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Noonan Syndrome

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Abstract

Noonan syndrome is a common genetic disorder that causes multiple congenital abnormalities and a large number of potential health conditions. Most affected individuals have characteristic facial features that evolve with age; a broad, webbed neck; increased bleeding tendency; and a high incidence of congenital heart disease, failure to thrive, short stature, feeding difficulties, sternal deformity, renal malformation, pubertal delay, cryptorchidism, developmental or behavioral problems, vision problems, hearing loss, and lymphedema. Familial recurrence is consistent with an autosomal dominant mode of inheritance, but most cases are due to de novo mutations. Diagnosis can be made on the basis of clinical features, but may be missed in mildly affected patients. Molecular genetic testing can confirm diagnosis in 70% of cases and has important implications for genetic counseling and management. Most patients with Noonan syndrome are intellectually normal as adults, but some may require multidisciplinary evaluation and regular follow-up care. Age-based Noonan syndrome-specific growth charts and treatment guidelines are available.

Noonan syndrome is a common genetic disorder with multiple congenital abnormalities. It is characterized by congenital heart disease, short stature, a broad and webbed neck, sternal deformity, variable degree of developmental delay, cryptorchidism, increased bleeding tendency, and characteristic facial features that evolve with age. Molecular genetic testing can confirm the diagnosis in most cases, which has important implications for genetic counseling and management. Physicians should know how to diagnose Noonan syndrome because patients who have it require monitoring for a large number of potential health conditions. Age-appropriate guidelines for the management of Noonan syndrome are available.¹

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Epidemiology

Noonan syndrome is characterized by marked variable expressivity, which makes it difficult to identify mildly affected individuals. The incidence is one in 1,000 to 2,500 live births for severe phenotype, but mild cases may be as common as one in 100 live births.² Familial recurrence is consistent with an autosomal dominant mode of inheritance, but de novo mutations are more common, accounting for 60% of cases.³ There is no known predilection by race or sex.

Clinical Presentation

The diagnosis of Noonan syndrome depends primarily on the identification of characteristic clinical features (Table 1¹⁻¹¹). The clinical spectrum in children is well described, but few studies have examined medical complications in adults.³ Figures 1 through 4 show the cardinal phenotypic features of Noonan syndrome based on patient age.⁶⁻⁸

Nonspecific prenatal anomalies common among patients with Noonan syndrome include increased nuchal translucency, polyhydramnios, and abnormal maternal serum triple screen (α -fetoprotein, human chorionic gonadotropin, and unconjugated estriol). The most common morphologic fetal anomaly is hydrothorax. Diagnosis of cardiac anomalies is rarely made prenatally. The diagnosis of Noonan syndrome should be considered in all fetuses with a normal karyotype and increased nuchal translucency, especially when cardiac anomaly, polyhydramnios, and/or multiple effusions are observed.¹²

Patients who have Noonan syndrome display clinical similarities to patients who have Turner syndrome (EDITOR'S NOTE: see <http://www.aafp.org/afp/2007/0801/p405.html>). However, patients can be easily differentiated because with Turner syndrome, only females are affected (because of the loss of the X chromosome), left-side heart defects are common, developmental delay occurs less often, renal anomalies are more common, and primary hypogonadism causes amenorrhea and sterility.^{8,13} A number of other partially overlapping syndromes, such as cardio-facio-cutaneous syndrome, Watson syndrome, Costello syndrome, neurofibromatosis 1, and LEOPARD syndrome (lentiginos, electrocardiogram conduction abnormalities, ocular hypertelorism, pulmonary stenosis, abnormal genitalia, retardation of growth, and sensorineural deafness), also can be differentiated from Noonan syndrome based on clinical features.^{8,14}

Diagnosis

Early, accurate diagnosis of Noonan syndrome is important because each patient requires an individual treatment regimen and has a different prognosis and recurrence risk. A scoring system has been devised to help diagnose patients with the condition (Table 2¹⁵).

