We first started thinking that cortex must be important in Parkinson's disease when Alan R. Crossman did some experiments in rats, showing transient reductions in 6-hydroxydopamine-induced spontaneous turning behavior after cortical lesions [1]. The experiments were elegant, but the lesions were large and did not block the turning, suggesting a kind of competition rather than a causal influence of cortex in the turning behavior. Similar conclusions plagued the many attempts to decide which of the brainstem pathways were the substrate of the turning behavior that followed the destruction of dopamine cells unilaterally, for review see, Arbuthnott and Wright [2]. However, as we finished a study of the anatomy of the basal ganglia [3] we concluded that the final output from the striatum came through the output nuclei of the basal ganglia: the globus pallidus pars interna (entopeduncular nucleus in rodents) and the substantia nigra pars reticulata, to a small nucleus in the ventral thalamus (ventromedial -VM- in the rat). Tracing the output from that nucleus brought us back to layer 1 of the cortex, close to where the search started in layer V [3]. This result had the basal ganglia appearing to be a loop 'linking' layer 5 to the superficial level of the cortex. Not a very likely scenario, nevertheless it did prepare us to look for an involvement of cortex in the consequences of dopamine destruction. Furthermore, the evidence was already there, the striatonigral spiny projection neurons (SPNs) that carried the first stage of the basal ganglia output, have cortical synapses on the spines [4]. When we studied the electron microscopic (EM) anatomy of the striatum without dopamine, there were obvious differences in those SPN spines [5-7]. There were fewer of them: we counted them stereologically in serial EM sections and found statistically fewer spines when the dopamine had been removed. As the theory about the differences in the two output pathways from the striatum developed, we started a long series of experiments where we identified the cells on which the spines were counted. By then, we were not alone and the final publication brought together the laboratories of Susan R. Sesack, Ariel Y. Deutch, Jim D. Surmeier, and ourselves [8]. It may be that we missed some dopamine D1 cells that were also denuded of spines [9], but the major effect was robust across all our studies. Therefore, damage to the dopamine input to the striatum, somehow spread to the cortical synapses on the spines of the SPNs. We did most of the work on rats but we also checked that the effect occurred in Parkinsonian patients. In fact, in post mortem human brain the effects were even more marked, with a 27% reduction in spine numbers compared with the 15% in the rats [10].

Our results had two consequences that had to be explored: firstly there were many more spines lost than originally had dopamine synapses on them: an idea that supported Fuxe and Agnati’s idea about volume transmission of dopamine signaling [11,12]. Secondly, such a denervation must have consequences for the cortical involvement in basal ganglia functions and indeed, dopamine functions. A series of experiments that were the beginning of a long-term collaboration between myself and Jeff Wikens showed that in slices of the rat corticostriatal system we could convert the usual long-term depression that followed tetanic trains of cortical input, into long-term potentiation of the corticostriatal synapses [13]. This introduction did little except prepare us to look for the major action of dopamine to be expressed in the corticostriatal synapses, an idea that had been around for some time [14-17]. On the other hand, a

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more convincing argument resulted from the discussion with Dieter Jaeger that saw me working with his team on the substrate of deep brain stimulation (DBS) [18]. The justification had many aspects, but as neurophysiologists we had struggled with the problem of what was stimulated by electrical stimuli. Exploring this problem and considering that effective DBS delivers 2.5-3.5 V of 60-90 µs pulses at high frequency (130-180Hz) [19], results from excitability 'strength-duration' curves were interesting: a 50 µs pulse was very unlikely to stimulate anything that was not myelinated, therefore an action on the subthalamic nucleus (STN) cells themselves, was an improbable result and magnified since those cells could fire faster than the 130Hz. Therefore, we agreed that the likeliest substrate for stimulation was the presence of corticofugal fibres that run past the simulating electrodes. Our experiments showed that stimulation in the STN of rats, clearly induced antidromic potentials intracellularly recorded from layer V cells in the motor cortex. Furthermore careful analysis of the evoked potentials, showed that the activity was centered in layer V and that a possible mechanism of action might be the interruption of the synchronized oscillatory activity in cortex by the antidromic driving [20].

