THE SUBTHALAMIC NUCLEUS AND THE EXTERNAL PALLIDUM: TWO TIGHTLY INTERCONNECTED STRUCTURES THAT CONTROL THE OUTPUT OF THE BASAL GANGLIA IN THE MONKEY

E. SHINK,* M. D. BEVAN,t J. P. BOLAMt and Y. SMITH*~

*Centre de Recherche en Neurobiologie, Hôpital de l’Enfant-Jésus and Université Laval, Québec, Canada
†MRC Anatomical Neuropharmacology Unit, and University Department of Pharmacology, Mansfield Road, Oxford OX1 3TH, U.K.

Abstract—The aim of the present study was to elucidate the organization of the interconnections between the subthalamic nucleus and the two segments of the globus pallidus in squirrel monkeys. By making small deposits of tracers in the two segments of the globus pallidus, we demonstrate that interconnected neurons of the subthalamic nucleus and the external pallidum innervate, via axon collaterals, the same population of neurons in the internal pallidum. Furthermore, this organizational principle holds true for different functional regions of the pallidum and the subthalamic nucleus.

Injections of biotinylated dextran amine were made in the dorsal (associative), ventrolateral (sensorimotor) and rostromedial (limbic) regions of the internal pallidum. Following these injections, there were rich clusters of labelled terminals in register with retrogradely labelled perikarya in related functional regions of the subthalamic nucleus and the external pallidum. At the electron microscopic level, the majority of labelled terminals in the external pallidum displayed the ultrastructural features of boutons from the subthalamic nucleus and were non-immunoreactive for GABA, whereas those in the subthalamic nucleus resembled terminals from the external pallidum and displayed GABA immunoreactivity. In both cases, the synaptic targets of the labelled terminals included labelled neurons. These observations suggest that the biotinylated dextran amine injected in the internal globus pallidus was transported retrogradely to perikarya in the external pallidum and the subthalamic nucleus and then anterogradely, via axon collaterals, to the subthalamic nucleus and the external pallidum respectively. This suggestion was supported by injections of biotinylated dextran amine or Phaseolus vulgaris-leucoagglutinin in regions of the external pallidum that corresponded to those containing retrogradely labelled cells following injections in the internal pallidum. The clusters of labelled cells and varicosities that resulted from these injections were found in regions of the subthalamic nucleus similar to those labelled following injections in the internal globus pallidus. Furthermore, terminals from the external pallidum and the subthalamic nucleus converged on the same regions in the internal globus pallidus.

The results of the present tracing study define the basic network underlying the interconnections between the external segment of the globus pallidus and the subthalamic nucleus, and their connections with the output neurons of the basal ganglia in primates. Copyright © 1996 IBRO. Published by Elsevier Science Ltd.

Key words: basal ganglia, pallidum, pallidosubthalamic pathway, subthalamopallidal pathway, tract-tracing method, post-embedding immunocytochemistry.

On the basis of electrophysiological and anatomical data, models of the functional organization within the basal ganglia thalamocortical circuit have been suggested.5,5 These models state that the striatal output reaches the internal pallidum (GPI) via two different pathways, a “direct” inhibitory projection from the striatum to the GPI, and an “indirect” pathway, which involves a GABAergic projection from the striatum to the external pallidum (GPe), an inhibitory projection from the GPe to the subthalamic nucleus (STN) and an excitatory projection from the STN to the GPI. Recent anatomical data revealed the existence of another branch of the indirect pathway which includes a direct projection from GPe to GPI. 8,30,34,39,40,64,65,66,71 According to these models,2,2 the output of the basal ganglia is under the control of inhibitory and excitatory influences arising from the direct and the indirect pathways, respectively. By virtue of their polarities, activation of the direct pathway facilitates movement by disinhibition of thalamocortical neurons, whereas activation of the...
indirect pathway inhibits movements by increasing the inhibition of thalamocortical cells. A balance between the activity of the two pathways is thus essential for the normal control of motor activities.

However, the exact mechanism by which the direct and indirect striatal afferents to GPi interact to control motor behaviour is poorly understood. The anatomical organization of the direct striatal output to GPi has been thoroughly studied in various species and appears to be highly specific. However, the organization of the indirect pathway is still obscure. The major reason for this is the small size and the heterogeneous functional organization of the STN. The combination of these two features complicates the interpretation of data obtained after injections of anterograde tracers in the STN. In these anterograde tracing studies, the injections of tracers resulted in a diffuse and largely unspecific labelling of terminals that extended throughout the entire extent of the pallidal complex in primates. On the basis of these findings, the STN was considered as a source of non-specific excitatory influences on the globus pallidus. Another possible interpretation of these data is that, in fact, the projection is highly specific but the injections involved neighbouring neuronal groups that projected to different regions of the globus pallidus. The relative size of the injection sites in the STN compared to the size of the STN itself makes this possibility very likely. Other investigators addressed this issue by means of retrograde transport from the globus pallidus to the STN, but in these cases the injection sites covered large areas of the GPe and GPi. We have therefore re-examined the organization of the interconnections between the two segments of the globus pallidus and the STN in primates. Since recent studies have emphasized the importance of the projection from the GPe (or globus pallidus in non-primates) to the GPi (or entopeduncular nucleus in non-primates), one of the aims of the present study was to examine the location of neurons in the STN that innervate these interconnected regions. This was achieved by placing iontophoretic deposits of biotinylated dextran amine (BDA) in interconneced regions of the GPi and GPe.