Until recently, diagnosis was made solely on the basis of clinical features, but molecular genetic testing can provide confirmation in 70% of cases.⁸ Noonan syndrome is caused by mutations in the RAS/mitogen-activated protein kinase (MAPK) pathway, which is essential for cell cycle differentiation, growth, and senescence.¹⁴ Approximately one-half of the

known mutations are in the protein tyrosine phosphatase nonreceptor, type 11 (PTPN 11) gene.⁸

Genetic Counseling

Most cases are sporadic. In familial cases, autosomal dominant inheritance is confirmed. The risk of Noonan syndrome developing in the sibling of an affected person is 50% if the parent is affected, but is less than 1% if the parent is unaffected. Risk of transmission to the offspring of an affected individual is 50%.⁸ Preimplantation genetic diagnosis can be offered in familial cases with known mutations.⁸

Management

Intellectual and physical abilities are normal in most adults with Noonan syndrome, but some may require multidisciplinary evaluation and regular follow-up care.¹⁶ Recently, management guidelines were developed by American and European consortia, and management is optimized by adherence to age-specific guidelines that emphasize screening and testing for common health issues.^{1,17} Referral to a clinical geneticist for assistance in the diagnosis and management of Noonan syndrome, including determining the appropriateness and sequence of genetic testing, may be helpful.^{1,8} Clinical growth charts are also available via a European network at <http://www.dyscerne.org>. Table 3 lists system-based management guidelines to assist physicians caring for patients with Noonan syndrome and their families.^{1,8,17} Table 4 describes when Noonan syndrome should be suspected.

Resources

The following education, support, referral, and research websites are useful for physicians and for their patients who have Noonan syndrome:

- GeneTests (www.genetests.org)
- Genetics Home Reference (<http://ghr.nlm.nih.gov/condition/noonan-syndrome>)

Genomics Glossary

Cardio-facio-cutaneous syndrome: Characterized by cardiac abnormalities, distinctive craniofacial appearance, and cutaneous abnormalities.

Costello syndrome: Characterized by growth problems, developmental delay or intellectual disability, coarse facial features, curly or sparse fine hair, soft skin with deep palmar and plantar creases, papillomata of the face and perianal region, diffuse hypotonia, joint laxity, and cardiac disease.

LEOPARD syndrome: An acronym for the cardinal features of lentiginos, electrocardiogram conduction abnormalities, ocular hypertelorism, pulmonary stenosis, abnormal genitalia, retardation of growth, and sensorineural deafness.

Neurofibromatosis 1: Characterized by multiple café au lait spots, axillary and inguinal freckling, neurofibromas, Lisch nodules, gliomas, and osseous lesions.

Preimplantation genetic diagnosis: A procedure used to decrease the chance of a particular genetic condition for which the fetus is specifically at risk by testing one cell removed from early embryos conceived by in vitro fertilization and transferring to the mother's uterus only those embryos determined not to have inherited the mutation in question.

RAS/mitogen-activated protein kinase (MAPK) pathway: Essential in the regulation of the cell cycle, differentiation, growth, and cell senescence, all of which are critical to normal development. Mutations in the genes that encode for the pathway have profound effects on development.

Turner syndrome: A chromosomal abnormality in which one X chromosome is absent; 45,X karyotype.

Variable expressivity: The range of signs and symptoms that can occur in different persons with the same genetic condition.

Watson syndrome: An autosomal dominant disorder characterized by pulmonary stenosis, café au lait spots, decreased intellectual ability, and short stature. Most affected individuals have a large head, Lisch nodules, and neurofibromas.

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This article is one in a series coordinated by the National Human Genome Research Institute, National Institutes of Health, Bethesda, Md. A glossary of genomics terms is available at <http://www.aafp.org/afp/genglossary>.

This clinical content conforms to AAFP criteria for continuing medical education (CME).
See CME Quiz Questions on page 6.

