Cardiofaciocutaneous Syndrome and the Dermatologist’s Contribution to Diagnosis

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PRACTICE POINTS
- RASopathies, a class of developmental disorders, are caused by mutations in genes that encode protein components of the RAS/mitogen-activated protein kinase pathway. Cardiofaciocutaneous (CFC) syndrome is a RASopathy.
- Skin manifestations may help in differentiating CFC syndrome from other RASopathies.

To the Editor:
RASopathies, a class of developmental disorders, are caused by mutations in genes that encode protein components of the RAS/mitogen-activated protein kinase (MAPK) pathway. Each syndrome exhibits its phenotypic features; however, because all of them cause dysregulation of the RAS/MAPK pathway, there are numerous overlapping phenotypic features between the syndromes including cardiac defects, cutaneous abnormalities, characteristic facial features, neurocognitive impairment, and increased risk for developing some neoplastic disorders.

Cardiofaciocutaneous (CFC) syndrome is a RASopathy and is a genetic sporadic disease characterized by multiple congenital anomalies associated with mental retardation. It has a complex dermatological phenotype with many cutaneous features that can be helpful to differentiate CFC syndrome from Noonan and Costello syndromes, which also are classified as RASopathies.

A 3-year-old girl presented with skin xerosis and follicular hyperkeratosis of the face, neck, trunk, and limbs (Figure 1). Facial follicular hyperkeratotic papules on an erythematous base were associated with alopecia of the eyebrows (ulerythema ophryogenes). Hair was sparse and curly (Figure 2A). Facial dysmorphic features included a prominent forehead with bitemporal constriction, bilateral ptosis, a broad nasal base, lip contour in a Cupid’s bow, low-set earlobes with creases (Figure 2B), and a short and webbed neck.

Congenital heart disease, hypothyroidism, bilateral hydronephrosis, delayed motor development, and seizures were noted for the first 2 years. Brain computed tomography detected a dilated ventricular system with hydrocephalus. There was no family history of consanguinity.

Pregnancy was complicated by polyhydramnios and preeclampsia. The neonate was delivered at full-term and was readmitted at 6 days of age due to respiratory failure secondary to congenital chylothorax. Cardiac malformation was diagnosed as the ostium secundum atrial septal defect and interventricular and atrioventricular septal defects. Up to this point she was being treated for Turner syndrome.

The RASopathies are a class of human genetic syndromes that are caused by germ line mutations in genes that encode components of the RAS/MAPK pathway.1 There are many syndromes classified as RASopathies (Table).2,3

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Figure 1. Follicular hyperkeratosis of the limbs (A and B) and trunk (C).

Figure 2. Sparse and curly hair (A) and low-set earlobes with creases (B).

### RASopathies and Involved Genes

<table>
<thead>
<tr>
<th>Syndrome</th>
<th>Gene(s)</th>
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<tbody>
<tr>
<td>Autoimmune lymphoproliferative syndrome type IV (RAS-associated autoimmune leukoproliferative disorder)</td>
<td>NRAS, KRAS</td>
</tr>
<tr>
<td>Capillary arteriovenous malformation</td>
<td>RASA1</td>
</tr>
<tr>
<td>Cardiofaciocutaneous syndrome</td>
<td>BRAF, KRAS, MAP2K1, MAP2K2</td>
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<tr>
<td>Costello syndrome</td>
<td>HRAS</td>
</tr>
<tr>
<td>Hereditary gingival fibromatosis 1</td>
<td>SOS1</td>
</tr>
<tr>
<td>Legius syndrome</td>
<td>SPRED1</td>
</tr>
<tr>
<td>LEOPARD syndrome*</td>
<td>PTPN11 (type 1), RAF1 (type 2), BRAF (type 3)</td>
</tr>
<tr>
<td>Neurofibromatosis type 1</td>
<td>NF1</td>
</tr>
<tr>
<td>Noonan syndrome</td>
<td>PTPN11, SOS1, SOS2, RAF1, BRAF, KRAS, NRAS, RIT1, LZTR1</td>
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</tbody>
</table>

*aLEOPARD indicates multiple lentigines, electrocardiographic conduction abnormalities, ocular hypertelorism, pulmonic stenosis, abnormal genitalia, retardation of growth, and sensorineural deafness.

Data from Tidyman and Rauen.²
Cardiofaciocutaneous syndrome (Online Mendelian Inheritance in Man [OMIM] 115150) is a genetic disorder first described by Reynolds et al and is characterized by several cutaneous abnormalities, cardiac defects, dysmorphic craniofacial features, gastrointestinal dysmotility, and mental retardation. It occurs sporadically and is caused by functional activation of mutations in 4 different genes—BRAF, KRAS, MAP2K1, MAP2K2—of the RAS extracellular signal-regulated kinase molecular cascade that regulates cell differentiation, proliferation, and apoptosis.