The results of this study have been presented in abstracts.

**EXPERIMENTAL PROCEDURES**

Six adult male squirrel monkeys (Saimiri sciureus) were used in the present study. Each animal was anaesthetized with a mixture of ketamine hydrochloride (Ketaset, 70 mg/kg, i.m.) and xylazine (10 mg/kg, i.m.) before being fixed in a stereotaxic frame. The depth of anaesthesia was determined by monitoring heart rate, respiration rate, muscle tone, as well as corneal and toe-pinching reflexes. The surgical and anaesthesia procedures were performed according to the guidelines of the Canadian Council on Animal Care. In four animals, BDA was injected bilaterally in different regions of the pallidal complex, while in another animal, a unilateral injection of BDA was performed in GPi. In the last animal, BDA was injected in both pallidal segments on one side of the brain and *Phaseolus vulgaris* lectin (PHA-L) was delivered in GPe on the other side. The stereotaxic coordinates were derived from the atlas of Emmers and Akert. The two tracers were injected iontophoretically using the same parameters. The BDA (Molecular Probes, Eugene, OR, U.S.A., 5% in distilled water) and PHA-L (2.5% solution in phosphate buffer, 0.01 M, pH 8.0) were loaded in glass micropipettes (tip diameter 20-30 μm) and delivered with a 7-μA positive current for 20 min by a 7 s on/7 s off cycle. For the first two days after the surgery, the animals received injections of analgesic (buprenorphine; 0.01 mg/kg, i.m.) twice daily.

After seven to 10 days, the animals were deeply anaesthetized with an overdose of pentobarbital and perfuse-fixed with 500-700 ml of cold oxygenated Ringer solution followed by 1.21 of fixative containing 3% paraformaldehyde and 0.5% glutaraldehyde in phosphate buffer (PB; 0.1 M, pH 7.4). The unbound aldehydes were then washed out by perfusing with 11 of cold PB (0.1 M, pH 7.4). After perfusion, the brains were dissected out from the skull, cut in 10-mm-thick blocks in the transverse plane and stored in phosphate-buffered saline (PBS, 0.01 M, pH 7.4) until sectioning. The blocks were then cut in 60-μm-thick transverse sections with a vibrating microtome, collected in cold PBS and treated for 20 min with sodium borohydride (1% in PBS). The sections were repeatedly washed in PBS and processed to reveal the tracers according to protocols suitable for light and electron microscopy.

**Histochemical localization of biotinylated dextran amine**

**Light microscopy.** A series of sections was washed in PBS and processed to reveal the transported BDA. They were placed in an avidin-biotin-peroxidase solution (ABC; Vector Laboratories, Burlingame CA, U.S.A.: 1:100 in PBS including 1% bovine serum albumin (BSA) and 0.3% Triton X-100) for 12-16 h at room temperature. They were then washed in PBS and Tris buffer (0.05 M, pH 7.6) before being placed in a solution containing 3,3'-diaminobenzidine tetrahydrochloride (DAB, 0.025%; Sigma Chemicals, St Louis, MO, U.S.A.), 0.01 M imidazole (Fisher Scientific, Nepean, Ontario, Canada) and 0.006% hydrogen peroxide for 10-15 min. The reaction was stopped by repeated washes in PBS. The sections were then mounted on to gelatin-coated slides.
Fig. 1.
Electron microscopy. A series of sections including the pallidal complex or the STN were processed for the localization of the transported BDA in the electron microscope. The sections were placed in a cryoprotectant solution (PB, 0.05 M, pH 7.4, containing 25% sucrose and 10% glycerol) for 20–30 min. After having sunk, they were frozen at −80°C for 20 min. They were then thawed and washed many times in PBS before being processed to localize BDA according to the protocol described above, except that no Triton X-100 was added to the ABC solution and the incubation was carried out at 4°C for 48 h. Once the BDA was localized, the sections were washed in PB (0.1 M, pH 7.4), postfixed in osmium tetroxide (1% in PB; 0.1 M, pH 7.4) for 20–30 min, and dehydrated in a graded series of alcohols and propylene oxide. Uranyl acetate was added to the 70% ethanol (40 min) to improve the contrast in the electron microscope. The sections were then embedded in resin (Durcupan ACM, Fluka) on microscope slides and cured for 48 h at 60°C.

Immunohistochemical localization of Phaseolus vulgaris leucoagglutinin

The sections including the PHA-L injection site in GPe as well as a series of sections containing GPe and STN were prepared for the localization of PHA-L in the light microscope. They were first incubated with a goat anti-PHA-L antiserum (Vector Labs; 1:2000 in PBS containing 0.3% Triton X-100 and 1% normal rabbit serum) for 16 h. This was followed by an incubation with biotinylated rabbit anti-goat IgG (Vector Labs; 1:200 in PBS; 0.3% Triton X-100/1% normal rabbit serum) for 2 h and then an incubation in ABC (1:100 in PBS/0.3% Triton X-100/1% BSA) for two additional hours. The incubations were all carried out at room temperature. The peroxidase bound to PHA-L was revealed with DAB (see above). Once the reaction was completed, the sections were mounted on gelatin-coated slides, dehydrated and a coverslip was applied with Permount.

