Critical Congenital Heart Disease Detection in the Screening Era: Do Not Neglect the Examination!

Deepika Sankaran, MD1  Heather Siefkes, MD, MSCI2  Frank F. Ing, MD, FACC, MSCI3
Satyan Lakshminrusimha, MD1  Francis R. Poulain, MD1

1 Division of Neonatology, Department of Pediatrics, University of California, Davis, Sacramento, California
2 Division of Pediatric Critical Care Medicine, Department of Pediatrics, University of California, Davis, Sacramento, California
3 Division of Pediatric Cardiology, Department of Pediatrics, University of California, Davis, Sacramento, California

Abstract

Pulse oximetry oxygen saturation (SpO2)-based critical congenital heart disease (CCHD) screening is effective in detection of cyanotic heart lesions. We report a full-term male infant with normal perfusion who had passed the CCHD screening at approximately 24 hours after birth with preductal SpO2 of 99% and postductal SpO2 of 97%. Detection of a loud systolic cardiac murmur before discharge led to the diagnosis of pulmonary atresia (PA) with ventricular septal defect (PA-VSD) by echocardiogram. The infant was transferred to a tertiary care center after initiation of prostaglandin E1 (PGE1) therapy. Throughout the initial course, he was breathing comfortably without respiratory distress or desaturations on pulse oximetry. We believe that this is the first documented report of PA missed by CCHD screening. Thorough and serial clinical examinations of the newborn infant proved vital in the timely diagnosis of this critical disease. We review the hemodynamics and the recent literature evaluating utility of CCHD screening in the diagnosis of PA-VSD. Pulse oximetry–based CCHD screening should be considered a tool to enhance CCHD detection with an emphasis on detailed serial physical examinations in newborn infants.

Keywords
► newborn
► congenital heart disease
► pulse oximetry
► CCHD screening
► pulmonary atresia
► PA-VSD
► oximetry screening
► false negative
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► critical congenital heart disease

Congenital heart diseases (CHDs) are the most common congenital malformations with an incidence of 5 to 10 per 1,000 live births, and a quarter of them qualify as critical CHDs (CCHD). Prior to universal oxygen saturation (SpO2)-based CCHD screening, up to 10% of all infant deaths were attributed to CHDs. Although a substantial number of infants with CCHDs could be detected by screening, an equal number of infants with CCHD may be missed. Acyanotic lesions, such as coarctation of the aorta, are commonly missed and present clinically within the first 2 weeks with rapid deterioration after the ductus arteriosus (PDA) closes. On the other hand, there are no published reports of pulmonary atresia (PA) that were missed by oximetry screening.
Patient Presentation

A term male infant was delivered in a community hospital by cesarean section due to category-II fetal heart tracing. The antenatal history was unremarkable except for gestational hypertension. The mother had normal fetal anatomy ultrasound at 20 weeks of gestation. The baby had routine care in the delivery room with Apgar scores of 7 and 9. He had normal physical examination except for left-sided undescended testis. The baby was breastfeeding well, had normal vital signs, and the subsequent physical examinations on the day of birth did not demonstrate any cardiac murmur.

Twenty-four hours after birth, the infant passed SpO₂-based CCHD screening with preductal and postductal SpO₂ of 99 and 97%, respectively. At 27 hours after birth, he was noted to have a grade 3 of 6 systolic murmur heard all over the precordium best heard in the left second intercostal space, with normal first and single second heart sounds. Detailed examination showed normal cardiac apical impulse on the left side, normal peripheral pulses, no bruit, clear breath sounds bilaterally, and no hepatomegaly. His respiratory rates and heart rates were normal at this time (Fig. 1). Echocardiogram was performed due to the murmur and revealed PA with a ventricular septal defect (PA-VSD, Fig. 2). The aorta was overriding the VSD and the flow across the patent ductus arteriosus (PDA diameter of 0.4 cm) was left to right. The infant was started on intravenous prostaglandin E1 (PGE1) therapy at 0.025 µg/kg/min and was transferred to a level-IV neonatal intensive care unit.

During and after the transfer, he continued to have SpO₂ greater than 95% while breathing comfortably in room air (Fig. 1) when the SpO₂ was measured using three different pulse oximeters at the community hospital (Masimo SET 2329 sensor with a GE B450 monitor), during the transfer (Masimo 4003 sensor with a Phillips Intellivue MP 70 monitor) and at the tertiary hospital (Masimo 4003 sensor with a
Zoll X Series monitor). The baby had normal capillary refill, peripheral pulses, and blood pressures. At 38 hours after birth, the infant had respiratory distress that briefly required nasal continuous positive airway pressure of 5 cm H₂O with 21% inspired oxygen that was quickly weaned off within the next few hours to room air. His chest radiograph showed normal lung fields with mild prominence of pulmonary vascularity (►Fig. 3). On the day 2 after birth, the baby underwent cardiac catheterization with the intent to stent the ductus. Unfortunately, the PDA was too large for a stent and hence a Blalock–Taussig (BT) shunt was placed to secure pulmonary flow. The 12-lead electrocardiogram (EKG) at 5 days after birth showed upright T-waves on lead V1 (►Fig. 4) suggestive of right ventricular hypertrophy (RVH) with a strain pattern.