Patient information: A handout on this topic, written by the authors of this article, is available at <http://www.aafp.org/afp/2014/0101/p37-s1.html>. Access to this handout is free and unrestricted.

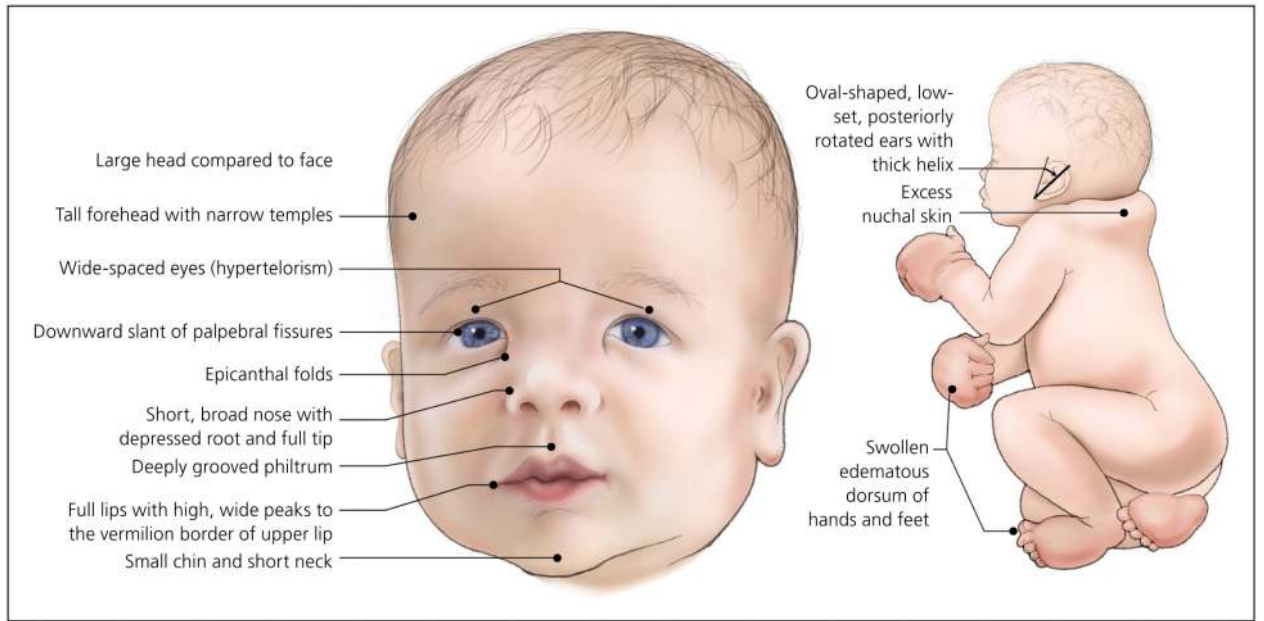


Figure 1.
Newborn with Noonan syndrome.