Analysis of the material

The sections containing the injection sites were examined with a Nikon light microscope equipped with a camera lucida to determine the extent of the tracers in their respective targets. The sections of GPe and STN prepared for electron microscopy were chosen from the case illustrated in Fig. 6, whereas the sections selected in GPe were taken from the case illustrated in Fig. 5. Once located, regions that contained rich plexuses of BDA-positive terminals were cut out from the slides and glued to the top of resin blocks with cyanoacrylate glue. Serial ultrathin sections were then cut on an ultramicrotome (Reichert, Ultracut E), collected on a series of sections containing GPi and STN were prepared for the localization of the transported BDA in the electron microscope. The sections were placed in a cryoprotectant solution (PB, 0.05 M, pH 7.4, containing 25% sucrose and 10% glycerol) for 20–30 min. After having sunk, they were frozen at −80°C for 20 min. They were then thawed and washed many times in PBS before being processed to localize BDA according to the protocol described above, except that no Triton X-100 was added to the ABC solution and the incubation was carried out at 4°C for 48 h. Once the BDA was localized, the sections were washed in PB (0.1 M, pH 7.4), postfixed in osmium tetroxide (1% in PB; 0.1 M, pH 7.4) for 20–30 min, and dehydrated in a graded series of alcohols and propylene oxide. Uranyl acetate was added to the 70% ethanol (40 min) to improve the contrast in the electron microscope. The sections were then embedded in resin (Durcupan ACM, Fluka) on microscope slides and cured for 48 h at 60°C.

Post-embedding immunocytochemistry for GABA

A series of sections including the STN or GPe were processed for post-embedding immunocytochemistry for GABA. The post-embedding immunogold procedure was carried out with an antiserum raised in rabbit against GABA. The production, characterization and specificity of this antiserum have been described in detail elsewhere.

Briefly, a series of adjacent ultrathin sections were preincubated for 10 min in Tris-buffered saline (TBS; 0.05 M, pH 7.6) containing 0.01% Triton X-100. This was followed by an overnight incubation at room temperature with the primary antiserum (1:5000 dilution) diluted in TBS/0.01% Triton X-100. The sections were then washed three times (2 × 10 min and 1 × 30 min) in TBS/0.01% Triton X-100 followed by TBS (0.05 M, pH 8.2) for 10 min. They were then incubated with the gold-conjugated goat anti-rabbit IgG (BioCell; 1:25 in TBS 0.05 M, pH 8.2) for 90 min at room temperature, washed in distilled water and stained with uranyl acetate (1% in distilled water) for 90 min. Finally, after having been washed in distilled water and stained with lead citrate, they were examined with a Philips EM 300 electron microscope.

Control experiments

The specificity of labelling was tested by incubation with solutions in which the primary antiserum was replaced with non-immune rabbit serum. After such incubation, the tissue was devoid of gold particles indicating that the GABA immunolabelling was due to the primary antibodies. Another series of control grids was incubated with GABA antiserum that had undergone liquid-phase pre-adsorption with structurally related amino acids conjugated to ethanamine with glutaraldehyde. Each antiserum was preadsorbed against taurine, GABA, glutamate and glutamine conjugates. After such incubation, the tissue was almost completely devoid of gold particles in the cases where the GABA antiserum was pre-adsorbed with GABA glutaraldehyde conjugates, whereas pre-adsorption of the antiserum with other amino acids conjugates had no effect on the intensity of staining. These control experiments indicate that the GABA immunostaining described in the present study is specific.

RESULTS

Location of injection sites

In order to analyse the interconnections between the GPe, STN and GPi, the following approaches were used. Firstly, we injected BDA in different regions of the GPi. Secondly, we determined the location of the retrogradely labelled cells in the GPe.
Fig. 2.
following these injections. Thirdly, we delivered BDA or PHA-L in those regions of the GPe that were found to contain retrogradely labelled cells. Using this approach, we succeeded in injecting interconnected pools of neurons in the dorsal third (1 in GPi, 2 in GPe) (Figs 1, 3, 4) and the ventrolateral region (2 in GPi, 3 in GPe) (Figs 2, 5, 6, 7) of the GPi and GPe. In another case, BDA was injected in the ventromedial part of the GPi, without injection in the corresponding region of the GPe (Fig. 8). Most of the animals received injections of BDA, except in one case where PHA-L was delivered in the ventral part of the GPe (Fig. 7). In general, the injections were confined to their respective targets except for a slight contamination of the dorsal part of the pallidum after injections in the ventral GPi (Figs 2A, 5E, 8D). In the cases that received injections of BDA in the dorsal third of the GPe, the injection sites damaged the medullary laminae (Figs 1B, 4C).

