Large Angle Congenital Esotropia in a Child With Kabuki Syndrome: A Case Report

Talaat Hamdi 1, Ahmed Basheikh 2

1. Ophthalmology, Jeddah University, Jeddah, SAU 2. Ophthalmology, King Abdulaziz University, Jeddah, SAU

Corresponding author: Talaat Hamdi, tjhamdi@gmail.com

Abstract

Kabuki syndrome is an uncommon genetic disease associated with skeletal, cardiac, neurological, and ocular manifestations. Strabismus is an ophthalmic manifestation of Kabuki syndrome; however, it is infrequently documented in detail. We report a case of Kabuki syndrome in a patient who presented with large angle congenital esotropia. This case report highlights the importance of early eye examinations and subsequent interventions in patients diagnosed with Kabuki syndrome.

Introduction

Kabuki syndrome is a rare congenital genetic disease with an estimated prevalence of 1 in 32,000 Japanese births [1]. The name originates from the characteristic faces present in patients, which are similar to those of actors in traditional Japanese Kabuki theater. Other names that have been used include Nihkawa-Kuroki syndrome and Kabuki make-up syndrome. This disease is believed to be caused by mutations in the KMT2D and KDM6A gene, with multiple patterns of inheritance, including both autosomal dominant and X-linked inheritance patterns [1, 2]. In Saudi Arabia, Kabuki syndrome is not well reported due in part to rarity as well as lack of disease awareness. To our knowledge, this is one of a few case reports that document a large congenital esotropia in a child with Kabuki syndrome in Saudi Arabia.

Case Presentation

A two-year-and-ten-month-old boy presented with notable developmental delay, including requiring support to sit, lack of social smile, inability to recognize voices, and inability to pass objects between hands. The child was born at 39 weeks gestational age through planned cesarean section. The patient first presented to the hospital at three months of age with a weight of 4.3 kg, height of 50 cm, and head circumference of 40 cm. At the time of the present evaluation, his height was 89 cm (2nd percentile, UK-WHO growth chart) and weight was 10 kg (less than the 0.4th percentile), indicating severe growth retardation.

Perinatal history was unremarkable, and the family history was negative with no history of consanguinity. The patient was admitted to the hospital twice due to atypical febrile convulsions, was diagnosed with hypothyroidism, and prescribed thyroxine. The patient presented with dysmorphic features including down seated posture, prominent ears, hyperpigmented areas of the skin, high arched palate, wide space between the left metatarsal bones, and downturned mouth. Echocardiography showed a small restrictive ventricular septal defect (VSD) that was asymptomatic. Genetic team consultation was performed, which confirmed the diagnosis of Kabuki syndrome.

Ophthalmologic examination conducted by a pediatric ophthalmologist at around eight months of age showed a large congenital esotropia (ET) with a variable angle, and the Krimsky test showed about 70 prism diopters (PD) at near in the primary position. The patient’s visual acuity showed a mild grimace to light about each eye separately, with no fixation, and no following of objects. Retinal examination showed no aberrant abnormality besides mild disc pallor. The pupils were reactive with no afferent pupillary defect. Atropine refraction at eight months showed OD: +0.75 D sphere +0.75 D cylinder X 100, OS: +0.25 D sphere +0.50 D cylinder X 85. Repeat of atropine refraction at two years of age showed OD: -4.50 D sphere +1.25 D cylinder X 75, OS: -3.75 D sphere +0.75 D cylinder X 115 with myopic astigmatism that was corrected with glasses. Vision at the time of presentation was central, steady, and maintained in both eyes; the extraocular muscle ductions and versions showed minimal abduction defects in both eyes. The Krimsky test with correction showed around 45-50 PD at near. Anterior segment and retinal examination remain within normal limits at two years of age. Other ocular findings seen in the patient were long palpebral fissures, sparse arched eyebrows, eversion of the outer third of the lower eyelids, epicanthal folds, and bilateral ptosis as shown in Figure 1.

Open Access Case Report
DOI: 10.7759/cureus.8445

How to cite this article
Hamdi T, Basheikh A (June 04, 2020) Large Angle Congenital Esotropia in a Child With Kabuki Syndrome: A Case Report. Cureus 12(6): e8445. DOI 10.7759/cureus.8445
FIGURE 1: Large esotropia in Kabuki syndrome patient.