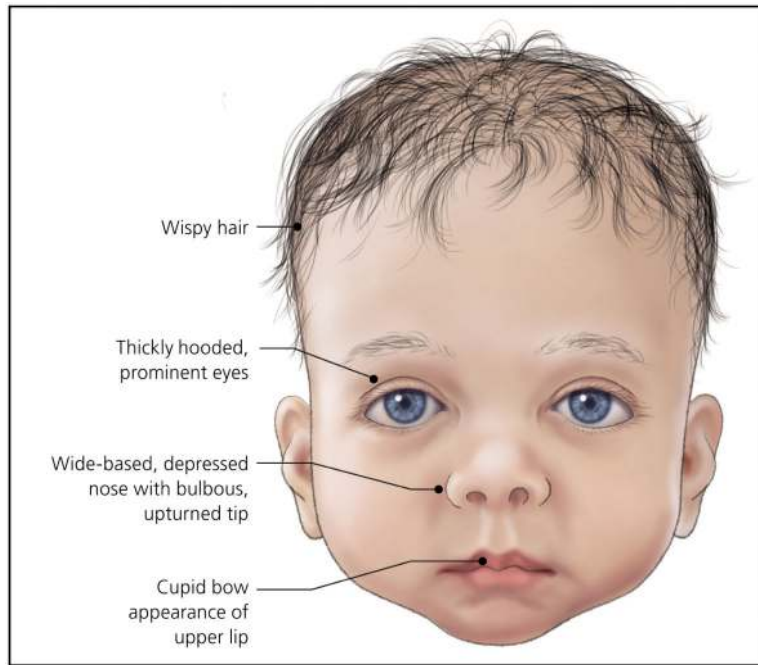


Figure 2.
Infant with Noonan syndrome.

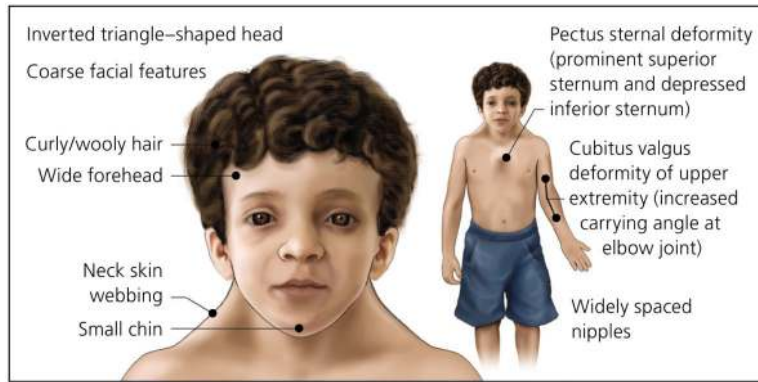


Figure 3.
Child/adolescent with Noonan syndrome.

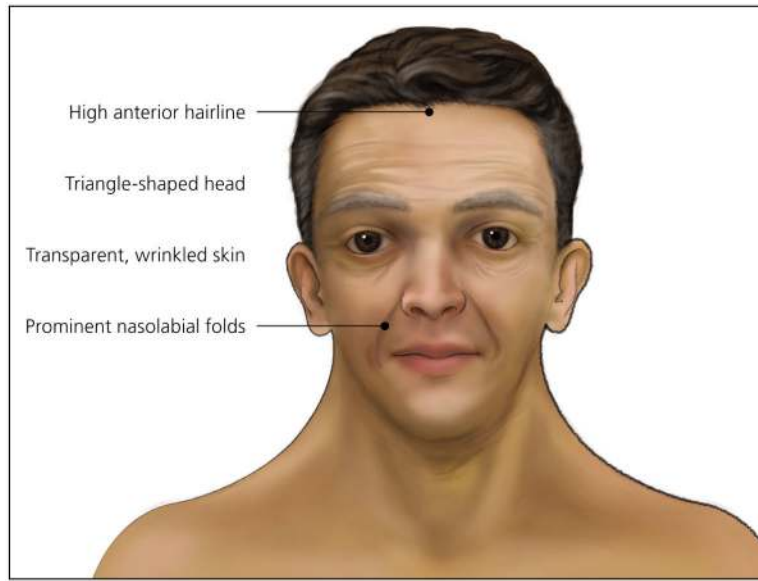


Figure 4.
Adult with Noonan syndrome.