Pattern of labelling following deposits of tracers in the internal and external pallidum

Each of the injections in either the GPi or the GPe resulted in a similar pattern of labelling in the corresponding division of the pallidal complex and the STN.

Deposits of tracer in the GPi led to the labelling of dense clusters of axons and axonal varicosities that were largely in register with labelled neurons in both the GPe and the STN (Figs 1A,C,E, 2A,C,E, 9A,B, 10A,B). Only a few of the labelled neurons were found outside the region of axon labelling. In both structures the labelled cells were often stained in a Golgi-like manner and were interspersed with unlabelled neurons (Figs 9A,B, 10A,B). The perikarya and proximal dendrites of the labelled and unlabelled neurons were often closely apposed by the labelled axonal varicosities (Figs 9A,B, 10B).

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Correspondence and topography of labelling in the subthalamic nucleus after biotinylated dextran amine injections in interconnected regions of the internal and external pallidum

The injection sites in the GPi or GPe and the subsequent labelling in the pallidum and the STN are illustrated schematically in Figs 3-8. After BDA injections confined to the dorsal third of the GPi (Figs 1A, 3E), a cluster of labelling was found in the central part of the STN (Figs 1C, 3G-J). Apart from a few scattered perikarya, the labelled varicosities and cell bodies were in complete register at different rostrocaudal levels of the STN (Fig. 3G-J). These injections in the dorsal GPi also resulted in two clusters of labelled varicosities in register with retrogradely labelled cells that were confined to the dorsal third of the GPe (Figs 1E, 3C,D). The labelling was predominantly in sectors of the GPe located 0.5–1.0 mm more rostral than the injection site. The rostral and caudal poles of the GPe were devoid of labelling (Fig. 3A,F).

Injections of BDA into the sector of the GPe where neurons were labelled after injections in the dorsal third of the GPi (compare Figs 1B and 1E, 3C and 4C) led to dense aggregates of labelling in the STN and the GPi. In the STN, clusters of labelled varicosities were in register with labelled perikarya and were confined to the sector of the nucleus that was labelled after the BDA injection in the GPe (compare Figs 1C and 1D, 3H,I and 4H,I), except that there was a slight lateral shift. More labelled cells were found outside the main cluster of labelling after this injection than after injections in the GPi (Figs 1C,D, 3G-J, 4G-J). The injection in the GPe also led to a dense cluster of labelling in the GPi (Figs 1F, 4E) that was located precisely where the BDA injection site was in the preceding case (compare Figs 1A and 1F, 3E and 4E).

In another case (not shown), the BDA injection was larger and more medial in the dorsal part of the GPe and involved the internal medullary lamina. The pattern and location of labelling in the STN corresponded to that described for the preceding case (Fig. 4G-J), except that the labelled structures were slightly shifted ventromedially. In the GPe, the pattern of labelling was similar to that obtained after the lateral injection (Figs 1F, 4E).

The BDA injections confined to the ventrolateral two-thirds of the GPi (Figs 2A, 5E) led to dense aggregates of labelling in the dorsolateral region of the STN, but excluding the lateralmost part of the nucleus (Figs 2C, 5G-J). The majority of labelled cells in the STN were in register with the labelled varicosities although a few were located more ventrally (Figs 2C, 5H,I). In the GPe, a dense cluster of labelled cells and varicosities was confined to the central part of the structure (Figs 2E, 5C,D). Like the injections in the dorsal third of the GPi, the densest zones of staining in the GPe were 0.5–1.0 mm more rostral than the injection site in the GPi (Fig. 5C,D).

An injection of BDA confined to the ventral region of the GPe that contained retrogradely labelled cells after BDA injection in the ventrolateral GPi (compare Figs 2B and 2E, 5C and 6C) led to a cluster of labelling in the dorsolateral STN (Figs 2D,F, 6G,I). This labelling was in the same region of the STN that was labelled after tracer injections in the ventrolateral GPi (compare Figs 2C and 2D, 5H,I and 6H,I), except that the rostromedial pole contained labelled elements after injection in the GPe but not after that
Fig. 3. Schematic drawing through the globus pallidus (A–F) and the STN (G–J) of the squirrel monkey to illustrate the distribution of labelled fibres (sinuous lines), terminals (small black dots) and perikarya (red dots) after BDA injection in the dorsal third of GPi (E). One red dot indicates three retrogradely labelled cells. Note the close overlap between the retrogradely labelled cells and the labelled fibres and terminals in GPe and STN. The anteroposterior coordinates are indicated in parentheses for each drawing.
Fig. 4. Schematic drawing through the globus pallidus (A–F) and the STN (G–J) of the squirrel monkey to illustrate the distribution of labelled fibres (sinuous lines), terminals (small black dots) and perikarya (red dots) after BDA injection in the dorsal third of GPe (C). One red dot indicates three retrogradely labelled cells. Note that the location of the injection site corresponds to that where the retrogradely labelled cells were found after injections in the dorsal part of GPi (see Fig. 3). The anteroposterior coordinates are indicated in parentheses for each drawing.
Fig. 5. Schematic drawing through the globus pallidus (A–F) and the STN (G–J) of the squirrel monkey to illustrate the distribution of labelled fibres (sinuous lines), terminals (small black dots) and perikarya (red dots) after BDA injection in the ventrolateral two-thirds of GPi (E). One red dot indicates three retrogradely labelled cells. The anteroposterior coordinates are indicated in parentheses for each drawing.
Fig. 6.
in the GPi (compare Figs 5G and 6G). None of the retrogradely labelled cells were outside the cluster of labelled varicosities following the injection in the GPe (Figs 2D,F, 6G–J). A rich cluster of labelled varicosities occurred in the region of the GPi where the injection of BDA was made in the preceding case (compare Figs 5E and 6E) except that the labelling extended slightly more in the rostromedial and mediolateral planes. Similar patterns of labelling occurred following PHA-L injections in the same regions of the GPe (compare Figs 6C and 7C) except that the labelling was not as extensive in the GPi (compare Figs 6 and 7).

