin at her current dose of 7.5 mg daily
- Because anticoagulation can lead to fetal...
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to experience mild dyspnea with activity. By the 34th week of pregnancy, she notes mild lower-extremity edema. She says she has no paroxysmal nocturnal dyspnea, palpitations, cough, or chest pain. Her blood pressure is 114/70 mm Hg. Serial echocardiograms and NT-proBNP measurements remain unchanged, despite her persistent and worsening symptoms.

Based on the patient's current symptoms and echocardiogram, what is the most appropriate next step in her management?

- Start a diuretic to address signs of heart failure
- Perform right-heart catheterization to assess heart failure
- Perform transesophageal echocardiography to evaluate for prosthetic valve thrombosis
- No change in management at this time; continue to monitor closely

A change in management is not necessary at this time, but the patient should continue to be monitored closely.

A mechanical heart valve is associated with elevated maternal and fetal risk

Symptoms such as shortness of breath, orthopnea, lower-extremity edema, and sinus tachycardia may be considered normal in some women during pregnancy owing to expected hemodynamic changes. Likewise, echocardiographic chamber size, transvalvular flow velocity, and valvular regurgitation are also expected to increase as pregnancy progresses. It can be difficult to differentiate these symptoms and imaging changes from cardiac decompensation in women with heart disease.²⁻⁵

Serial NT-proBNP measurement can be helpful in differentiating whether worrisome symptoms and echocardiographic changes should be considered pathologic. On the other hand, reduced ventricular function is never considered normal during pregnancy. As such, it is important to evaluate each patient's cardiac function before determining whether a change in management strategy is necessary.

**CASE CONTINUED: A HEALTHY BABY**

Given her stable cardiac imaging and NT-proBNP measurements, it is deemed unlikely that the patient's symptoms represent cardiac decompensation. She continues to be monitored closely without additional medical therapy.

As pregnancy progresses, she develops severe pelvic pain due to suspected symphysis pubis separation, as well as hemorrhoids, for which she is managed supportively.

Warfarin is continued until 35 weeks of gestation, at which time she is transitioned to low-molecular-weight heparin.

At 37 weeks, she presents with spontaneous rupture of membranes while fully anticoagulated with low-molecular-weight heparin. Despite counseling regarding bleeding risks, she opts for a primary cesarean delivery because of her concerns about the potential of worsening symptoms related to pubic symphysis separation, as she cannot be managed with neuraxial anesthesia owing to her anticoagulated state.

She undergoes primary cesarean delivery under general anesthesia. The procedure is uncomplicated and results in the delivery of a healthy infant boy with Apgar scores of 9 (of a maximum possible 10) at both 1 and 5 minutes.

Her postpartum course is uneventful. Anticoagulation with intravenous heparin is reinitiated 12 hours after delivery. Warfarin is restarted on postoperative day 3, while continuing intravenous heparin until a therapeutic INR is achieved. She is discharged on postoperative day 7 receiving warfarin (INR 2.5) and aspirin 81 mg daily. A nonhormonal intrauterine device is recommended for contraception.
PREGNANCY IN CONGENITAL HEART DISEASE

- Patients with CC-TGA can carry a pregnancy successfully without complication. The most common pregnancy-associated cardiovascular complication in patients with CC-TGA is supraventricular arrhythmia. They are also at risk for worsening systemic ventricular dysfunction and decompensated heart failure.

- Managing anticoagulation in women with a mechanical heart valve to prevent valve thrombosis during pregnancy can be difficult. When 5 mg or less of warfarin is required to achieve therapeutic anticoagulation, it should be continued even in the first trimester. If low-molecular-weight heparin is needed, anti-Xa levels should be followed closely to ensure therapeutic anticoagulation.

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