FACTS
Precious Information
Pulse Oximetry Screening for Critical Congenital Heart Disease

OVERVIEW
Congenital heart defects are malformations of the heart or major blood vessels that occur before birth. In many cases, however, hospital staff may not identify these defects and outwardly healthy infants may be admitted to nurseries and discharged from hospitals before signs of disease are detected.

Occurring in 8 out of 1,000 live births, congenital heart defects account for 27% of infant deaths that are caused by birth defects. A quarter of infants who have congenital heart defects will be diagnosed with critical congenital heart disease (CCHD), a life threatening condition that requires surgery or catheter intervention within the first year of life. Failure to detect CCHD and late detection of CCHD may lead to serious morbidity or death.

Fortunately, an emerging body of evidence suggests that measuring blood oxygen saturation can lead to early diagnosis and detection of CCHD. Once detected, many heart defects can be surgically repaired. It is estimated that 85% of neonates who undergo surgery for CCHD will reach adulthood.

RECOMMENDED UNIFORM SCREENING PANEL FOR NEWBORNS
Newborn screening is a well-established state-based public health program that involves testing all infants for metabolic, hormonal, genetic, and developmental disorders. Each year, more than 98% of newborns are screened across the United States for these disorders.

In 2002, the Health Resources and Services Administration (HRSA) commissioned the American College of Medical Genetics to develop a list of conditions that all states could consider including in their screening programs. This list is called the Recommended Uniform Screening Panel and it currently advises all states to mandate testing for 31 core disorders and 26 secondary disorders. Creation of the Recommended Panel has led to greater uniformity among states in their adoption of screening programs. New conditions for screening are frequently nominated for inclusion in the Panel.

Recently, the U.S. Secretary of Health and Human Services endorsed the addition of CCDH screening to the Recommended Uniform Screening Panel for newborns. The Secretary’s Committee on Heritable Disorders in Newborns and Children recommends that hospitals use a specific type of test called pulse oximetry to screen infants for CCHD.

CUSTOMARY SCREENING PRACTICE
Several tools are regularly used to identify infants who have heart defects.

- Prenatal ultrasounds performed 18-20 weeks into a pregnancy can reveal anatomical abnormalities. Routine prenatal ultrasounds, however, detect less than 50% of CCHD, and rates of detection depend on differing levels of access to prenatal ultrasound and degree of practitioner training.
- After birth, infants are physically examined by primary care providers both before hospital discharge and in routine follow-up visits. Physical exam results may lead clinicians to perform additional tests, including chest radiographs, echocardiograms, and pulse oximetry.

Although prenatal ultrasounds and postnatal physical exams successfully detect many heart defects, they are not sufficient to diagnose all cases of CCHD. New research suggests that when all infants are screened using pulse oximetry in conjunction with the routine practices, CCHD can be detected over in over 90% of newborns.

PULSE OXIMETRY SCREENING
Pulse oximetry screening is a low-cost, non-invasive and painless bedside diagnostic test that can be completed by a technician in as little as 45 seconds. Pulse oximetry testing is conducted to estimate the percentage of hemoglobin in the blood that is saturated with oxygen. When the screening identifies newborns with low blood oxygen concentration, additional testing can be completed to detect heart defects or other life-threatening conditions that could have gone undetected.
FACT SHEET: Pulse Ox Screening for CCHDs

Many studies show that pulse oximetry screening for CCHD has a less than one percent chance of giving false positive results.\(^1\) False positive screening results for CCHD can still offer information to doctors: roughly 25% of infants identified as having low blood oxygen without CCHD may be diagnosed with other conditions that require medical intervention.\(^2\)

The American Heart Association (AHA), the American Academy of Pediatrics (AAP), and the American College of Cardiology Foundation (ACCF) recently outlined recommendations for a standardized pulse oximetry screening approach and diagnostic follow-up.\(^3\) According to these recommendations, screening should be performed on asymptomatic newborns after 24 hours of life in order to avoid false-positive results.\(^4\)

When pulse oximetry screening identifies newborns with low blood oxygen levels, echocardiography can be used to definitively diagnose heart defects.\(^5\) The AHA/AAP/ACCF recommendations emphasize that echocardiograms should be interpreted by pediatric cardiologists.\(^6\) Studies have shown that underserved and rural areas can use telemedicine to access pediatric cardiologists for CCHD diagnosis.\(^7\)\(^8\)

Pulse oximeters are available in most neonatal units, and hospital staff are well trained in how to perform pulse oximetry screening.\(^9\)\(^10\) A recent cost-effectiveness analysis estimated that universal newborn pulse oximetry screening would cost just under $4 per infant.\(^11\) Although there are monetary costs associated with false positive results from pulse oximetry screening, these costs may be partially or fully offset by early diagnosis of infants with CCHD before they become ill and/or incur irreversible damage. Research suggests that the cost savings associated with early detection of a single case of CCHD could exceed the costs associated with screening 2,000 infants.\(^12\) Many clinicians and experts agree that the benefits of detecting CCHD far outweigh the costs incurred by the screening itself.\(^13\)

Although there is not a clear way to bill insurers for pulse oximetry screening at this time, many other routine newborn tests, including hearing screenings, are frequently included in the bundle of services that hospitals provide to infants prior to discharge.\(^14\)

**STATE POLICY APPROACHES TO PULSE OXIMETRY SCREENING**

States across the nation are beginning to work to implement the Secretary's recommendation to screen all newborns for CCHD.

State policies have a substantial effect on newborn screening rates. Research shows that screening rates are significantly higher in states that have passed test-specific legislation than in states without these laws.\(^15\) While some individual providers or hospital systems may initiate voluntary pulse oximetry screening, legislative action is the only way to ensure equitable and uniform CCHD screening for all newborns.

**THE AHA ADVOCATES**

The AHA is committed to advancing public policies that will allow children and adults with heart defects to live longer and fuller lives. These policies include:

- State adoption of mandatory CCHD screening using pulse oximetry for all newborns;
- The collection of screening data to be used for surveillance, evaluation and continuous quality improvement of CCHD screening;\(^16\)
- The development, dissemination, and validation of screening standards for CCHD;
- The continued development of FDA’s guidance document regarding the safety and effectiveness of pulse oximeters.\(^17\)

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**References**

12. Secretary’s Advisory Committee on Heritable Disorders in Newborns and Children. Recommended Uniform Screening Panel. Health Resources and Services Administration; 2011–1–2.