In the single case that received an injection of BDA confined to the ventromedial region of the GPi (Fig. 8), a dense cluster of labelling was found in the dorsomedial part of the STN (Fig. 8G–I). In contrast to injections in the more lateral part, the injection located more medially in the GPi led to labelling in the rostral pole of the STN that was generally placed more medially (compare Figs 5H,I and 8G–I). In both cases, a few retrogradely labelled cells were found more ventral than the main cluster of labelling (Fig. 8G–I). In the GPe the cluster of labelled axons, varicosities and cells was found in the rostroventral region just caudal to the anterior commissure (Fig. 8B).

Ultrastructural characteristics of labelled terminals in the subthalamic nucleus and external pallidum after injections of biotinylated dextran amine in the internal pallidum

In order to determine the exact source of the labelled boutons in the STN and GPe after BDA injections in the GPi, we analysed their ultrastructural features, determined their pattern of synaptic innervation and tested whether they were immunoreactive for GABA. In the electron microscope, the three types of labelled terminals that were examined. The remaining elements in contact with the type 1 boutons (Fig. 9D) and formed symmetric synapses with dendrites (50%) and perikarya (50%). None of the elements in contact with the type 1 boutons were labelled with BDA. A third category, comprising labelled terminals (7%); n = 39), referred to as type 3 boutons, contained ovoid synaptic vesicles, formed symmetric synapses with unlabelled dendritic shafts and displayed GABA immunoreactivity (Fig. 10C).

Ultrastructural features of labelled terminals in the internal pallidum after biotinylated dextran amine injections in the external pallidum

In the light microscope, two types of varicosities appeared to compose the clusters of labelling in the GPi after injections in the GPe (Fig. 11A,B). The first category consisted of large-sized boutons (2–4 μm in diameter) that formed pericellular baskets around the perikarya and proximal dendrites of GPi neurons (Fig. 11B). The second class comprised medium-sized varicosities (0.5–2.5 μm in diameter) that were associated preferentially with dendritic shafts (Fig. 11B). In the electron microscope, the three types of labelled boutons described above were found. The type 1 terminals (n = 47; 44%; 11D,E) and the type 2 (n = 53; 50%; Fig. 11C) terminals accounted for 94% of the 106 labelled terminals that were examined. The remaining terminals (n = 6; 6%) displayed the ultrastructural features of the type 3 boutons (Fig. 10C).

DISCUSSION

The neuronal network underlying the interconnections between the GPe and the STN and the relationship of these structures with the GPi has been the subject of many studies. 1,27,29,38,45,47, 50,56,57,61,65. Elucidation of this network is of critical importance in our understanding of how the “indirect” pathways influence neurons of the GPi and hence the output of the

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Fig. 6. Schematic drawing through the globus pallidus (A–F) and the STN (G–J) of the squirrel monkey to illustrate the distribution of labelled fibres (sinuous lines), terminals (small black dots) and perikarya (red dots) after BDA injection in the ventral part of GPe (C). One red dot indicates three retrogradely labelled cells. Note that the location of the injection site corresponds to that where the retrogradely labelled cells were found after injection in the ventrolateral region of GPi (see Fig. 5C). The anteroposterior coordinates are indicated in parentheses for each drawing.
Fig. 7.
basal ganglia. By the use of a tracer that is transported in both retrograde and anterograde fashions we describe the basic circuit underlying these interconnections. The results demonstrate that the interconnections between the two divisions of the pallidal complex and the STN are much more specific than previously suspected. The basic circuit is such that interconnected groups of neurons in the GPe and STN innervate, via axon collaterals, the same population of neurons in the internal pallidum, i.e. output neurons of the basal ganglia. A basic principle that emerged is that the topology of the tight interconnections respects the functional organization of both pallidal segments and the STN (Fig. 13). These findings have important implications for the understanding of the neuronal mechanisms that underlie the roles of the basal ganglia in the control of motor activity in primates.