Table 1

Clinical Features of Noonan Syndrome

<p>Cardiovascular^{4,5} Hypertrophic cardiomyopathy Pulmonary stenosis, often with a dysplastic valve</p> <p>Dental/oral^{1,2} Articulation difficulty High arched palate Malocclusion Micrognathia</p> <p>Dysmorphic facial features^{6,7} See Figures 1 through 4</p> <p>Ears⁸ Hearing loss</p> <p>Eyes^{2,9} Anterior segment problems (prominent corneal nerves, cataract, anterior stromal dystrophy) Nystagmus Ptosis, hypertelorism, and epicanthal folds Refractive error Strabismus</p> <p>Gastrointestinal^{3,4} Feeding difficulties (poor sucking function, prolonged feeding time, recurrent vomiting and reflux)</p>	<p>Genitourinary^{2,8} Cryptorchidism Female fertility is normal Males can have fertility issues (e.g., defective spermatogenesis caused by cryptorchidism, gonadal dysfunction due to impaired Sertoli cell function) Malformations (renal pelvis dilation, solitary kidney, duplex collecting system) Puberty can be delayed in both sexes</p> <p>Growth^{2,4,8} Birth weight and length are normal Failure to thrive and short stature (50% to 70% of patients with Noonan syndrome) Mean final adult height is 63 to 66 inches (160 to 168 cm) in males and 59 to 61 inches (150 to 155 cm) in females</p> <p>Hematologic^{1,3,4,10,11} Increased bleeding tendency (due to factor deficiency, quantitative or qualitative platelet defect) Leukemia Myeloproliferative disorder</p> <p>Lymphatic⁸ Lymphedema</p>	<p>Neurologic^{1,8} Behavioral conditions (stubbornness, irritability, body image problems, poor self-esteem) Central nervous system malformation Early motor milestones delay (hypotonia and joint laxity) Learning difficulties Mild intellectual disability (33% of patients with Noonan syndrome) Most individuals have normal intelligence Speech disorders</p> <p>Skeletal^{2,4} Cubitus valgus Spinal abnormality (scoliosis, talipes equinovarus) Sternal deformities (pectus carinatum superiorly, pectus excavatum inferiorly)</p> <p>Skin conditions^{2,4} Dystrophic nails Extra prominence on pads of fingers and toes Follicular keratosis Hyperelastic skin Moles Multiple lentiginos Nevi Thick curly hair or thin sparse hair</p>
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Information from references 1 through 11.

Table 2

Diagnostic Criteria for Noonan Syndrome

Feature	A = Major	B = Minor
1. Facial	Typical facial dysmorphism (facial features vary with age and are described in Figures 1 through 4)	Suggestive facial dysmorphism
2. Cardiac	Pulmonary valve stenosis, hypertrophic cardiomyopathy, and/or electrocardiographic results typical of Noonan syndrome	Other defect
3. Height	< 3rd percentile	< 10th percentile
4. Chest wall	Pectus carinatum/excavatum	Broad thorax
5. Family history	First-degree relative with definite Noonan syndrome	First-degree relative with suggestive Noonan syndrome
6. Other features	All of the following: intellectual disability, cryptorchidism, and lymphatic vessel dysplasia	One of the following: intellectual disability, cryptorchidism, or lymphatic vessel dysplasia

NOTE: Noonan syndrome is considered present if the patient has typical facial dysmorphism plus one feature from categories 2A through 6A or two categories from features 2B through 6B, or has suggestive facial dysmorphism plus two features from categories 2A through 6A or three features from categories 2B through 6B.

Adapted with permission from van der Burgt I, Berends E, Lommen E, van Beersum S, Hamel B, Mariman E. Clinical and molecular studies in a large Dutch family with Noonan syndrome. *Am J Med Genet.* 1994;53(2):190.