As a RASopathy, CFC syndrome is a member of a family of syndromes with similar phenotypes, which includes mainly Noonan and Costello syndromes. Psychomotor retardation and physical anomalies, the common denominator of all syndromes, may be explained by the effects of the mutations during early development.

In CFC, relative macrocephaly, prominent forehead, bitemporal constriction, absence of eyebrows, palpebral ptosis, broad nasal root, bulbous nasal tip, and small chin commonly are found. The eyes are widely spaced and the palpebral fissures are downward slanting with epicanthic folds.

Follicular keratosis of the arms, legs, and face occurs in 80% of cases of CFC and ulerythema ophryogenes with sparse eyebrows in 90% of cases. Sparse, curly, and slow-growing hair is found in 93% of patients. Xerotic scaly skin, hyperkeratosis of the palms and soles, infantile hemangiomas, and multiple melanocytic nevi also may occur.

Cardiac abnormalities are seen in 75.7% of patients. Other features include mental retardation, delayed motor development, and structural abnormalities in the central nervous system, as well as seizures and electroencephalogram abnormalities. Unlike Noonan and Costello syndromes, it is unclear if patients with CFC syndrome are at an increased risk for cancer.

Noonan syndrome (OMIM #163950) is a disorder characterized by congenital heart defects, short stature, skeletal abnormalities, distinctive facial dysmorphic features, and variable cognitive deficits. Other associated features include cryptorchidism, lymphatic dysplasia, bleeding tendency, and occasional hematologic malignancies during childhood. This syndrome is related to mutations in the PTPN11, SOS1, SOS2, RAF1, BRAF, KRAS, NRAS, RIT1, and LZTR1 genes. The typical ear shape and placement in Noonan syndrome is oval with an overlapped helix that is low set and posteriorly angulated, which is uncommon in CFC syndrome. Noonan syndrome is characterized by an inverted triangular face; hypertelorism; blue or blue-green iris color; webbed neck; limited skin involvement, mainly represented by multiple nevi; and a much milder developmental delay compared to CFC and Costello syndromes.

Costello syndrome (OMIM #218040) is a rare condition comprised of severe postnatal feeding difficulties, mental retardation, coarse facial features, cardiovascular abnormalities (eg, pulmonic stenosis, hypertrophic cardiomyopathy, atrial tachycardia), tumor predisposition, and skin and musculoskeletal abnormalities. Costello syndrome is clinically diagnosed. This syndrome shows coarse facies with macrocephaly, downward-slanting palpebral fissures, epicanthic folds, bulbous nose with anteverted nostrils and low nasal bridge, full cheeks, large mouth, thick lips, large tongue, nasal papillomas, cutis laxa, low-set ears, short neck, diffuse skin hyperpigmentation, ulnar deviation of the hands, and nail dystrophy that are not observed in CFC. It is now accepted that the term Costello syndrome should be reserved for patients with HRAS mutation because of the specific risk profile of these patients. Remarkably, patients with Costello syndrome are at increased tumor risk (eg, rhabdomyosarcoma, neuroblastoma, bladder carcinoma).

The diagnosis of CFC syndrome is purely clinical: There have been many attempts to delineate the syndrome, but none of the described traits are pathognomonic. In 2002, Kavamura et al created the CFC index, a useful diagnostic approach based on 82 clinical characteristics and their frequencies in the CFC population.

Skin abnormalities are helpful manifestations to differentiate CFC syndrome from Noonan and Costello syndromes. Patients with CFC syndrome present with follicular hyperkeratosis and absent eyebrows. Absent eyebrows, narrowed temples, and Cupid's bow lip are hallmark features of CFC syndrome and are absent in Noonan and Costello syndromes. The presentation of palmar plantar hyperkeratosis also is a differentiating feature; in patients with Costello syndrome, it is found outside the pressure zones, whereas in those with CFC syndrome, it is present mainly in the pressure zones. Dermatologists can assist geneticists in the differential diagnosis of these syndromes.

The treatment of disorders with follicular plugging and xerosis is challenging. Emollients with urea, glycolic acid, and lactic acid could improve the appearance of the skin. Treatment with mutated MEK gene inhibitors is under investigation to restore normal development of affected embryos with CFC. This case and theoretical data show that skin manifestations can be helpful to differentiate CFC syndrome from other RASopathies such as Noonan and Costello syndromes.
REFERENCES


