Pallidal activity during dystonia: somatosensory reorganisation and changes with severity

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Abstract
A woman with progressive, medically intractable right upper limb dystonia underwent a pallidotomy with only transient improvement. During the procedure her dystonia became more severe as she repeatedly made a fist to command in order to provoke dystonia transiently (movement provoked dystonia). Comparisons within cells in the internal segment of the globus pallidus (Gpi) disclosed that the firing rate was the same at rest, with making a fist, and during movement provoked dystonia. However, the firing rate compared between cells decreased significantly throughout the procedure as the patient made a fist repeatedly. During the second half of the procedure the firing rate of cells in the Gpi was similar to that in hemiballismus. The proportion of cells in the Gpi which responded to sensory stimulation was significantly higher in dystonia (53%) than in hemiballismus (13%). These results suggest that pallidal activity can correlate inversely with the severity of dystonia, perhaps due to activity dependent changes in neuronal function resulting from repeated voluntary movement.

Keywords: globus pallidus; Parkinson's disease; apomorphine; dystonia; plasticity

Dystonia is a movement disorder characterised by sustained muscle contractions leading to twisting movements and abnormal postures.1 The neuronal mechanism of dystonia is unknown but has been hypothesised to fit the hyperkinetic model of movement disorders.2–5 Pallidotomy is reported to be an effective treatment for dystonia and offers the opportunity to study neuronal activity in the disease.6–8 We report a patient in which a progressive decrease in firing rates in the internal segment of the globus pallidus (Gpi) correlated with worsening dystonia during a pallidotomy, as the patient made repetitive voluntary movements.

Case report and methods
A 45 year old right handed woman presented to focal dystonia. An MRI and an evaluation for secondary causes of dystonia were unremarkable. Seven years after onset the dystonia had become incapacitating and refractory to medical treatment with levodopa/carbidopa, clonazepam, trihexyphenidyl, and botulinum toxin; she had been on no medications for the 3 months before the procedure. Either at rest or after right arm movement (for example, making a fist) the arm often moved slowly from its anatomical position to posture characterised by shoulder adduction and internal rotation, elbow flexion, and wrist flexion so that the fist was held below the opposite axilla (dystonic posture). She underwent a stereotactically guided pallidotomy, with microelectrode exploration lasting 245 minutes before a 6 mm diameter lesion was made in the Gpi. Postoperatively she had moderate improvement in the dystonia which lasted for 10 days.

Microelectrode guided pallidotomy was carried out as previously described.5 10 The microelectrode signal was amplified (DAM-80, WPI, Sarasota, Florida, USA), analog filtered (−6 dB below 300 and above 10 000 Hz) and stored on a video tape recorder (Channel risetime 25 µs: Vetter Model 4000, Rebersberg, Pennsylvania, USA). The activity in the patient with dystonia (three trajectories, Gpi-17 cells, (Gpe)-20) was compared11 with that in other patients undergoing pallidotomy including activity (146 trajectories, Gpi-934, Gpe-1127) in 40 consecutive “control” patients with medically intractable bradykinesia, fluctuations, and dyskinesias of Parkinson’s disease (age range 43–74, Hoehn and Yahr “off” 2.5–4), and in one patient with hemiballismus (three trajectories, Gpi-13, Gpe-12).9 Five of the patients with Parkinson’s disease (age range 43 to 69 years; “off” Hoehn and Yahr stage 3 to 4) were studied intraoperatively both “off” and “on” —that is, after apomorphine—as previously described.9 Patients were classified as Parkinson’s “on” over the period of time after administration of apomorphine during which the rate of movements increased, tone decreased, and the patient felt “on”, compared with the period before apomorphine was administered and “off” after these measures returned to baseline. The number of finger taps in 10 seconds in these patients was significantly lower (t test, p<0.001)
Pallidotomy was performed under local anaesthetic block. The locations of the optic tract, internal capsule, cellular, and acellular regions were determined and compared with atlas maps to establish the location of the Gpi. Responses to sensory stimuli were sought for all cells in the GPi and GPe during passive movements of upper and lower limbs, both ipsilateral and contralateral to the recording site. Lesions were made within the part of the Gpi where cells responded to passive movements of the limbs.

Postoperatively, neuronal activity was studied for all isolated cells that could be recorded for 120 s including intervals with the arm at rest, during making a fist, and after the arm had achieved the final dystonic posture. The action potentials of single neurons were discriminated and digitised at 10 kHz by a standard shape fitting package (Explorer, Brainwave, Thornton, Colorado, USA) and times of occurrence of action potentials were stored at a clock rate of 10 000 Hz. All quantitative results are firing rates of isolated single neurons calculated as number of action potentials divided by sampling epoch, reported as the median and range and analysed by non-parametric statistical methods (Kruskal-Wallis ANOVA, Statistica, Statsoft, Oklahoma City, OK, USA).

**Results**

The figure illustrates that Gpi firing rates in the patient with dystonia at rest (40.8/s, range 20.1–53.6, n=16) were significantly different from Parkinson’s “off” (91/s, range 84.5–103.8, n=21) but were not significantly different from hemiballismus (30/s, range 18.6–41.0, n=13, p>0.05) or Parkinson’s “on” (44.3/s, range 33.9–64.3, n=17, p>0.05). The low frequency modulation and pauses in the
spike train seemed to be less pronounced in the patient with dystonia than in the patient with hemiballismus but more common than in patients with Parkinson's disease, either “on” or “off”.

