Short report

Oculomotor control in Gilles de la Tourette syndrome

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SUMMARY Saccadic eye movements, fixation and smooth pursuit were studied in 28 children with Gilles de la Tourette syndrome and found to be normal. A link has been postulated between Gilles de la Tourette syndrome and other movement disorders. The results obtained in the present series do not support this hypothesis.

Gilles de la Tourette syndrome is a disorder characterised by multiple motor and vocal tics. Patients are able to suppress their tics for minutes to hours. Over periods varying from weeks to months, the intensity of tics shows a waxing and waning course. The onset of symptoms is in childhood.1-3 Forced gaze deviation and blepharospasm, common in progressive supranuclear palsy and in Huntington’s disease,4,5 are frequently observed in Gilles de la Tourette syndrome. It has been suggested that this common symptomatology may indicate a link between the syndrome and progressive supranuclear palsy and Huntington’s disease.4 Moreover, oculomotor abnormalities are known to be an important cause of failure in ability to perform tasks requiring eye-hand coordination.5 As neuropsychological examination has revealed disturbances of such abilities in Gilles de la Tourette syndrome,7 oculomotor was studied in this condition.

Patients

Twenty-eight children (21 boys, 7 girls) with Gilles de la Tourette syndrome were examined (mean age, 11:2 years; age range, 8 to 15 years). The patients entering the study came from the Department of Neuropsychiatry, Psychiatric Centre “Rosenburg” The Hague, the outpatients clinics of the Sophia Children’s Hospital Rotterdam, and the Department of Neuropsychiatry of the University Hospital Leiden. All patients met the DSM III (DSM III, 1980)8 diagnostic criteria and the criteria defined by Shapiro.1 Nineteen patients used low doses of the following drugs: clonidine (12 patients), pimozide (7), haloperidol (1), fluphenazine (1) or pipamperonchloride (dipiperon) (1). Three patients were on a combination of drugs and the remaining nine patients were medication free.

Methods

Eye movements were measured with a newly developed infrared reflection technique combined with a computerised portable eye movement processing system.9-12 The eye movement measuring device was incorporated in a spectacle frame attached to the head. Accordingly, eye position was measured with respect to head position. Head movement did not interfere with the measuring properties of the technique. With a bandwidth of D.C. to 100 Hz (—3 dB) the resolution of the IR technique is 2 min arc in both horizontal and vertical directions. The system is linear within 3% for eye-rotation up to 20° horizontally and up to 14° vertically, measured from the centre position. The drift of the IR-technique is negligible, so that it does not disturb the DC mode used to record eye position. The movement of both eyes was measured simultaneously while patients were sitting in a room with low level of ambient illumination. Whenever patients were not able to prevent head movements, despite a support at the back of their head, one of the investigators gently stabilised the head with his hands. In this set-up, the target for saccades consisted of an array of red gallium-arsenide light-emitting diodes (LED) of wavelength 635 nm, mounted in a black cylindrical screen. Each diode subtended a visual angle of 0.16° at a viewing distance of 1.5 m. If
Normal saccades towards a visual target in a patient with Gilles de la Tourette syndrome. Upper trace: target, amplitude 20°. Lower trace: saccadic eye movements with normal latency and normal peak velocity. Saccades to the right: upward deflection, saccades to the left: downward deflection.

Fig 1


Results

During the test performance, one patient did not cooperate and was therefore omitted. Of the single unwanted saccade and only so if the final target position was reached. The number of the saccades was counted and expressed as a percentage of the total number of distracting stimuli presented.

Normal values of saccadic latency, peak velocity and frequency of unwanted saccades were compared with normal values previously established in our laboratory. Normal values of saccadic eye movements are as follows: latency maximal (mean +2SD) 270 ms, peak velocity minimal (mean -2SD) 135°/s (6-7° amplitude), 260°/s (13-4° amplitude) and 300°/s (20-1° amplitude). Frequency of unwanted saccades towards distracting visual stimuli is 10% or less of the frequency of the distracting stimuli.

Fixational square-wave jerks larger than 1° in amplitude and at a frequency above 10/min were considered abnormal.14 They were counted and expressed as number/s. Macro-square-wave jerks were defined as having an amplitude above 7°. Despite the occurrence of square-wave-jerks during fixation, reliable calibration of eye position was possible in all patients.

Patients were kept alert by verbal encouragement. After each tenth stimulus, a short break was introduced to avoid fatigue.

In the statistical evaluation the sign test and the Spearman rank correlation test were used.

Smooth pursuit was evoked by a target moving along a line through the fovea. A laser beam was reflected on to a screen by a mirror galvanometer mounted on an optical scanner servomotor. The scanner system is driven by software running in a Z80-based microprocessor system. Sinusoidal stimuli with peak-to-peak amplitude of 24° and with frequencies of 0-1, 0-3, 0-5 and 0-7 Hz were selected. Smooth pursuit gain of less than 0-95 was considered abnormal.

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Results

During the test performance, one patient did not cooperate and was therefore omitted. Of the
remaining 27 patients, latency and peak velocity of voluntary saccades and of visually evoked saccades were normal. No multiple step saccades and no hypermetria were seen. Fixation was normal; no square wave jerks and no unwanted saccades towards distracting stimuli during fixation occurred.

Smooth pursuit showed no abnormalities. Smooth pursuit gains were normal for all the target frequencies tested (fig 2).

No differences were found between medicated and unmedicated patients.

Discussion

Saccadic eye movements, fixation of a midline and an eccentric target, and smooth pursuit were normal in our children. Of particular interest in our patients is the absence of unwanted saccades towards distracting visual stimuli. The substantia nigra pars reticulata inhibits saccade related cells in the superior colliculi and in this way prevents the occurrence of unwanted saccades towards visual stimuli.13 14 This inhibitory function of the substantia nigra pars reticulata is thought to be the main basal ganglia output modulating saccadic eye movements.15 In contrast to Huntington's disease, where unwanted saccades towards distracting visual stimuli occur with abnormally high frequency,8 9 18 the inhibitory neostriatal output appears to be normal in Gilles de la Tourette syndrome. Therefore, despite the fact that the clinical findings in Gilles de la Tourette syndrome in some aspects may be comparable to those in Huntington's disease and progressive supranuclear palsy,4 our results do not support a link between Gilles de la Tourette syndrome and these movement disorders. Furthermore the disabilities revealed in neuropsychological tests in Gilles de la Tourette syndrome3 appear not to be due to oculomotor disturbances.

A variety of centrally acting drugs has been shown to impair various aspects of oculomotion in man.19 Low doses of neuroleptics as given to our patients appear to have had no effect on eye movement parameters.

References