Congenital facial diplegia

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Congenital facial paralysis may occur alone or in association with other anomalies (Duke-Elder, 1964). The most common association is with the ocular motor nerves, particularly the VIth (Moebius syndrome).

Henderson (1939) reviewed 61 cases, showing a high incidence of motor palsies and other defects.

Case report
This patient, a boy aged 4 years, was referred to the Paediatric Department by his general practitioner who had noticed his stiff upper lip, and that he was unable to smile or wrinkle his face. In infancy he had had difficulty in sucking a bottle, but could feed from the breast. He was later able to swallow using a cup, without dribbling, but could not suck from a straw. He had had recurrent ocular infections until he was 2 or 3 years old.

Family history
There was no family history of muscular or neurological disease, and no consanguinity. There were no complications during pregnancy and the confinement was normal.

Examination
The patient had an expressionless face with no movement of the facial muscles (Buster Keaton facies) (Fig. 1). He was unable to close his eyes fully, smile, frown, raise his eyebrows, or purse his lips to whistle. Bell’s phenomenon was observed on attempted closure of the eyes (Fig. 2). The mouth

FIG. 1 General appearance at age 5

FIG. 2 Open eyelids on attempting to close the eyes

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Congenital facial diplegia was always slightly open and the upper lip prominent. The only movement of the mouth was that due to opening the jaw and to contraction of platysma in the neck. He had no difficulty in swallowing. The tongue was normal in size and could be protruded normally with no sign of twitching. The remainder of the examination gave normal results. Ocular movements were full. There was no nystagmus. Convergence was normal. There was watering of both eyes, but no discharge. Hearing was normal. The tear ducts were syringed and found to be patent.

TREATMENT
The symptoms were relieved by prescribing glasses with side-pieces to prevent dust or dirt from entering the eyes.

Summary
A case of congenital facial diplegia is described in which there was no involvement of the ocular muscles and no systemic defects.

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References
HENDERSON, J. L. (1939) *Brain*, 62, 381