**Interpretation of labelling: possible sources of labelled boutons in the subthalamic nucleus and external pallidum**

As mentioned in the introduction, the data obtained so far relating to the organization of the subthalamopallidal projection in primates must be interpreted with caution because of technical problems inherent to the approaches used to investigate this issue. In the present study, these problems were overcome by the use of the neuronal tracer BDA. The fact that BDA can be injected iontophoretically enabled us to deliver relatively small quantities of tracer that were mostly confined to the different functional territories of the GPe and GPI. After injections of BDA in the GPs, dense plexuses of labelled terminals in register with retrogradely labelled perikarya were found in the STN and GPe. The presence of retrogradely labelled perikarya in these structures is consistent with previous studies showing the existence of projections from the STN and GPe to the GPI. However, the occurrence of labelled varicosities could not be explained by the anterograde transport of BDA from the GPI to the STN and GPe as these projections are thought to be very sparse if they exist at all. The most likely explanation for this labelling is that following injections of BDA in the GPI, the tracer was transported retrogradely to STN and GPe neurons, and then anterogradely, via axon collaterals, to terminal fields in the GPe and STN, respectively (Fig. 12A). Thus, we propose that most of the labelled terminals in the STN are derived from neurons in the GPe which retrogradely transported the BDA that was delivered in the GPI, and then antero-

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Fig. 7. Schematic drawing through the globus pallidus (A–F) and STN (G–J) of the squirrel monkey to illustrate the distribution of labelled fibres (sinuous lines) and terminals (small black dots) after PHA-L injection in the ventral part of GPe (C). Note that the location of the injection site corresponds to that where the retrogradely labelled cells were found after injection in the ventrolateral region of GPI (see Fig. 5C). The anteroposterior coordinates are indicated in parentheses for each drawing.
Fig. 8. Schematic drawing through the globus pallidus (A–F) and the STN (G–J) of the squirrel monkey to illustrate the distribution of labelled fibres (sinuous lines), terminals (small black dots) and perikarya (red dots) after BDA injection in the ventromedial region of GPi (D). One red dot indicates three retrogradely labelled cells. The anteroposterior coordinates are indicated in parentheses for each drawing.
results of this part of the study indicate that this is indeed the case. Thus, the deposits of BDA or PHA-L in regions of the GPe that contained retrogradely labelled cells after the GPi injections, led to clusters of labelled terminals in regions of the STN that corresponded closely to those labelled after injections in the GPi (compare Figs 1C and 1D; 2C and 2D). As one would predict, the deposits of tracer in the GPe also led to rich plexuses of labelled terminals in the regions of the GPi that had received the injections.

Fig. 9. Photomicrographs (A,B) and electron micrographs (C,D) showing various features of labelled structures in the STN after BDA injections in the ventrolateral part of the GPi. The micrographs in A and B were taken from the cluster of labelling shown in Fig. 2C. Note that BDA-labelled varicosities are apposed to the surface of both labelled (arrowheads in B) and unlabelled perikarya (arrowhead in A). The arrows in A indicate retrogradely labelled cells that are not contacted by BDA-positive terminals. (C,D) The two types of terminals visualized in the STN after BDA injections in the GPi. (C) A type 2 terminal that forms an asymmetric synapse (arrowheads) with a BDA-labelled dendritic shaft (d/BDA). The type 2 terminals accounted for 11% of the labelled boutons in the STN after BDA injections in the GPi. (D) Illustrates an example of a type 1 terminal that represents 89% of the total number of labelled boutons seen in the STN. The labelled terminal forms a symmetric synapse (arrows) with a dendritic shaft (d1). In a section that has been processed for the post-embedding immunohistochemical localization of GABA, the BDA-containing terminal is associated with a large number of gold particles which indicates that it displays GABA immunoreactivity (indicated by G). Another terminal in the same section (G1) is also immunoreactive for GABA whereas the bouton indicated by "G-" does not display immunoreactivity. The dendritic shafts (d) in the neuropil are non-immunoreactive for GABA. Scale bars = 100 μm (A); 20 μm (B); 0.5 μm (C, also applies to D).
Fig. 10. Photomicrographs (A,B) and electron micrographs (C,D) showing various features of labelling in the GPe after injection of BDA in the ventrolateral two-thirds of the GPi (see Fig. 2A,E). (A) Shows a higher power view of the cluster of retrogradely labelled cells in register with BDA-containing varicosities indicated by an arrow in Fig. 2E. (B) Example of a retrogradely labelled cell in the core of the cluster of labelling in the GPe. The dendritic shaft of the labelled cell is apposed by numerous BDA-containing varicosities. (C) Examples of BDA-containing type 3 terminals apposed to the surface of a dendritic shaft (d). This type of terminal represents the least frequent population (7%) of labelled boutons encountered in the GPe after BDA injections in the GPi. These boutons display the ultrastructural features of striatal terminals and are immunoreactive for GABA. Note that the density of gold particles associated with these terminals is significantly higher than that overlying another bouton (indicated by "G-”) in the neuropil. (D) An example of a BDA-labelled type 2 terminal that forms an asymmetric synapse (arrowhead) with a dendritic shaft (d). This type of terminal accounts for 82% of the labelled boutons found in the GPe after BDA injections in the GPi. In a GABA-immunostained section, these terminals are devoid of gold particles, which indicates that they are non-immunoreactive for GABA ("G-”). Note that other terminals in the neuropil are strongly immunoreactive for GABA ("G"). Scale bars = 0.25 mm (A); 20 μm (B); 0.5 μm (C, also applies to D).