Two other pieces of evidence encouraged us, the first was the discovery of human data of antidromically generated slow waves in patients undergoing DBS, as recorded during surgery for the placement of electrodes [21-23], and the second a publication using not electrical but optogenetic manipulations of STN in 6-hydroxydopamine-lesioned mice. Exciting or inhibiting STN neurons did not recover the animals, but high frequency stimulation of the cortical layer V cells in Thy1-cre rodents did release the animals from the results of the lesions, by relieving bradykinesia and increasing distance and speed of locomotion [24]. Optogenetic techniques have been improved but the result is still the same [25], except that the important part of the stimulation might be the ‘hyperdirect pathway’ from cortex directly to the STN [26-27], rather than the whole corticofugal axon bundle. In the meanwhile, we had shown with different collaborators that the threshold for recovery of motion in rats made akinetic by dopamine blockade, was the same as that of the cortical antidromic wave [28,29]. Finally, in freely moving animals, we could demonstrate...
the synchronization of cortical neuronal firing in lesioned animals and the desynchronization that resulted from stimulation in STN, that also drove the cortical layer V neurons antidromically. The antidromic activity was stochastic because of on-going activity in freely moving animals, but high rates of antidromic driving, effectively desynchronized the electroencephalogram (EEG) [30]. Later there were some modelling studies that supported the likelihood that the antidromic activity was most effective in desynchronizing the EEG oscillations, [31,32]. We wrote a summary of this series of experiments for a conference in Dusseldorf in 2017, with the idea that not many people had heard of the results [33]. Since then, there have been several research papers about the ‘early response’ from the cortex to single pulses of DBS [34,35] with the extra, that even effective pallidal stimulation for Parkinson’s disease, also involved the same ‘fast response’ and there is even a suggestion that without frontal cortical invasion, DBS is less effective [36]. Not surprisingly the EEG field evoked responses have been used as a means of identifying the effect STN site for stimulation [37].

An interesting engineering development might make for a critical test of the idea, by placing a pair of stimulating electrodes that would make stimulation of the ‘internal capsule’ more difficult [38]. Clinical papers have investigated the best site in the STN for DBS [39] and two recent papers have suggested that its efficacy may separate Parkinson's disease into categories by length and severity of the disease [40], or even of disease subtypes while off medication according to the Unified Parkinson’s disease Rating Scale [41]. There has been a resurgence of information about beta power modulation within high beta frequency bands [42-45], and in deep brain structures during DBS [46-48]. There have also been interesting ideas both in modelling papers reviewed recently by Humphries et al. [49] and recordings in vitro by Aparicio-Juarez et al. [50], where cortical stimulation is sufficient to re-normalize, at least partially, the abnormal firing patterns in the cortex to single pulses of DBS [34,35] with the extra, that even effective pallidal stimulation for Parkinson's disease, also involved the same 'fast response' and there is even a suggestion that without frontal cortical invasion, DBS is less effective [36]. Not surprisingly the EEG field evoked responses have been used as a means of identifying the effect STN site for stimulation [37].

The influence of Parkinson’s disease in cortical physiology and anatomy has been evaluated in some studies that have emphasised the role of motor cortex in the disease [55-57,47,41]. These studies indicate that as the disease progresses there are consequences for the cortex. Can cortical stimulation replace DBS? Probably not, although the original test in monkeys seemed promising [58-59] and there is a reasonable positive review article [60] although it cannot be said to be a mainstream treatment.

Finally, the cortical involvement in Parkinson's disease has as long a history as the discovery of cognitive effects of the disease. To my surprise and far from my cautious attribution of cortical disturbances to the symptoms of the disease, Jose A. Obeso has recently suggested cortical malfunction as the cause [61,49]. Such an initial influence of cortex on the symptoms has the obvious advantage of helping with the problem of the early stage of the disease in patients. The initial signs are nearly always localized and often to one side and to a few motor acts. It is hard to see how the diffusely projecting dopamine system could result in such localized dysfunction. On the other hand, cortical lesions certainly have well described local actions as for example seen following a stroke. There remains only the way to link the cortical malfunction with the dopamine system and Obeso’s surprising and thought provoking suggestion is that local overactivity in corticofugal pathways might pass on, or perhaps even initiate, synuclein overproduction in dopamine terminals that could be passed back to destroy the whole neuron. There are gaps for sure, but this explanation of a cortical source of the disease, rather than the symptoms appearing after the dopamine loss, seems reasonable and may even suggest new therapies.

Competing Interests

The authors declare that they have no competing interests.

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