Table 3

Guidelines for Management of Noonan Syndrome

System-based intervention	Clinical issue	Timing of intervention	Comments
Auditory	Hearing loss	Hearing tests in infancy and/or at diagnosis Annual hearing test throughout childhood	—
Cardiovascular	Congenital heart defects	At the time of diagnosis and regular follow-up with cardiologist Children and adults without heart disease on initial evaluation should have cardiac reevaluation every five years	Cardiac consultation, echocardiography, and electrocardiography
Dental	Dental malocclusion	Oral examination by primary care physician at each visit Dental referral between one and two years of age, with annual dentist visits thereafter	—
Dermatologic	See Table 1 for common skin findings	Referral as indicated	—
Developmental	Behavioral disorders Developmental delay Learning difficulties Mild intellectual disability Speech disorders	Development assessment annually (beginning in second half of first year of the patient's life or at diagnosis) Baseline neuropsychologic assessment at primary school entry	If screening results are abnormal, consider appropriate intervention
Endocrine	Delayed puberty Hypothyroidism Short stature	Referral as indicated	Endocrine evaluation for growth hormone therapy to treat short stature and hormone therapy for delayed puberty Thyroid function tests if patient shows signs and symptoms of hypothyroidism
Gastroenterologic	Feeding difficulties	Referral as indicated	Appropriate therapeutic intervention as needed (e.g., swallowing study, speech therapy, reflux studies)
Genetic	To confirm diagnosis, genetic counseling and genotype–phenotype correlation	Referral as indicated	—
Growth	Failure to thrive Short stature Slow growth	At the time of diagnosis, three times a year for the first three years, then annually	Monitor and plot growth on Noonan syndrome age-based growth charts
Hematologic	Increased bleeding tendencies (factor deficiency, quantitative or qualitative platelet defect)	Baseline coagulation screen at diagnosis or after six to 12 months of age if screening was initially performed during infancy or whenever diagnosis is made	If bleeding symptoms: First-tier: Baseline coagulation screen (complete blood count, prothrombin time, activated partial thromboplastin time) Second-tier (in consultation with hematologist): Specific factor assay and platelet function studies
Lymphatic	Lymphatic vessel dysplasia, hypoplasia, or aplasia	Referral as indicated	Refer patients with lymphedema to lymphedema clinic (contact National Lymphedema Network; www.lymphnet.org)
Metabolic	Failure to thrive Inadequate weight gain	Referral as indicated	Dietary assessment and nutrition intervention
Neurologic	Arnold-Chiari malformation Craniosynostosis	Referral as indicated	Physician should have a low threshold for investigating neurologic symptoms (electroencephalography, magnetic

System-based intervention	Clinical issue	Timing of intervention	Comments
	Headaches		resonance imaging of the brain)
	Hydrocephalus		
	Seizures		
Ocular	Dysmorphic findings Vision problems	Detailed eye examination in infancy and/or at diagnosis Eye reevaluation as indicated if abnormal or every two years thereafter	—
Pregnancy	Congenital heart defects Effusion Hydrops fetalis Increased nuchal translucency Polyhydramnios Renal anomalies	Per obstetrician's recommendation	Chorionic villus sampling or amniocentesis for diagnosis if indicated Fetal ultrasonography and echocardiography
Renal	Renal anomalies (e.g., pyeloureteral stenosis, hydronephrosis)	Renal ultrasonography at the time of diagnosis	Increased risk of urinary tract infection if structural anomaly found, consider antibiotic prophylaxis and evaluation by nephrologist
Reproductive	Female fertility normal ⁸ Male fertility problems related to defective spermatogenesis caused by cryptorchidism or gonadal dysfunction due to impaired Sertoli cell function	Orchiopexy if testes undescended by one year of age	Fertility clinic evaluation in males
Skeletal	Pectus deformity of sternum Spinal abnormality	Annual examination of chest and back	If screening abnormal, consider radiography of the spine
Surgical and anesthesia risk	Increased risk of complications due to a cardiac disorder, increased bleeding tendency, or craniofacial and/or vertebral anomalies	Referral as indicated	See cardiovascular, hematology, and skeletal management recommendations

Information from references 1, 8, and 17.

Table 4

When to Suspect Noonan Syndrome

Noonan syndrome should be considered in anyone who presents with two or more of the following:

Characteristic facial features (Figures 1 through 4)	Short stature
Developmental delay and/or learning disability	Typical chest deformity
Heart defect	Undescended testes
Pubertal delay and/or infertility	First-degree relative who has Noonan syndrome or any of the above features