Electron-microscopic analysis of these terminals revealed two major populations. One type possessed the characteristics of terminals derived from the globus pallidus and were presumably labelled by the anterograde transport of the tracer from the GPe to the GPi. The second class displayed the ultrastructural features of terminals derived from the STN; which implies that they were presumably labelled by retrograde transport of tracer from the GPe to the STN and then anterograde transport to collaterals in the GPi. This phenomenon has also been observed in the entopeduncular nucleus.
following tracer deposits in the globus pallidus of rats.

These findings, together with previous data obtained with other neuronal tracers in the basal ganglia and other brain regions (see above), lead us to conclude therefore, that the axons and terminals that become labelled in the STN after injections of BDA in the Gpi are the collaterals of GPe neurons that retrogradely transported the tracer and then distributed it throughout their axonal fields (Fig. 12A). Similarly, we conclude that the terminals labelled in the GPe after the tracer deposits in the Gpi are derived from neurons in the STN that retrogradely transported the tracer to their somata and then anterogradely transported it, via long collaterals, to the Gpe (Fig. 12A). Although the transport of BDA along the axon collaterals of retrogradely labelled cells could make the interpretation of the data obtained with this tracer difficult, our findings demonstrate that BDA also offers unique possibilities for studying the relationships between interconnected brain structures such as the pallidal complex and the STN. However, a prerequisite for such an application of BDA is a thorough knowledge of (i) the basic circuitry of the neuronal network under investigation, and (ii) the ultrastructural features and neurotransmitter content of the labelled terminals. It should be noted that, in addition to the main population of terminals in both the STN and GPe that were labelled following tracer deposits in the Gpi, a small proportion of labelled terminals in each structure displayed ultrastructural features different from the major group of labelled boutons. In the STN, a few labelled terminals (type 2 boutons) contained round synaptic vesicles and formed asymmetric axodendritic synapses. Based on these ultrastructural features, it is likely that these terminals arose from intrinsic axon collaterals of retrogradely labelled STN neurons, the existence of which has been previously demonstrated in the rat.37 In the GPe, some labelled terminals displayed the ultrastructural features of pallidal boutons (type 1 terminals). This population of boutons may have been anterogradely labelled from the Gpi19,30 or may arise from intrinsic axon collaterals of retrogradely labelled GPe neurons.39,67,68 A very small proportion of the labelled terminals in the GPe (type 3 boutons) contained ovoid synaptic vesicles, formed symmetric synapses and displayed GABA immunoreactivity. These characteristics are typical of those described on many occasions for the boutons of striatal projection neurons.6,8,12,57,60 Since it is known that a proportion of neurons in the striatum project to the Gpi and send axon collaterals to the Gpe,30,32,50 it is likely that the third type of terminals became labelled by retrograde transport from the Gpi and then anterograde transport, via axon collaterals, to the GPe.

**Neuronal network interconnecting the external pallidum and subthalamic nucleus with the internal pallidum**

The present findings demonstrate a highly ordered and specific relationship between neurons of the pallidal complex and neurons in the STN. The specificity of the relationship is depicted by the close overlap between fields of terminals and retrogradely labelled cells in the STN and GPe after injections of tracers in the Gpi. Thus, BDA injections in the Gpi led to retrograde labelling of populations of neurons in both the GPe and STN that were embedded in dense networks of terminals derived from the STN and GPe, respectively. This reciprocity also occurred at the synaptic level. Many of the neurons in the STN that were retrogradely labelled from the Gpi received synaptic input from terminals that were the collaterals of pallidal neurons that innervated the same region of the Gpi. Similarly, neurons in the Gpe that were retrogradely labelled from the Gpi received synaptic input from the axon collaterals of subthalamic neurons that innervated the common region of the Gpi.

The deposits of neuronal tracers in different regions of the internal and external segments of the globus pallidus and the subsequent anterograde and retrograde labelling that we observed demonstrate that the organizational principle of this neuronal network holds true for different functional regions of the GPe, STN and Gpi. Our data thus confirm and extend previous findings relating to the topographical organization of the subthalamo-pallidal and the pallidosubthalamic projections in primates11,65,66 and demonstrate that the functional specificity is maintained throughout the pallido-subthalamo-pallidal system. Thus, populations of neurons within sensorimotor, cognitive and limbic territories in the GPe are reciprocally connected with populations of neurons in the same functional territories of the STN and the neurons in each of these regions then, in turn, innervate the same functional territory of the Gpi.1,2,25,26,63,74,75

An additional interesting and important conclusion of the present study is that neurons in the STN possess axon collaterals that innervate both segments of the pallidal complex. Although previous retrograde labelling studies in monkeys have suggested that projections to the GPe or Gpi largely arise from different populations of neurons in the STN,11,50 these conclusions were based on the analysis of retrograde labelling in the STN after large injections of horseradish peroxidase11 or fluorescent dyes50 in one or the other pallidal segment. The fact that these injections were large and, as is now apparent, did not involve interconnected regions of the GPe and Gpi may explain the discrepancy between our findings and those obtained in these studies. However, our data do not necessarily indicate that all STN neurons project to both pallidal segments. After BDA
Organization of the pallido-subthalamo-pallidal system

Fig. 12. Schematic diagrams that show the possible sources of labelling in the pallidal complex and the STN following injections of BDA in the GPi (A) or GPe (B). The arrows indicate the directions that the tracer is expected to have been transported to produce the observed results. (A) After injections in GPi, BDA is transported retrogradely to GPe and then anterogradely, via axon collaterals, to the STN and GPi respectively. (B) After injections in the GPe, BDA is transported retrogradely to the STN and then anterogradely, via axon collaterals, to the GPi. BDA is also transported anterogradely to the GPi and STN.