Photograph of the patient in this case demonstrates the characteristic facial features associated with Kabuki syndrome, including long palpebral fissure, eversion of the lateral one-third of the lower eyelid, arched eyebrows with sparseness in the lateral one-third, low set ears and depressed nasal tip. In addition, a large angle esotropia that has been present since birth is shown.

Discussion

Kabuki syndrome presents with a wide variability in clinical signs. Nonetheless, based on the analysis of 62 patients by Niikawa et al., the diagnosis of Kabuki syndrome requires five fundamental features: 1) characteristic face (100%) with long palpebral fissures, eversion of the lateral one-third of the lower eyelid, arched eyebrows with sparseness in the lateral one-third, prominent large ears and depressed nasal tip; 2) skeletal abnormality (92%); 3) dermatoglyphic abnormality (93%); 4) mental retardation (92%), and 5) short stature [3]. These features were all observed in this patient and led to a diagnosis of Kabuki syndrome.

Kabuki syndrome is associated with many ocular findings with prevalence ranges of 38% to 61% [4]. These include but are not limited to sparse hair in the lateral part of the eyebrows, strabismus, ptosis, amblyopia, refractive errors, sixth cranial nerve palsy, blue sclera, cataracts, retinal pigmentary changes, coloboma of the iris and retina, congenital corneal staphyloma, and Peter's anomaly [2, 4-7].

A previous expansive literature review found that strabismus was present in about 22% of 300 patients with a confirmed diagnosis of Kabuki syndrome [8]. However, these cases lack detailed documentation and only three cases reported the degrees of deviation most of which were found to be at small angles (12-25 PD). Additionally, these cases described older age patients, ranging from five to nine years of age [9]. Sharma and Dave reported one case of esotropia subsequently treated with 4-mm bimedial rectus recession with an approximate angle of deviation of 25 PD [9]. We report a case with a larger angle strabismus at a young age at approximately 70 PD, which decreased over time with full refractive error correction to approximately 45-50 PD.

Conclusions

In conclusion, we advocate the need for careful monitoring of the case before intervention as the strabismus angle might change over time from the first presentation. Besides, we emphasise the important need for an early eye examination in patients diagnosed with Kabuki syndrome to prevent visual impairment.

Additional Information

Disclosures

Human subjects: Consent was obtained by all participants in this study. Conflicts of interest: In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that no financial support was received from any organization for the submitted work. Financial relationships: All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. Other relationships: All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.
References

1. Milunsky JM, Huang XL: Unmasking Kabuki syndrome: chromosome 8p22-8p23.1 duplication revealed by comparative genomic hybridization and BAC-FISH. Clin Genet. 2003, 64:509-516. 10.1046/j.1399-0004.2003.00189.x
2. Matsumoto N, Niikawa N: Kabuki make-up syndrome: a review. Am J Med Genet C Semin Med Genet. 2003, 117:57-65. 10.1002/ajmg.c.10020
3. Niikawa N, Kuroki Y, Kaji T, et al.: Kabuki make-up (Niikawa-Kuroki) syndrome: a study of 62 patients. Am J Med Genet. 1988, 31:565-589. 10.1002/ajmg.1320310312
4. Cheon CK, Ko JM: Kabuki syndrome: clinical and molecular characteristics. Korean J Pediatr. 2015, 58:517-324. 10.3345/kjp.2015.58.9.517
5. Tanaka R, Takenouchi T, Uchida K, et al.: Congenital corneal staphyloma as a complication of Kabuki syndrome. Am J Med Genet A. 2012, 158:2000-2002. 10.1002/ajmg.a.35453
6. Bogershausen N, Wollnik B: Unmasking Kabuki syndrome. Clin Genet. 2013, 85:201-211. 10.1111/cge.12051
7. Dentici ML, Di Pede A, Lepri FR, et al.: Kabuki syndrome: clinical and molecular diagnosis in the first year of life. Arch Dis Child. 2015, 100:158-164. 10.1136/archdischild-2015-305858
8. Wessels MW, Brooks AS, Hoogeboom J, Niermeijer MF, Willems PJ: Kabuki syndrome: a review study of three hundred patients. Clin Dysmorphol. 2002, 11:95-102. 10.1097/00019605-200204000-00004
9. Sharma P, Dave V: Esotropia in Kabuki syndrome. J Pediatr Ophthalmol Strabismus. 2010, 48:1-3. 10.3928/01913913-20100524-05