Wildervanck or cervico-oculo-acoustic syndrome and MRI findings

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In 1952, Wildervanck described the first case of what he styled the cervico-oculo-acoustic (COA) syndrome. This comprises Klippel Feil’s (KF) anomaly (congenitally fused cervical vertebrae), congenital sensorineural deafness and Duane’s retraction syndrome (deficient abduction with retraction on adduction). Since that original paper, there have been further reports describing this triad, either completely or incompletely. A further case of this syndrome is reported and the first report of MRI head scan findings in this condition is presented. In addition, the origin of mirror movements observed as part of the KF syndrome are discussed.

Case Report
A 50 year old woman was admitted for the further evaluation of episodic right-sided facial weakness, associated with vague head pains. She had been well until March 1988, when she began to experience increasing tinnitus and deafness in her right ear. Two months later, she presented with sudden onset of right sided weakness which included the face. A CT head scan revealed a lacunar infarct in the left corona radiata, poorly pneumatized left mastoid air cells and generalised atrophy of the posterior fossa structures including the brainstem, with a large fourth ventricle. Her right sided weakness improved substantially but she developed further intermittent episodes of mild right-sided facial weakness and vague head pains. By the time she was seen in our unit, these symptoms had resolved completely and her only spontaneous complaint was of increased stiffness of the right arm.

The patient was known to have had abnormal eye movements since birth and was found to be completely deaf in the left ear on a routine school hearing examination at the age of 12. There was no other history of note. There was a family history of vascular disease; her mother had two cerebrovascular accidents (CVA), the first in her 60s and one of her brothers had died of a myocardial infarct at the age of 38. The patient had smoked at least 20 cigarettes a day before her CVA.

On examination, the patient was of short stature (1.47m) with the clinical features of the KF syndrome. The general medical examination was otherwise unremarkable. She had bilateral Duane’s syndrome, with absent abduction, retraction on adduction and narrowing of the palpebral apertures. She was orthotropic in the primary position and did not have diplopia. The oculo-cephalic reflexes on lateral gaze were absent but preserved small amplitude optokinetic nystagmus (OKN) was found. Up-gaze was normal, but downgaze was impaired although OKN was present. She was unable to converge. She had minimal right facial weakness and was totally deaf in the left ear, but hearing appeared to be relatively preserved in the right ear on bedside testing. The only abnormalities in the limbs were an increase in tone of the right arm, and mirror movements.

The combination of physical signs supported a diagnosis of Wildervanck’s syndrome, and an unrelated resolved right hemiparesis secondary to lacunar infarction. A repeat CT head scan did not reveal any change, four vessel aortic arch digital subtraction angiography demonstrated no significant pathology and an echocardiogram was normal. A full haematological and biochemical screen was normal. Pure tone audiometry confirmed complete deafness in the left ear, but also showed mixed high frequency loss in the right ear. A cervical spine radiograph revealed extensive developmental abnormalities in the upper and mid cervical spine extending down to C6 compatible with the KF anomaly. An MRI head scan (fig) showed marked atrophy of the brain stem from the pons to the midline cervical junction. There was a lesion measuring 1–2 mm in diameter in the pons to the left of the midline consistent with an infarct: the previously observed corona radiata infarct was not visible. The cerebellum was smaller than normal and the fourth ventricle was correspondingly dilated. There was failure of segmentation of the bodies of C3 and C4 with marked basilar invagination more pronounced to the left of the midline with indentation of the atrophied brain stem anteriorly.

Discussion
Hearing loss with KF anomaly, and hearing loss with an abducens nerve palsy had been documented before Wildervanck reported the triad of signs which now bears his name. There is a gradual transition in the clinical features between complete and incomplete forms of COA syndrome. Only one third of patients with KF anomaly have hearing impairment and this may be purely conductive, sensorineural or mixed. In COA syndrome, the unilateral or bilateral congenital
hearing loss is usually severe and of neurosensory origin though mixed hearing loss has been reported. The K-F syndrome may overlap with the stigmata of COA, Turner's and Noonan's syndromes and Wildervanck's syndrome is the most common multiple congenital abnormality found in association with Duane's syndrome. There is a tenfold female preponderance in COA syndrome. The cause is unknown, but Wildervanck suggests that it is due to polygenic heredity with sex limitation, though McKusick raises the possibility of sex linked dominance with fatal effect in the male. Chromosomal analysis has revealed normal patterns in females. A chromosome ring was demonstrated in one case though this may have been the consequence of pre-natal radiation exposure.

There are few complete cases of COA syndrome in the British literature. We report a further case which exhibited all the features of this syndrome. This is the first report to include MRI changes showing marked cerebellar and brain stem atrophy with basilar invagination and failure of vertebral segmentation. Our patient also demonstrated prominent mirror movements and has been investigated extensively with regard to the pathogenesis of this phenomenon. Necropsy of a patient with K-F anomaly and mirror movements revealed failure of pyramidal decussation in the cervical cord. This would leave little direct connection between the motor cortex of each hemisphere and its respective contralateral spinal cord. To generate movement of the contralateral extremities, the use of less specific alternate pathways such as those originating in the supplementary motor cortex would be required, and these would presumably give rise to bilateral activity. This case, however, was associated with cerebellar tonsil herniation and lack of fusion of the dorsal halves of the neural plate. Our patient did not have MRI evidence of such abnormalities and the left corona radiata infarct associated with right sided pyramidal signs provided evidence for pyramidal tract decussation. Electromyographic (EMG) recordings were made from both upper limb muscles and the results were compared with those obtained in normal control subjects. Limbal percutaneous electrical or magnetic brain stimulation applied to either hemisphere elicited bilateral and symmetrical short latency muscle responses in relaxed intrinsic hand muscles in our patient, while in normal subjects, unilateral brain stimulation only elicited contralateral muscle responses. Cross correlation analysis of ongoing single and multiunit needle EMGs recorded between muscles of left and right hands revealed a central peak in the cross-correlogram. No cross-correlogram peaks were found between left and right hand muscles in normal subjects. The number and time course of the central peaks in the cross-correlograms constructed between the firing of motor units on opposite sides of the body in the patient were similar to those found in cross-correlograms constructed between the firing of motor units from muscles on the same side of the body in the patient and in normal subjects. We concluded that the mirror movements in our patient resulted from motor commands that are distributed to motor neuron pools on the left and right side of the spinal cord via abnormally branched fast-conducting corticospinal tract fibres. The finding of bilateral long latency components of the cutaneous reflex in our patient strengthens the view that these components result from activity in a transcortical reflex pathway.

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