

Important Information for Parents of Infants Who Have Not Yet Had the Newborn Screening Test



Newborn Screening Branch
Genetic Disease Screening Program
www.dhs.ca.gov/nbs



California Department of Public Health

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*For more information about the Newborn Screening Program
and the most current list of diseases that can be detected
through the Program, visit our website at*
www.dhs.ca.gov/nbs

The California Newborn Screening Test

Newborn screening can save your baby's life or prevent serious brain damage. Newborn screening can identify babies with certain diseases so that treatment can be started right away. Early identification and treatment can prevent mental retardation and/or life-threatening illness.



Does My Baby Need To Have A Newborn Screening Test?

Yes, to protect the health of all its newborns, California state law requires that your baby must have the Newborn Screening (NBS) Test before leaving the hospital. This test screens for specific diseases in the following groups:

■ Metabolic ■

chemical reactions in the body to create energy and build tissue

■ Endocrine ■

hormones that affect body functions

■ Hemoglobin ■

red blood cells that carry oxygen

■ Other Genetic Diseases ■

Cystic Fibrosis

In California there will be about 750 babies identified with one of these diseases each year. This means about 1 out of every 760 babies tested will have one of these diseases.

What Is Screening?

Screening is the testing of a group of people to identify those who are at risk for having a specific disease even though they may seem healthy. Newborn screening identifies most, but not all, of the babies who have one or more of the many diseases screened for by the California Program. Not every baby with a positive screening test will have one of these diseases. Further testing and evaluation by the baby's health care provider or a specialist are needed to make the diagnosis.



Is My Baby Too Old For Newborn Screening?

No. If your baby is less than a year old, he or she can still be tested. However, since babies can look healthy and still have a disorder, it is critical to have your baby tested as soon as possible. The earlier babies with any of these conditions receive treatment, the better they will do.

How Is The Test Done?

A few drops of blood taken from the baby's heel are put on special filter paper. The blood is then sent to a state-approved lab for testing.



Is The Test Safe?

Yes, this is a simple and safe test. Over 13 million California newborns have had blood collected by heelstick without any harm to the newborn.

Can I Say No To The Test?

You can only say no if screening is in conflict with your religious beliefs or practices. You must then sign a special form. It states that not having the test done can result in serious illness or permanent damage to your child. It also states that you accept responsibility should this occur.



Is The Test Accurate?

Yes. The blood is sent to a state-approved lab for testing. The state checks the work of the testing labs closely to make sure the results are reliable. It is rare that a baby with one of the more common diseases is not found through a positive newborn screening test. For a few rare diseases, the test may find only a small number of the babies affected.

Where Can I Have The Screening Test Done?

Many hospitals and health department clinics provide this service. Call your health care provider or local health department to find out where to go for testing.



How Can I Get The Results?

If the test is positive, you will be contacted within a few days after having the test done. If the test is negative, it takes about two weeks for doctors to get a copy of the results. You can get your baby's test results from your doctor or clinic. If your doctor does not have the results, he/she can contact the Newborn Screening Program to request a copy.

If you move after the test is done, make sure your baby's doctor, midwife or clinic has your new address and phone number in case they need to contact you about your baby's results.

What Do I Do If The Baby's Results Are Positive?

If the results are positive, more tests will be needed. You should receive a phone call and/or letter about what to do next. After further testing, many babies who have a positive first test are found **not** to have a disease. However, you must have your baby re-tested because babies who do have one of these diseases benefit from early treatment.

Early Treatment Can Prevent Serious Problems

If these diseases are not found and treated soon after birth, the baby can have serious health problems or even die. Early treatment can prevent many of these problems.

Metabolic Diseases

Metabolic diseases affect the body's ability to use certain parts of food for growth, energy, and repair. The parts include **amino acids** from proteins, **fatty acids** from fats and **organic acids** from proteins, fats, and sugars. To break down or convert these substances, certain proteins called enzymes must be present. When there aren't enough of the needed enzymes, some substances build up in large amounts and may be harmful to the body. Metabolic diseases have varying degrees of severity. If identified early, many of these conditions can be treated before they cause serious health problems. Treatment may include close monitoring of the person's health, medication, dietary supplements, and/or special diets.



These are some of the metabolic diseases screened for by the NBS Program:

- **Phenylketonuria (PKU)** – Babies with PKU have problems when they eat foods high in protein such as milk (including breast milk and formula), meat, eggs, and cheese. Without treatment, babies with PKU develop mental retardation and/or have other health problems. A special diet can prevent these problems.
- **Galactosemia** – Babies with this disease cannot use some of the sugars in milk, formula and breast milk, and other foods. Without treatment, babies with galactosemia can become very sick and die. A special infant formula and diet can help prevent these problems.
- **Maple Syrup Urine Disease (MSUD)** – Babies with MSUD have problems using fats and protein. Without treatment, MSUD can cause mental retardation or death. Treatment with a special diet can prevent these problems.
- **Medium-Chain Acyl-CoA Dehydrogenase Deficiency (MCADD)** – Babies with MCADD are unable to convert fat into energy. Without treatment, babies can have seizures, extreme sleepiness, coma, and even die. Treatment may include frequent feedings and a special diet.
- **Homocystinuria** – Babies with this disease have problems breaking down protein. Without treatment, babies can have delayed development, eye problems, and/or other health problems. A special diet can help prevent these problems.
- **Biotinidase Deficiency** – Babies with this disease cannot use or recycle biotin (vitamin B) from their diet. Without treatment, babies with this disorder can have seizures, mental retardation, vision problems, hearing loss, and/or other health problems. Treatment includes daily biotin supplements.