His clinical course was complicated by difficulty in tolerating feeds. He was found to have malrotation for which he underwent Ladd’s procedure with gastrostomy tube placement at 4 weeks and an H-type tracheoesophageal fistula that was surgically ligated at 5 weeks of age. On genetic evaluation, his microarray was normal and his spine radiograph was normal. The renal ultrasound showed grade one hydronephrosis. The infant was suspected to have VACTERL (vertebral anomalies, anal atresia, cardiac defects, tracheoesophageal fistula, renal anomalies and limb anomalies) association. He was discharged home at 8 weeks of age.

Fig. 2 Images from echocardiogram on admission to tertiary hospital. (A) Aorta valve (AoV; red arrow) overriding the interventricular septum (red asterisk); the overriding aortic valve position allows blood from both the right ventricle (RV) and the left ventricle (LV) to flow into the aorta thus creating complete mixing of oxygenated and deoxygenated blood. (B) Color flow Doppler showing a tortuous patent ductus arteriosus (PDA) (red arrow) originating off the underside of the distal aortic arch. (C) Color flow Doppler showing PDA flow (red arrow) to both the right pulmonary artery (RPA) and left pulmonary artery (LPA). (D) Short axis view showing the atretic pulmonary valve (red arrow) adjacent to the aortic valve seen en-face. (E) Four-chamber view (2D and color flow) showing an atrial septal defect. Note the septum bows right to left due to higher pressures in the right atrium (RA) with potential shunting of blood from the RA to the left atrium (LA; red arrow). 2D, two-dimensional.

Fig. 3 Chest and abdomen radiograph of the infant at 2 days after birth notable for absence of pulmonary congestion or cardiomegaly.
During his outpatient cardiology follow-up, an aberrant right subclavian artery and partial anomalous pulmonary venous return (PAPVR) of the right-sided pulmonary veins were detected, and a complete repair of PAPVR along with that of PA-VSD was performed at 5 months of age.

Discussion

The term PA-VSD has been used to describe a group of CHDs with the common features of absence of intraluminal continuity between the pulmonary artery and the right ventricle, and the presence of a nonrestrictive interventricular septal defect. The pulmonary valve is atretic and the pulmonary trunk may also be hypoplastic. The perimembranous VSD (in PA-VSD) is often associated with an overriding aorta and described as "tetralogy type VSD," a type of anterior malalignment defect. This condition could be associated with DiGeorge's syndrome, VACTERL association, and occasionally with tracheomalacia or bronchomalacia. In PA-VSD, the source of pulmonary blood flow (Qp) is from the systemic circulation and is dependent on either major aortopulmonary collateral arteries (MAPCAs) in two-thirds of the cases or on the PDA. MAPCAs represent segmental branches of the descending aorta or indirect branches arising from the aortic arch vessels. When MAPCAs are absent, PGE1 therapy is initiated to maintain patency of the ductus arteriosus followed by placement of a ductal stent or a BT shunt to ensure adequate Qp for oxygenation of blood. Complete repair is performed at a later time.

The absence of direct ventricular connection to the pulmonary arterial system in PA-VSD results in complete mixing of the systemic and pulmonary venous blood with identical oxygen saturations in both the pulmonary arteries and the aorta. In a complete mixing lesion, the systemic oxygen saturation is dependent on the ratio of Qp and systemic (Qs) blood flows. Depending on the mixed venous saturation (50–70%), a Qp:Qs ratio of 1 to 2:1 results in systemic oxygen saturation of 75 to 85%, since the pulmonary venous oxygen saturation is usually 100%. At lower Qp:Qs ratios, a small increase in the ratio results in a large increase in SpO2, whereas at higher Qp:Qs ratios, a similar increase in the ratio would result in a smaller increase in SpO2. The infant discussed here had saturations that were consistently greater than 95%, which would entail Qp:Qs ratio of at least 4–6:1 (theoretically, one aliquot of mixed venous saturation of 70% and four aliquots of pulmonary venous saturation of 100%, Qp:Qs of 4:1 would results in SpO2 = (70 + 100 + 100 + 100 + 100)/5 = 94%). Such high Qp:Qs ratios would be expected to lead to pulmonary congestion presenting with respiratory distress and prominent pulmonary vascularity on chest radiograph. Our patient did not have evidence of respiratory or hemodynamic stress. We speculate that newborn infants can tolerate much higher pulmonary blood flow for several hours prior to becoming symptomatic. This may be due to higher pulmonary vascular resistance in the early postnatal period that dramatically falls at births and continues to fall over the next few days. We consider the theoretical possibility of preferential streaming of oxygenated blood to systemic circulation and deoxygenated blood to pulmonary circulation. The later detection of PAPVR may partially explain the presence of higher SpO2 in our patient due to higher mixed venous oxygen saturation without requiring a higher Qp.

