The KBG Foundation is dedicated to providing support, assisting in research programs and advocating to raise awareness about the syndrome.

To stay up to date and join the private community follow us on Facebook & Twitter.

@KBGFdn #kbgfdn #everylinkmatters

What is KBG Syndrome?

INTERNATIONAL KBG SYNDROME AWARENESS DAY JUNE 19TH
What is KBG Syndrome?

KBG Syndrome is caused by a mutation in the ANKRD11 gene at location 16q24.3 (maybe more) which creates a shortening of a scaffolding protein in that region.

This short protein may be completely ineffectual or may be somewhat functioning which could contribute to the variety and severity of the symptoms.

**Appearance**

Nearly all patients present with large upper front teeth, bushy eyebrows and triangular faces. The 'KBG appearance' is quite distinctive and is usually the first noticeable trait along with developmental delay and other bone anomalies.

**Incidence**

New research suggests that less than 200 individuals worldwide have been diagnosed with KBG Syndrome. For no known reason, males seem to be more affected than females.

**Prognosis**

KBG has an excellent prognosis with no known impact on longevity.

**Treatment**

There is no singular treatment for KBG Syndrome but is determined on a case by case basis as symptoms arise.

**Common Traits/Symptoms**

- Facial dysmorphism
- Autistic Characteristics
- Abnormal hair implantation (low in front or back of skull)
- Brachy-clinodactylyous 5th finger (short/curved pinky)
- Macrodontia (large teeth) with oligodontia (absence of more than 6 teeth)
- Abnormal EEG with or without seizures
- Cognitive deficits/psychomotor delay
- Anteverted nostrils (upturned nose)
- Short femoral necks/hip dysplasia
- Cutaneous syndactyly, toes II/III
- Palatal defects (including uvula)
- Webbed/short neck
- Mild synophrys
- Brachycephaly/turricephaly
- Sternum abnormalities
- Cryptorchidism
- Abnormal ribs/vertebrae
- Epicanthal folds
- Delayed bone age
- Abnormal spine curvature
- Ptosis
- Prominent/high nasal bridge
- Long philtrum
- Short hand tubular bones
- Hearing loss
- Wide eyebrows
- Wormian bones in skull
- Thin upper lip
- Prominent/anteverted ears
- Strabismus
- Congenital heart defects

**1975**

The KBG Phenotype is defined by Herrmann, Pallister, Tiddy, Opitz
PMID: 1218237

**2004**

Genetic inheritance confirmed
PMID: 21782149

**2011**

ANKRD11 genetic mutation identified
PMID: 21782149

**2012**

16q24.3 Gene location reported
PMID: 22307766

**2014**

C-terminus leads to KBG mutation
PMID: 25413698

**2015**

KBG Foundation