Greek Warrior Helmet Facies (Wolf-hirschhorn Syndrome)

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Abstract

Wolf-Hirschhorn syndrome (WHS) is caused by a chromosomal deletion of the band 4p16.3 with characteristic craniofacial features -'Greek warrior helmet' facies (prominent glabella, hypertelorism, broad beaked nose and frontal bossing), high-arched eyebrows, protruding eyes, epicanthal folds, short philtrum, distinct mouth with downturned corners, micrognathia, dysplastic ears, preauricular tags. Till date there are very few case reports of Wolf-Hirschhorn syndrome.Here we report a case that had characteristic dysmorphic facies (Figure1) ‘Greek warrior helmet’ and was diagnosed as a case of WHS. But presence of Meningo-encephalocele and lissencephaly is rarely reported in literature in association with Wolf-hirschhorn syndrome till date.

Introduction

Wolf-Hirschhorn syndrome (WHS) is caused by a chromosomal deletion of the band 4p16.31 and was first described in 1961 by the Herbert L. Cooper and Kurt Hirschhorn2 and thereafter by Ulrich Wolf, and Hirschhorn and their co-workers in 1965 in the articles in the German scientific magazine ‘Humangenetik’3-4. Wolf-Hirschorn syndrome is a congenital malformation syndrome characterized by pre- and postnatal growth deficiency, developmental disability of variable degree, characteristic craniofacial features -‘Greek warrior helmet’ facies (prominent glabella, hypertelorism, broad beaked nose and frontal bossing), high-arched eyebrows, protruding eyes, epicanthal folds, short philtrum, distinct mouth with downturned corners, micrognathia, dysplastic ears, preauricular tags), and a seizure disorder. The typical craniofacial phenotype in combination with mental retardation, seizures, congenital heart defects, genital and renal anomalies is indicative of the diagnosis6-7. Most of patients with WHS have a de novo deletion, usually on the paternal chromosome8. De novo unbalanced translocations have also been described in 1.6% of WHS patients9. The translocation t (4;8)(p16;p23)10 may be the most frequent after t(11q;22q), which is the most common reciprocal translocation in humans11. It can be demonstrated by FISH(Fluorescence in situ hybridization) technique in cytogenetically normal parents of a affected offspring12. A familial translocation is responsible for only 5 – 13 % of cases13. WHS is mostly maternally inherited with a 2:1 ratio female to males14.

The Case

Here we report a case of female neonate born of a term gestation to a 24 yrs old second gravida. Baby was low birth weight (2kgs) and presented with dysmorphic facies and respiratory distress along with scalp deformity.
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It was a case of normal delivery and antenatal history was uneventful. However there was history of previous child's death in early neonatal period.

Baby was tachypneic (respiratory rate of 80/min with bilateral subcostal suction) and needed oxygen, 10% dextrose as intravenous fluid and kept under radiant warmer on servo-control mode.

On examination baby had characteristic dysmorphic facies- prominent glabella, hypertelorism, broad beaked nose and frontal bossing, high-arched eyebrows, protruding eyes, epicanthal folds, short philtrum, distinct mouth with downturned corners, and micrognathia- ‘Greek warrior helmet’ (Figure 1). Baby had polydactyly and syndactyly (Figure 2) of great toe of left foot and syndactyly (Figure 3) of ring finger of right hand. (Figure 4)

On palpation there was scalp defect with coming out of meninges and brain matter through the defect-meningo encephalocoele. (Figure 5). Baby showed generalized hypotonia. Baby developed seizure at 2 hours of age and capillary blood glucose was 86gms/dl, calcium 9gms/dl and seizure was controlled by injection phenobarbitone. Baby developed intractable convulsions which was not controlled with multiple anticonvulsants (phenobarbitone, Phenytoin, Lorazepam, Midazolam) and succumbed to death within nine hours of birth.

Investigations: CT scan of brain (Figure 6) showed evidence of wide (22mm) bony gap seen in occipito-parietal region with associated meningo-encephalocoele along with multiple septate cystic areas within it. There was evidence of multiple air-bubbles seen within the herniated component and also within brain parenchyma of occipital lobe. Cerebral sulci are effaced due to lissencephaly. No significant grey-white differentiation seen with loss of differentiation of different deep grey-matter regions. Apart from frontal horn of left ventricle, rest of ventricular system is not identified. Deformity of facial bone with micrognathia is suggestive from CT scan of brain.

Non-contrast CT scan (NCCT) of thorax (Figure 7) and abdomen (Figure 8) shows ill-defined areas of consolidation with air-bronchogram and associated ground-glass haze, seen to occupy almost whole of both lung fields, sparing anterior aspects of bases- suggestive of hyaline membrane disease. Mediastinal vascular structures were normal. No significant abnormalities were seen in abdominal structures from NCCT except hepatomegaly. Echocardiography showed no congenital structural abnormality.
Fig 3: Showing Syndactyly of the Right Hand

Fig 4: Whole Body Showing Generalised Hypotonia

Fig 5: Showing Meningo-Encephalocele

Fig 6: C T Scan Showing Meningo-Encephalocele

Fig 7: C T Scan Thorax

Fig 8: C T Scan Abdomen
Discussion

Wolf-hirschhorn syndrome is a rare 4p deletion syndrome with characteristic facial traits, CNS malformations (30%), low birth weight, short stature, muscular hypotonia, seizures (50-100%), congenital heart defects, colobomata of iris, genito urinary anomalies (25%), deafness, abnormalities of dermatoglyphics and skeletal abnormalities (60-70%). Battaglia evaluated 15 patients of which (33.3%) had heart lesions; (47%) had orofacial clefts; (87%) had a seizure disorder. A literature review of 22 reports of neuroimaging findings in WHS indicated that the most common findings were corpus callosum abnormalities (71%), focal white matter signal abnormalities (46%), lateral and third ventricle enlargement (42%), white matter volume reductions (42%), and periventricular cysts (29%). In our case there is presence of low birth weight, dysmorphic facies, syndactyly and polydactyly, generalized hypotonia, seizure, CNS anomaly in the form of meningo-encephalocele with lissencephaly.

Since the first Case report in 1961, about 159 cases have been published to date. Due to limited facilities for cytogenetic studies in India as well as in other developing countries, confirmatory diagnosis cannot be established in majority of cases. This may be the reason for the less number of published cases.

Conclusion

In a child with a clinical entity of mental retardation, associated with dysmorphic features and congenital anomalies, cytogenetic studies may help establish a specific diagnosis, which can predict the clinical outcome. There is no treatment available for this rare genetic syndrome but a multidisciplinary team approach is required including rehabilitation, speech communication therapy and genetic counselling. The risk of a parent having another child is unlikely unless the parent is a carrier of the disorder. Thus, parental advice for prenatal diagnosis should also be taken into consideration where facilities allow.

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