Throughout the exploration of the basal ganglia (245 minutes) the patient with dystonia repeatedly (n=39) made a fist for 1–2 minutes in response to command. As illustrated in the figure, firing rates were not significantly (Kruskal-Wallis ANOVA) different between rest (45.9/s, range 20.1–53.6), making a fist (33.7/s, range 19.6–51.5), and the final dystonic posture (40.7/s, range 17.7–48.3). The dystonia became worse throughout the procedure as indicated by the surgeon's observation, the patient's self-report, and the significant (r=-0.89, n=37, p<0.001) decrease in the time between making a fist and achieving the final dystonic posture (fist to posture interval).

The firing rates in the Gpi decreased significantly as a function of time during the procedure in the patient with dystonia (r=-0.66, n=16, p<0.01) and as a function of the number of times a fist was made (r=-0.68, n=42, p<0.01). No significant change was seen in either Gpi activity or clinical state during the procedures in any of the six patients with Parkinson's disease in whom such analysis was performed. The fist to posture interval was directly related (r=0.50, n=16, p<0.05) to the firing rate in the Gpi. During the second half of the procedure in the patient with dystonia (time after beginning of procedure>180 min), firing rates in Gpi (16.9/s, range 12–22) were similar to those in the patient with hemiballismus (30/s, 18.6–40.8).

Firing rates in the GPe were lower in the patient with dystonia at rest (45.9/s, range 27.9–65.3, n=15, Kruskal-Wallis ANOVA, p<0.05) and in hemiballismus (18.1/s, range 7.5–26.5, n=12, p<0.001) than during Parkinson's “off” (53.4, range 44–80, n=17) or “on” (72.2, range 41.2–105.1, n=9). Firing rates in the GPe were not significantly different (Kruskal-Wallis ANOVA) between rest (45.9/s, range 27.1–65.3), making a fist (48.2/s, range 30–65.3), and the final dystonic posture (44.9/s, range 24.1–61.7). Firing rates in the GPe were not significantly related to fist to posture interval (r=0.31, m=15.8, p<0.25, n=16). Therefore, Gpi firing rates in dystonia decreased progressively throughout the procedure while those in Gpe were tonically less than in Parkinson's disease.

A higher proportion of cells in Gpi of the patient with dystonia responded to somatosensory stimulation (53%, 9/17), than in hemiballismus (13%, 2/15, p<0.05 Fisher's exact test), Parkinson's “off” (23/67, 34%, p>0.05) and Parkinson's “on” (6/19, 32%, p>0.05). The proportion of Gpi sensory cells was constant in the first (5/9) and second half (4/8) of the procedure in the patient with dystonia. In the two trajectories (one parasagittal plane) through Gpi cells responding to sensory inputs consistently had RFs including the dystonic upper limb (RFs on wrist or hand 5/17 cells). Similar consistency of RFs was never found in a single parasagittal plane of 40 patients with Parkinson's disease.

**Discussion**

The baseline firing rates of cells in the Gpi decreased into the range found in hemiballismus as the patient with dystonia made repetitive voluntary movements and the dystonia increased in severity. Clinically, the increase in dystonia with voluntary movement is so characteristic that the standard scale for dystonia rates both the disability from dystonia and the degree to which it is provoked by movement. The firing rates compared within cells in the Gpi were not significantly altered from resting levels during either making a fist or movement provoked dystonia. These results suggest that inputs to Gpi in dystonia may undergo activity dependent changes in relation to voluntary movement. During the second half of the procedure firing rates in Gpi are less than in Parkinson's disease ‘on’ or ‘off’, suggesting that as dystonia becomes more severe neuronal activity changes to fit the hyperkinetic model.

The number of cells with receptive fields was significantly higher (53%) in dystonia than in hemiballismus (13%). Furthermore, all cells with receptive fields were in the dystonic limb which indicates an enlarged sensory representation of the dystonic limb, similar to that found in the thalamus of patients with dystonia.

Lesions of the Gpi do not uniformly relieve dystonia, even after the Gpi has reorganised its sensory map, as in the present case (Lozano AM, personal communication, Vitek JL, personal communication, and see Jankovic et al, Vitek et al, and Vitek and Lenz). Therefore, sensory changes in the pallidum alone do not explain dystonia. However, they may distinguish dystonia from hemiballismus as features of both are consistent with the hyperkinetic model.

In a preliminary report of activity in the basal ganglia during dystonia firing rates of cells in the Gpi were found (48/s) to be similar to those seen in the patient and intermediate between those in hemiballismus and normal monkeys. Activity dependent changes in cellular firing were not noted. Decreased firing rates in the Gpi could explain the disinhibition of cortex in focal dystonia as a result of decreased γ-aminobutyric acid inhibition from the Gpi to the pallidal relay nuclei of the thalamus. The presence of dystonia with “normal” firing rates early in the procedure plus the failure of the lesion to relieve dystonia suggest that altered baseline firing rates are not the primary driver of dystonia in this case.

The inverse relation between dystonia and firing in the Gpi cannot be due to activity dependent changes occurring in the pallidum as within cells analysis disclosed that movements did not alter the firing of pallidal cells. Furthermore, altered activity in the Gpi is not explained by the indirect pathway as decreased firing rates seen in the Gpe would be predicted to increase firing rates in the Gpi via the indirect pathway. If the inverse pathway is excluded then the inverse relation may result...
from increased activity in the direct pathway from the putamen to the Gpi. The excitatory corticostriatal connection terminates on N-methyl-D-aspartate, (an excitatory amino acid neurotransmitter) receptors which have the potential to explain activity dependent changes in striatal activity.\(^5\)\(^{19–21}\) On this basis we hypothesise that the inverse relation between Gpi cellular firing and dystonia is the result of activity dependent changes in striatal activity transmitted to the Gpi through the direct pathway.

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