Endocrine Diseases

Babies with endocrine diseases make too much or too little of certain hormones. Hormones are produced by glands in the body and affect body functions.

- **Congenital Adrenal Hyperplasia (CAH)** – The adrenal glands of babies with this disease do not make enough of the key stress-fighting hormone cortisol. In about two-thirds of the cases, babies also do not produce enough of the salt-retaining hormone aldosterone. As a result, affected babies can develop dehydration, shock, and even death. Treatment with one or more oral medicines can help prevent these problems. Girls with this condition may have the additional problem of having masculine-looking external genitals, which can be corrected with surgery.
- **Primary Congenital Hypothyroidism** – Babies born with this disease do not have enough thyroid hormone. Without enough hormone, babies grow very slowly and have mental retardation. These problems can be prevented by giving the baby special thyroid medicine every day.

Hemoglobin Diseases

Hemoglobin is found in red blood cells. It gives blood its red color and carries oxygen to all parts of the body. Hemoglobin diseases often lead to anemia because they affect the type and amount of hemoglobin in the red blood cells. Treatment may include medication, folic acid and close monitoring of the child's health.

These are some of the hemoglobin diseases that are part of the newborn screening test:

- **Sickle Cell Anemia and other Sickle Cell Diseases** –

These diseases affect the type of hemoglobin in the baby's red blood cells. Babies with sickle cell disease can get very sick and even die from common infections. Many of the infections can be prevented with daily antibiotics. Ongoing health care and close monitoring help children with hemoglobin diseases stay as healthy as possible.

- **Hemoglobin H Disease** – This disease affects the amount of hemoglobin in the baby's blood. There is less hemoglobin, which results in smaller red blood cells. This also causes the cells to break down faster than normal. Babies with this disease can have mild to severe anemia, as well as other health problems. Treatment can include blood transfusions, taking folic acid, and avoiding certain medications and household products.

Cystic Fibrosis

This disease can affect many body organs, including the lungs and digestive system. In the first few months of life, a baby with CF can have poor absorption of milk or formula, slow growth, failure to thrive, recurrent lung infections, salty sweat, frequent runny stools, dehydration and life-threatening salt imbalance. Early treatment along with ongoing health care by a team at a special care center can alleviate many of these problems.

Will The Treatment Still Help If The Damage Has Already Occurred?

Yes, even delayed treatment can be helpful. It can prevent further damage. It could even save the life of your baby if he or she has one of these disorders. Have your baby tested as soon as possible. In addition, your baby will need regular well-baby health care. If you have any questions about your baby's health call your baby's health care provider or local health department.

Is Information About My Baby's Test Confidential?

Yes. There are serious penalties for any unauthorized release of private information collected during screening. For a copy of our Notice of Information and Privacy Practices please see pages 13-15 of this booklet.

How Much Does The Test Cost?

The cost is subject to change. Please check with your doctor, the hospital, or the NBS website for the current cost of the test. Medi-Cal, health plans and most private insurance will pay for the test. The cost is included in the hospital bill. You will not receive a bill from the NBS Program. If you have problems with your insurance, contact 1-800-927-HELP (1-800-927-4357) or if you have a prepaid health plan, contact 1-888-HMO-2219 (1-888-466-2219).

California law prevents insurance companies from refusing to issue or canceling a policy, or charging a higher rate or premium based on a person's genetic characteristics, including being diagnosed with one of the diseases found by newborn

screening. If you have any of these problems, call one of the numbers listed above. It is also illegal to refuse employment based on the results of a genetic test.

Does My Baby Need Any Other Blood Tests?

If there is something that you are worried about, or know of a disease that may run in the family, talk to your doctor about what other tests can be done. The Newborn Screening Program screens for the most common treatable diseases and includes almost all of the diseases a commercial newborn screening test would. The Program evaluates adding other diseases as new tests and treatments become available. However, newborn screening does not test for every disease that might be found.

Due to biological variability of newborns and differences in detection rates for the various disorders in the newborn period, the Newborn Screening Program will not identify all newborns with these conditions. A negative screening result does not rule out the possibility of a disorder. Parents should remain watchful for any signs or symptoms of these disorders in their child and consult a physician. In addition to screening, babies also need regular well-baby check-ups.



