USHER’S SYNDROME REVISITED

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ABSTRACT

Usher’s syndrome is a genetically inherited autosomal recessive disorder resulting in the double handicap of deafness and progressive blindness, known as retinitis pigmentosa. The disease is also associated with psychoses, mental retardation, and other major neurophysiological changes. It appears to be more common among Jewish individuals and consanguinous marriages. While it is rare in the general population (3 cases per 100,000 population), it is significantly prevalent among those who are deaf. Most patients are forced to give up their profession around age 30 or 40 or earlier, either because of advancing failure of sight leading to blindness at age 50 or 60, or due to the other disabilities of the condition.

Although a wide variety of treatments have been tried including surgery, endocrine therapy, vitamins, and transplants, at present the disease cannot be cured nor its course significantly altered. A program for prevention through high risk diagnostic screening, coupled with genetic counseling, is both feasible and practical.

In this report, we present two siblings with this syndrome, as well as a general review of the history and literature concerning this disorder.

MJIRI, Vol. 2, No. 2, 143-147, 1988

INTRODUCTION

Albrecht Von Graefe in Berlin in 1854 was the first to report the association of retinitis pigmentosa and congenital hearing loss. Usher in 1914, reporting a large series, added cataracts, speech disorders, and mental illness to the syndrome. Hallgren in 1954 presented 151 cases and added the finding of vestibular ataxia. Usher’s syndrome thus consists of retinitis pigmentosa and congenital and/or progressive sensorineural hearing loss, cataracts, speech disorders, and mental deficiency. Psychosis and vestibular ataxia are additional variable findings in Usher’s syndrome. The incidence of retinitis pigmentosa in congenitally deaf individuals has been variously reported to range from 3% to 10.4%, while hearing loss has been found in 3.4% to 6.3% of patients with pigment degeneration of the retina.

Liebreich in 1861 was one of the first to construct a series of pedigrees in which there appeared cases of deafness with retinitis pigmentosa in a survey in which he had examined 241 of the 341 deafmutes in Berlin and found 14 with retinal pigmentation. Stephenson and Chessman in Northern Ireland and the New York population study reported by Sank do not mention frequency of anomalies other than hearing impairment.

Etiology

While familial incidence of retinitis pigmentosa and deafness was noted by Von Graefe and Liebreich, Usher was the first to emphasize the familial nature of the syndrome and suggested that it constituted a specific genetic entity. Von Wibaut pointed out the striking preponderance of cases showing a recessive mode of inheritance as opposed to dominant inheritance, a hereditary pattern that is commonly seen in cases of isolated retinitis pigmentosa. Bell further emphasized the genetic aspects of the syndrome, and Lindenov presented evidence strongly supporting the contention that the two cardinal features of the sy-
drome were a pleotropic manifestation of a single recessive gene rather than effects of two separate genes. The hereditary pattern of Usher’s syndrome is usually autosomal recessive with 100% penetrance; the defect is at a single chromosomal locus.

Usher’s syndrome is believed to be transmitted by an autosomal recessive gene. Consequently, although it has been estimated that one out of 100 individuals is a carrier, the trait manifests in only three per 100,000. Other pertinent facts about the prevalence of Usher’s syndrome are that it appears to be more common among Jewish individuals and consanguinous marriages.

It is also known that deafness in not the most frequent degenerative process associated with retinitis pigmentosa. Some believe the genetic code for Usher’s syndrome begins to manifest itself during embryological development. The various lesions, although having different sites, are thought to have a common point of attack in the embryo.

Fraser, et al., describe Usher’s syndrome as a genetic error of metabolism and list two other syndromes (Pendred’s and Jervell Lang-Nielsen). It was theorized that the common feature of congenital deafness may be due to a hypersensitivity of the VIII nerve to metabolic and chemical toxins. Francois raised the issue of whether the action of the pathology of the gene is endocrinological, neural, or toxic; or whether it is simply based on diminution in the viability of retinal cells. There has been some debate over whether Usher’s syndrome is a bonafide genetic syndrome or three separate conditions.

**Clinical features**

The hearing loss is cochlear in type and is bilateral, often with mild to moderate preservation of low frequencies (from 125 to 500 Hz). Although the hearing loss is frequently congenital, it may not be recognized until the patient is one to two years old. It is generally severe and may be progressive. Vernon showed that from 5 to 10% of patients with congenital deafness have Usher’s syndrome.

The second cardinal feature of Usher’s syndrome, retinitis pigmentosa, frequently does not become apparent until late childhood or adolescence. The ocular symptom is usually night blindness. The visual fields are gradually restricted and visual acuity deteriorates. Cataracts are almost always located in the posterior cortical layers of the lens and are frequently observed as the disease progresses. Vision is further compromised in the fourth to sixth decades of life. Only minimal vision may remain, with progression of the visual loss to blindness. The earliest ophthalmologic sign is an abnormal finding on a dark adapted electroretinogram, which may permit the diagnosis of retinitis pigmentosa. In the presymptomatic state, the characteristic findings of the retina are a waxy, yellowish optic disk, narrow arteries, and bare corpuscular pigmentation in the periphery.

Not only does Usher’s syndrome involve a loss of vision and hearing, but Bossu and Luypaert found that most patients also lacked olfactory sensitivity. Thus, three of five special sensory modalities are involved. A loss of taste and smell has also been noted by the author among some genetically deaf children (a high risk group for Usher’s syndrome). The questions which arise from this and from Bossu’s finding concern not only the extent to which all sensory modalities may be involved, but also the possibility of a central lesion or at least a common degenerative process.

**Neuropathology and psychiatric findings**

Usher’s syndrome has special theoretical interest for the organically-oriented psychiatrist. The neurological degeneration associated with it involves the retina, a part of the brain proper. Possibly, the lesions causing hearing loss are central rather than peripheral. Consequently, an understanding of the degenerative process of Usher’s syndrome may have generality to other neurological and psychiatric diseases.

