Further refinement of the Usher 2A locus at 1q41

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Abstract
Usher syndrome (USH) is characterised by congenital sensorineural hearing loss and progressive pigmentary retinopathy. All three subtypes (USH1, USH2, and USH3) are inherited as recessive traits. People with Usher type 2 (USH2) have normal vestibular responses and moderate to severe hearing loss.

These syndromes have been found to be genetically heterogeneous, with a single locus for USH2 at 1q41 (USH2A), six loci for USH1, and one for USH3. Some USH2 families have been excluded from the 1q41 locus suggesting that a second, as yet unidentified, locus (USH2B) must exist. Linkage studies suggest that around 90% of USH2 families are USH2A.

Four USH2 families were analysed for linkage to markers flanking the USH2A locus. In one of these families a recombination event was observed in an affected subject which excludes the USH2A gene from proximal to the marker AFM143XF10 and defines this as the new centromeric flanking marker for the USH2A locus. A further recombination event in another patient from this family confirmed AFM144XF2 as the telomeric flanking marker.

The interval between these polymorphic markers is estimated to be 400 kb. This region is completely contained in each of three YACs from the CEPH library: 867g9, 919h3, and 848b9. This refinement more than halves the critical genetic interval and will greatly facilitate positional cloning of the USH2A gene.

Keywords: Usher syndrome; retinitis pigmentosa; genetics

Usher syndrome (USH) is characterised by congenital sensorineural hearing loss and progressive pigmentary retinopathy. In all cases the mode of inheritance is autosomal recessive. The frequency of USH in the USA has been estimated at around 5 per 100 000 people, and USH is thought to account for over half of all people with dual hearing and visual impairment.

A clinical classification of the phenotypic variants of Usher syndrome recognises the existence of three subtypes, Usher I, II, and III. Subjects with Usher syndrome type 1 (USH1) have profound sensorineural deafness and absent vestibular function. They do not benefit from traditional hearing aids and typically communicate by sign language. Subjects with Usher type 2 (USH2) have normal vestibular responses and moderate to severe sensorineural hearing loss, while those with Usher III experience progressive hearing loss.

These syndromes have been found to be genetically heterogeneous, with a total of nine genetic loci now reported. The gene MYO7A, located at 1q13, has been shown to be responsible for cases of Usher 1B. There are five additional loci for USH1, at 1q32 (USH1A), at 11p13 (USH1C), at 10q (USH1D), 21q (USH1E), and on chromosome 10 (USH1F). There is a single locus for USH2 at 1q41 (USH2A). USH3 has been linked to 3q25.

The original localisation of USH2A was to a 14.8 cM interval on 1q41 and this was later refined to a 2.1 cM region between markers D1S237 and D1S229. The construction of a yeast artificial chromosome (YAC) contig across this region, and the detection of recombination events in two separate families, permitted further refinement to a physical distance of 1.0 Mb between markers D1S474 and AFM144XF2.

Pieke-Dahl et al analyzed linkage to 1q41 markers in 29 Dutch families with clinical manifestations of USH2. Linkage to 1q41 was shown in 26 families indicating that around 90% of these families have USH2A. Several of the remaining families were excluded from the 1q41 locus suggesting that a second, as yet unidentified, locus (USH2B) must exist.

In the present study haplotype analysis of an inbred USH2A family has allowed us to determine a new centromeric flanking marker for this disorder and confirm AFM144XF2 as the telomeric flanking marker.

After gaining informed consent, genomic DNA was prepared from peripheral blood obtained from four unrelated USH2 families. Three of these were pedigrees from the Moorfields Eye Hospital retinal dystrophy register and one was ascertained in Pakistan. The patients in pedigree USH2(4) are of Sephardic Jewish extraction. Their parents, now dead, were uncle and niece. The three affected subjects were examined by two of the authors (DARB and ACB). All three had moderate congenital sensorineural hearing loss, without vestibular dysfunction, and had developed typical symptoms and signs of retinitis pigmentosa.

Non-radioactive PCR was performed in a 10 μl reaction with 300 ng of genomic DNA, 10 pmol of each primer, 200 μmol/l dNTPs, 1.5 mmol/l MgCl₂, and 1 unit of Taq DNA polymerase (Biotaq) in buffer provided by the manufacturer. A three stage PCR consisting of 35 cycles at 94°C, 50-62°C, and 72°C, each for
on data both from the Whitehead contig WC1-19 and the contig published by Sumegi et al., this region is covered by 11 YACs from the CEPH library: 867g9, 919h3, 848b9, 873a8, 836c7, 763d7, 762a6, 945f7, 798b4, 785h4, and 841g2. The critical genetic interval is completely contained on each of three of these YACs (867g9, 919h3, and 848b9).

On the contig WC1-19 there are three STSs (WI-3484, WI-3128, and WI-9496) within this interval, but no polymorphic markers that can be used in further refinement. At least four STSs on WC1-19 have been excluded by this refinement. WI-3484 and WI-3128 have no significant homology at the nucleotide level to any known genes or human cDNA sequences. WI-9496 is a partial cDNA (NCBI Entrez accession No Z39073) from a normalised infant brain library. Sequence database searching indicates homology between WI-9496 and several other human cDNA sequences including a human retina clone (Entrez accession No Y08993).

This refinement has more than halved the remaining genetic critical interval for this locus and will greatly facilitate the positional cloning of the USH2A gene.

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