Injections in either segment of the globus pallidus, a significant number of cells in register with the field of labelled terminals remained unlabelled in the STN. Although the projection site of these neurons was not determined, a possibility could be that these cells innervate exclusively the segment of the globus pallidus that was not injected with BDA. Of course, an alternative explanation is that these neurons project to other targets of the STN including the substantia nigra or the pedunculopontine nucleus. Intracellular filling of single STN neurons will resolve this issue. Similar findings were obtained in the GPe after BDA injections in the GPi. The fact that some GPe cells remained unlabelled in the core of the terminal field after injections in the GPi may indicate that not all GPe neurons project to both GPi and STN. These data are in keeping with those obtained in a recent intracellular study in the rat.

The organizational principle that we describe is not consistent with the current view (see Introduction) that the STN provides a widespread, “nonspecific” excitatory influence on the globus pallidus, rather, neurons in the STN are in a position to produce a very specific excitatory effect on small populations of related neurons in the GPe and GPi. It should be noted however, that following the deposits of tracer in the GPi occasional retrogradely labelled neurons were located outside of the region of axon and terminal labelling in the GPe and STN. It is unclear at present whether this observation simply reflects the fact that the tracer deposits extended in the dorsolateral plane, thus contaminating neighbouring functional regions or whether the tracer was taken up by distal segments of damaged axons that extend out of the injection site. Of course, another alternative is that the organizational principle that we have...
identified overlies an additional, albeit less prominent, system in which there is not a correspondence between functionally related neurons. The only way to fully solve this issue is to fill intracellularly single STN cells and trace their full axonal arborization in the GPi and GPe. However, because of the location and size of the STN, intracellular filling of cells in this nucleus is a very difficult task in the monkey brain.

Functional considerations

The new organizational principle of the interconnections between the GPe, STN and GPi that we have described has important implications for the understanding of the neuronal mechanisms by which the so-called indirect pathways²⁻¹⁴ are involved in the control of motor commands in primates and the interpretation of electrophysiological data. Firstly, it is clear that axons of subthalamic neurons are not widely spread in the GPe or GPi. In contrast, specific groups of functionally related STN neurons send projections that terminate with a high degree of specificity in restricted regions of the GPe and GPi. Thus, in the analysis of synaptic interactions between neurons in the pallidal complex and STN it is critical to select the interconnected functionally related regions of each structure. The failure to detect synchronization between pairs of STN and pallidal neurons in electrophysiological analyses in primates⁴ may simply reflect the failure to locate interconnected neurons rather than a diffuse connection between the STN and the pallidum. In keeping with this possibility, recent data showed that the activity of pallidal neurons in normal monkeys is non-synchronized, which indicates that the spiking activity of pallidal cells is not driven by common inputs.⁶ Secondly, the network that we describe provides an anatomical substrate for the complex excitatory and inhibitory responses that occur in output neurons of the basal ganglia following activation of the indirect pathway.²² Thus, although it is evident that a major excitatory "driving force" of GPi cells are neurons in the STN, it is likely that the same pool of neurons in the STN inhibit indirectly, the same neurons in the GPi by activation of the neurons in the GPe that project to the common region of the GPi. The neuronal mechanisms by which the opposite effects generated by activation of the STN–GPi and the STN–GPe–GPi projections are integrated at the level of single GPi cells remains to be established. The complex pattern of increased and decreased activity of the basal ganglia output neurons during a motor act¹⁰,²²,⁴²,⁴⁴ probably relies on precise temporal interactions between the activities of the inhibitory GPe and striatal afferents with that of the excitatory subthalamic input to single GPi neurons. Future electrophysiological data relating to the activity of interconnected pairs of GPe and GPi neurons following stimulation of the STN are essential to further our understanding of these issues.

CONCLUSIONS

The results of our study raise several important issues concerning the circuitry connecting the two pallidal segments and the subthalamic nucleus as well as the mechanisms by which information is processed in the basal ganglia. Firstly, the connections between the two segments of the pallidum and the STN show a high degree of specificity such that small groups of neurons in the GPe and STN innervate common regions in the GPi and are reciprocally connected at the synaptic level. Secondly, the topographical distribution of the interconnected neurons respects the known functional organization of the two segments of the globus pallidus and the STN. Thirdly, the findings imply that individual neurons in the GPe project via collaterals to both the GPi and STN and, similarly, that neurons in the STN project to both the GPi and GPe.

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