In the temporal bone, irregular degeneration of the stria vascularis, atrophy of the organ of Corti and the spiral ganglion, and both peripheral and central nerve degeneration with atrophy of the end branches of the labyrinthine vessels are evident on pathologic examination.

Another neurological structure affected by the disease is the vestibular apparatus. The reported prevalence of vestibular-based disturbances of equilibrium varies from 0-100%, with the most comprehensive study yielding a rate of 90%. Whether the site of lesion causing the balance disturbance is in the labyrinth or cerebellum is unclear, but the general consensus favors the labyrinth.

Psychiatric disorders are common is Usher’s syndrome. Hallgren found that of the 114 cases in whom he obtained or made a diagnosis, 23.2% (26 cases) were psychotic: 16 presented a schizophrenic-like picture, 3 showed depressive reactions, 2 had psychotic episodes and recovered, and 5 were hospitalized due to aggressiveness and agitation. Many reported auditory hallucinations. Paranoid tendencies and violence were common among these patients, and 13 of 26 psychotics were also mentally retarded. The overall prevalence of mental retardation in the total sample of 114 patients was 23.8%. The only remaining data on Usher’s syndrome and psychiatric illness comes from a survey of the total population in New York State Mental Hospit-
al. It was reported that 4.6% of all of the deaf patients hospitalized for mental illness had Usher's syndrome. Rainer and Streifler, et al obtained EEGs from 25 cases of retinitis pigmentosa with night blindness, and five relatives. Of these 30 cases, 14 had abnormal EEGs, consisting mostly of bilaterally symmetrical and synchronous slow waves.

**Prevention and management**

Despite circumstances which permit a significant reduction in the prevalence of this chronic, incapacitating disease, no preventive program has been developed. Five major steps in prevention are recommended by Vernon:

1. Education of professional persons who evaluate deaf and blind children about Usher's syndrome.
2. Diagnostic screening of all congenitally deaf children with an ophthalmic battery including ERG-EOG ophthalmoscopy examination, visual field tests, and dark adaptation measurements.
3. Evaluation of an audiological test battery plus vestibular function testing.
4. Similar ophthalmic and audiologic screening for relatives of deaf children identified as having Usher's syndrome.
5. Genetic counseling of identified individuals and their families.

**CASE REPORTS**

**Case 1**

A twenty-four year old white woman was referred to the ENT clinic because of bilateral hearing loss which had been noted at the age of three. The hearing loss had been progressive and she also complained of intermittent dizzy episodes with nausea. A review of symptoms at the time was negative.

The family history was positive; one sibling, an older brother, had a hearing loss. She had a past history of infectious hepatitis. An audiogram revealed a bilateral sensorineural hearing loss (Fig. 1). Otorhinolaryngologic examination was normal. The modified Kobrak ice caloric test was unremarkable. Blood tests including biochemistry, immuno-electrophoresis and urine analysis were all within normal limits. Plain roentgenograms of the petrous bones including Stenver's and transorbital views were normal. Eye examination including angiography and visual fields were reported as follows: arterial phase: generalized disturbance of the retinal pigment epithelium; the optic disc is well circumscribed, some areas of choroid retinal atrophy can be observed and remains non-fluorescent in other places; punctate remnants of the retinal pigment epithelium contrast with the intense background fluorescence.

Altogether gross attenuation of the vessels can be seen, which is nonfluorescent. In the peripheral retina, some bone-corpuscle pigmented aggregates can be seen.

The above observation shows retinitis pigmentosa of both eyes. If the two films of the angiography are compared, it shows roughly the same situation.

**Case 2**

A twenty-four year old brother of Case 1 was referred to the ENT clinic because of bilateral, progres-
Usher's Syndrome

Figure 2: Pape tone audiogram of patient 2

sive hearing loss accompanied with intermittent dizzi­
ness and nausea. The hearing loss was first noted at age
13. The family history was positive. His parents’ mar­
riage was not consanguinous. Two siblings out of six
had a hearing loss. His mother had a history of renal
disease. His father had developed squamous cell carci­
noma of the larynx which was treated with cobalt 60
irradiation. Audiometry revealed a severe bilateral
hearing loss (Fig. 2). The results of the remainder of the
ENT examination were unremarkable. Plain X-ray,
Stenver’s, transorbital and Towne’s views were all
normal. The patient noted night blindness and eye
exam included angiography and visual fields which
were reported as follows: arterial phase: generalized
disturbance of the retinal pigment epithelium; the optic
disc is well circumscribed; some areas of choroid retinal
atrophy can be observed, and remains non-fluorescent
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biochemistry and urine analysis were all within normal
limits. Modified Kobrak ice caloric testing was unre­
markable.

With the above findings, the diagnosis of Usher’s
syndrome was made. The following conclusions may be
drawn:

1. An autosomal recessive gene with 100% penetr­
ance located at a single chromosomal locus was found
to be responsible for the more profound hearing im­
pairment associated with Usher’s syndrome.

2. Without exception, all offsprings had Usher’s
syndrome when both parents had the syndrome, but no
case of deafness occurred among 52 offsprings when
only one parent had Usher’s syndrome, and the other
parent was deaf from some other autosomal recessive
gene.

3. With a few exceptions, known heterozygous
 carriers had a slight loss of hearing at all frequencies.
Signs of retinitis pigmentosa and cataract were found in
13% of all known heterozygous carriers.

4. The degree of hearing impairment was greater in
individuals with Usher’s syndrome than in those who
did not have the associated visual impairment.

5. Ophthalmological examinations are indicated for
all children with severe congenital hearing impairment
to aid genetic counseling.

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