Beginning in 2011, universal screening for CCHDs using pulse oximetry was adopted across the Unites States. It is estimated that the introduction of pulse oximetry screening has improved the detection of ducal dependent CCHDs from 77 to 92%. Ailes et al estimated that approximately 900 infants with nonsyndromic CCHD were likely to be detected by universal CCHD screening every year in the United States; however, an equal number was likely to be missed. The majority of missed cases of CCHD are lesions that compromise systemic blood flow. Every year, there are 550 babies born in the United States with PA which translates to approximately 1 in every 7,100 livebirths. About 60% of the cases of PA with or without VSD are diagnosed prenatally. The majority of prenatally undiagnosed infants with PA have been diagnosed by universal CCHD screening.
Fig. 5 Approach to an asymptomatic term neonate with a cardiac murmur. An easy to follow algorithm that describes an approach to the management of an asymptomatic term neonate with a cardiac murmur. References: Nadas, Brockmeier et al, and Koppel et al. CCHD, critical congenital heart disease; SpO₂, pulse oximetry oxygen saturation.
screening. In the series reported by de-Wahl Granelli et al and that by Zhao et al, none of the cases of PA (n = 3 and 30, respectively) resulted in false negative oximetry screening. Additionally, two infants with PA reported by Siefkes et al failed oximetry screening, and one of them also had low perfusion index values. In contrast, Prudhoe et al mentioned one case of PA who was discharged home undiagnosed but no further details regarding clinical course and time of diagnosis were provided.

The reliability of pulse oximetry in detecting CCHD depends on the accuracy of the measurements. Current recommendations are to use pulse oximeters that perform within a close and effective range as the Food and Drug Administration’s (FDA) trusted standard. There have been reports of discrepancies between SpO2 and SaO2 in newborn infants, with SpO2 being higher than SaO2 by 3%, while discrepancies up to 10% have been noted. This discrepancy may have played a role in our patient passing the CCHD screening. However, the infant had SpO2 of ~95% measured by three different pulse oximeters. Alternatively, a high Qp: Qs ratio would explain the high saturations that we measured in our case.

When oximetry fails, the diagnosis of CCHD depends on detection of abnormal physical findings. It is not uncommon that cardiac murmurs in newborns are often missed or dismissed as benign or innocent. A murmur is considered pathological (and requires further evaluation with a cardiology consultation, EKG, and echocardiogram) when it is a systolic murmur of grade ≥3 of 6 (loud murmur), any diastolic murmur, presence of either signs of congestive heart failure, or cyanosis. Turbulent flow across the PDA or increased flow across the ventricular outflow tracts may result in a murmur detected on physical examination. Increased Qp causes volume load on the heart, leading to respiratory and hemodynamic stress. The systolic murmur heard in our patient was probably secondary to the PDA (where disappearance of the diastolic component of the continuous murmur may be due to higher pulmonary diastolic pressures and lower systemic pressures) or increased flow across the aortic outflow tract. Vital signs reported in our case suggest that excess volume load can be well tolerated early in life, and not easily detected, at least before an infant is eligible for discharge. After the infant is discharged home, lethargy, cyanosis, or respiratory symptoms may alert the parents and the pediatrician; unfortunately, a few may present with sudden death when the ductus arteriosus closes spontaneously. An upright T-wave on lead V1 in an EKG after 48 hours is highly suggestive of right ventricular strain and would help identify right sided obstructive lesions as shown in our patient.

Conclusions

Although pulse oximetry screening prior to newborn discharge has enhanced timely detection of CCHDs, we need to recognize its limitations even in defects that are rarely missed by SpO2 screening. This report reinforces the importance of careful serial physical examinations of infants after birth and prior to their discharge. Pulse oximetry screening is meant to augment the detailed examination of newborns, a sizeable number of whom may have a CCHD.

Authors’ Contributions

D.S., F.R.P., H.S., F.F.I., and S.L. made substantial contributions to conception, design, and drafting of the manuscript. All the authors have critically revised and approved the final version of the manuscript. All authors agree to be accountable for all aspects of the work.

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Conflict of Interest

The authors have no conflict of interest to declare.

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