NOTICE OF INFORMATION AND PRIVACY PRACTICES

California Department of Public Health (CDPH) Genetic Disease Screening Program, Newborn Screening Program (Effective Date April 14, 2003)

This notice describes how personal and medical information about you or your newborn may be used and disclosed and how you can get access to this information. Please review it carefully.

Department's Legal Duty. Federal and State laws restrict the use, maintenance and disclosure of personal and medical information obtained by a State agency, and require certain notices to individuals whose information is maintained. In compliance with these laws, you and those providing information are notified of the following:

Department Authority and Purpose for the Newborn Screening Program. The CDPH collects information related to newborn screening as permitted in Health and Safety Code Sections 124980, 125000, 125001, 125025, and 125030. This information is collected electronically and includes such things as your name, address, medical care given to you and your newborn. Testing is required by law (Health and Safety Code Section 125000) and regulations (17 CCR 6500 through 6510) and if the required information is not provided, death or permanent handicaps for affected newborns could result. If you have religious objections to testing, you may say "no" to testing in writing and sign a form advising you that your hospital, doctor and clinic staff are not responsible if your baby develops problems because those disorders were not identified and treated early.

Uses and Disclosure of Health Information. The CDPH uses health information about you or your newborn for screening, to provide health care services, to obtain payment for screening, for administrative purposes, and to evaluate the quality of care that you or your newborn receives. Some of this information is retained for as long as 21 years. The information will not be sold.

The law also allows the Department to use or give out information we have about you or your newborn for the following reasons:

- For research studies unless you specifically request in writing that your information not be used.
- To organizations, which help us in our operations, such as collecting fees.

The information is otherwise confidential and will not be released without your written authorization. If you sign an authorization to disclose information you can later revoke that authorization to stop any future uses and disclosures by contacting the person listed below.

The Department may change its policies at any time subject to applicable laws and regulations. You may request a copy of our current policies or obtain more information about our privacy practices by contacting the person listed below or consulting our website at www.dhs.ca.gov/nbs. You may also request a paper copy of this notice.

Individual Rights and Access to Information. You have the right to look at or receive a copy (you will be charged) of your or your newborn's health information and receive a list of instances where we have disclosed health information about you or your newborn for reasons other than screening, payment or related administrative purposes. If you believe that information in your or your newborn's record is incorrect or missing, you have the right to request corrections. You have the right to make reasonable requests for us to contact you only in writing or at a different address, post office box, or telephone number.

You may request in writing that we restrict disclosure of your or your newborn's information for health care treatment, payment and administrative purposes. We are not required to agree to your request.

Complaints. If you believe that we have not protected your or your newborn's privacy or have violated any of your or your

newborn's rights you may file a complaint by calling or writing:
Privacy Officer, CDPH, P.O. Box 997413, Sacramento, CA
95899-7413, 916-445-4646 or (877) 735-2929 TTY/TDD. Or
visit our website at: www.dhs.ca.gov/privacyoffice/NPP/default.htm

You may also contact the Secretary of the Department of Health and Human Services, Office for Civil Rights at 50 United Nations Plaza, Room 322, San Francisco, CA. 94102, telephone (800) 368-1019. Or you may call the U.S. Office for Civil Rights at 866-OCR-PRIV (866-627-7748) or 866-788-4989 TTY.

The Department cannot take away your health care benefits or do anything to hurt you in any way if you choose to file a complaint or use any of the privacy rights in this notice.

Department Contact - The information on this form is maintained by the CDPH, Genetic Disease Screening Program. Address correspondence to the Chief of the Genetic Disease Screening Program, 850 Marina Bay Parkway, F175, Mail Stop 8200, Richmond, California, 94804 (510-412-1502).

Americans with Disabilities Act (ADA)
Notice of Information and Access Statement
Policy of Nondiscrimination on the Basis of Disability
And Equal Employment Opportunity Statement

The California Department of Public Health (CDPH) complies with all state and federal laws, which prohibit discrimination in employment and provide admission and access to its programs or activities.

The Deputy Director, Office of Civil Rights (OCR), CDPH has been designated to coordinate and carry out the department's compliance with nondiscrimination requirements. Title II of the ADA addresses non-discrimination and access issues regarding disabilities. To obtain information concerning the CDPH EEO Policies or the provision of the ADA and the rights provided, you may contact the CDPH OCR by phone at 916-440-7370, TTY 916-440-7399 or write to:

OCR, CA Department of Public Health
P.O. Box 997413, MS 0009
Sacramento, CA 95899-7413

Upon request, this document will be made available in Braille, high contrast, large print, audiocassette or electronic format. To obtain a copy in one of these alternate formats, call or write:

Chief, Newborn Screening Branch
850 Marina Bay Pkwy., F175, MS 8200
Richmond, CA 94804
Phone: 510-412-1502
Relay Operator: 711/1-800-735-2929

***The Genetic Disease
Screening Program wants
to provide quality services to
the families of California and
welcomes your***

Department of Public Health
Newborn Screening Program
850 Marina Bay Parkway, MS 8200
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