Blepharophimosis sequence and diaphragmatic hernia associated with interstitial deletion of chromosome 3 (46,XY,del(3)(q21q23))

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Abstract
A case of blepharophimosis, ptosis, and epicanthus inversus (BPES) associated with prenatally diagnosed diaphragmatic hernia and interstitial deletion of the long arm of chromosome 3, del(3)(q21q23), is reported. Comparison with other cases of BPES resulting from 3q rearrangements indicate that this disorder, previously assigned to 3q2, can now be more accurately mapped to 3q23.

The association of blepharophimosis, ptosis, and epicanthus inversus (BPES) with structural rearrangements of 3q has been reported for both familial and de novo translocations and several interstitial deletions, including one forming part of a complex de novo rearrangement involving two other chromosomes and one derived from a familial inserted translocation. We report a further case of a small de novo interstitial deletion resulting in the classic features of BPES complicated by diaphragmatic hernia detected prenatally. The missing chromosome segment can be unequivocally defined, having only minimal overlap with several previously described deletions, allowing BPES to be mapped conclusively to 3q23.

Case report
Ultrasound examination of the fetus of a 25 year old primigravida showed an appearance suggestive of diaphragmatic hernia at 18, 19-5, and 24 weeks' gestation. Lymphocyte chromosome preparations from a cordocentesis sample taken at 24 weeks' gestation showed a small interstitial deletion of the long arm of chromosome 3 involving loss of 3q22 and proximal q23, namely 46,XY,del(3)(q21q23). Both parents had normal karyotypes. The pregnancy was terminated at 25 weeks' gestation. Necropsy showed a male fetus, weight 728 g, foot length 50 mm, crown-rump length 235 mm, head circumference 215 mm (consistent with 25 weeks' gestation), with the facial features of BPES (fig 2), telecanthus (inner canthal measurement 20 mm, expected 15 mm), shortening of the palpebral fissures (10 mm, expected 13-14 mm), and a beaked nose. There were no upper eyelid folds. Internally, the left hemidiaphragm was almost entirely absent, with the stomach, most of the left lobe of the liver, and loops of the small and large bowel in the thoracic cavity. No other abnormalities were noted. The fetal karyotype was confirmed in cells cultured from placental villi. A cell line, reference F93/0233, is available from this laboratory.

Discussion
BPES is most commonly encountered as either a new mutation or in families with autosomal dominant inheritance. There are probably two distinct but related subtypes of the inherited form: type I, the most common, transmitted by males, where affected females are infertile, and type II, transmitted by both males and females. This variable phenotype has been interpreted as evidence for BPES being a contiguous gene syndrome. Several reports of the distinctive facial features of this condition associated with structural rearrangements involving 3q2 suggest this region as the likely location of both the sporadic and dominant inherited forms of the disorder.
There remains, however, some ambiguity as to detailed mapping within 3q2, owing to the restricted resolution observable in some of the published partial karyotypes, the descriptive limitations of ISCN for microdeletions, and the potential for reinterpretation of breakpoints.

The present case, and those of Williamson et al. and Fryns et al., show the unequivocal loss of a common segment involving bands q22–23 with the deleted chromosomes having very similar appearances in spite of minor variations in interpretation of breakpoints. Three more distal deletions reported by Alvarado et al., Martsoff and Ray, and Al-Awadi et al., all involve bands q23–25. The minimal area of overlap is thus within q23, which corresponds well with a de novo rearrangement, 46,XY,t(3;4)(q23;p15.2) observed by Fukushima et al. and a 46,XX,t(3;8)(q23;p21.1) karyotype described by Cabral de Almeida et al. Jewett et al. reported a smaller deletion involving loss of q22 only, which they interpreted as placing the gene at the q22–23 boundary, but unfortunately this publication is an abstract without a photograph for comparison.

Two reports, however, suggest a more proximal location on 3q. De Die-Smulders et al. reported a father and son with typical BPES, both with 46,XY,t(3;11)(q21;q23) karyotypes, although the quality of the published photograph is not incompatible with alternative breakpoints at 3q23 and 11q25. The case of Fujita et al. is less helpful. The authors claim to show an interstitial deletion, del(3)(q12q23), but if this patient has a simple deletion, then the breakpoints indicated on their photograph would be q12 and q21 (not q23), although q23 is arrowed on their ideogram. Again the quality of the published photographs is not incompatible with alternative breakpoints and it is not possible to exclude a more complex rearrangement of 3q.

We interpret these data as indicating location of BPES within 3q23, with preference for the more proximal part of that band. The unusual presentation, in our case, of diaphragmatic hernia detected prenatally has not been associated with other published examples of 3q deletions with BPES and is not a feature of the familial form of the disorder.

Recognition of further cytogenetic microdeletions involving 3q23, particularly the q22 interface, will allow the localisation of a minimal region of overlap and the eventual molecular identification of the contiguous gene sequence whose deletion or loss of function is responsible for BPES.

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