Are there three subdivisions in the primate subthalamic nucleus?

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Are there three subdivisions in the primate subthalamic nucleus?

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The subthalamic nucleus (STN) is a small but vitally important structure in the basal ganglia. Because of its central role in motor control, the STN is the target of deep-brain stimulation that alleviates severe motor symptoms in many levodopa-resistant patients with Parkinson’s disease. Unfortunately, deep-brain stimulation of the STN also comes at a cost. In particular, some patients with Parkinson’s disease experience a decline in cognitive functioning, whereas others may become depressed, experience hypersexuality, hypomania, and some even commit suicide (Burkhard et al., 2004; Temel et al., 2005).

These undesirable and perhaps unexpected side-effects of deep-brain stimulation suggest that the STN regulates not only motor behavior but also cognitive and emotional processes, a suggestion consistent with one of the most influential theories of the STN (Alexander and Crutcher, 1990). How does the STN carry out these separate duties? A simple and widely held assumption is that the STN is comprised of three anatomically distinct subparts, each of which is selectively connected to cognitive, limbic, and motor areas. An assumption, that seems to be supported by recent electrophysiological recordings in humans (Mallet et al., 2007; Greenhouse et al., 2011). To underscore the popularity of this assumption, we reviewed all 16 STN review articles published since 1985 (DeLong et al., 1985; Alexander et al., 1986, 1990; Alexander and Crutcher, 1990; Parent, 1990; Parent and Hazrati, 1993, 1995a,b; Joel and Weiner, 1997; Nakano, 2000; Rodriguez-Oroz et al., 2001, 2009; Hamani et al., 2004; Temel et al., 2005; Obeso et al., 2008; Krack et al., 2010) and found that 12 (75%) of these articles suggested that the STN is a tripartite system (DeLong et al., 1985; Parent, 1990; Parent and Hazrati, 1995a,b; Joel and Weiner, 1997; Nakano, 2000; Rodriguez-Oroz et al., 2001, 2009; Hamani et al., 2004; Temel et al., 2005; Obeso et al., 2008; Krack et al., 2010).

INTRODUCTION

The subthalamic nucleus (STN) is a small but vitally important structure in the basal ganglia. Because of its central role in motor control, the STN is the target of deep-brain stimulation that alleviates severe motor symptoms in many levodopa-resistant patients with Parkinson’s disease. Unfortunately, deep-brain stimulation of the STN also comes at a cost. In particular, some patients with Parkinson’s disease experience a decline in cognitive functioning, whereas others may become depressed, experience hypersexuality, hypomania, and some even commit suicide (Burkhard et al., 2004; Temel et al., 2005).

These undesirable and perhaps unexpected side-effects of deep-brain stimulation suggest that the STN regulates not only motor behavior but also cognitive and emotional processes, a suggestion consistent with one of the most influential theories of the STN (Alexander and Crutcher, 1990). How does the STN carry out these separate duties? A simple and widely held assumption is that the STN is comprised of three anatomically distinct subparts, each of which is selectively connected to cognitive, limbic, and motor areas. An assumption, that seems to be supported by recent electrophysiological recordings in humans (Mallet et al., 2007; Greenhouse et al., 2011). To underscore the popularity of this assumption, we reviewed all 16 STN review articles published since 1985 (DeLong et al., 1985; Alexander et al., 1986, 1990; Alexander and Crutcher, 1990; Parent, 1990; Parent and Hazrati, 1993, 1995a,b; Joel and Weiner, 1997; Nakano, 2000; Rodriguez-Oroz et al., 2001, 2009; Hamani et al., 2004; Temel et al., 2005; Obeso et al., 2008; Rodriguez-Oroz et al., 2009; Krack et al., 2010) and found that 12 (75%) of these articles suggested that the STN is a tripartite system (DeLong et al., 1985; Parent, 1990; Parent and Hazrati, 1995a,b; Joel and Weiner, 1997; Nakano, 2000; Rodriguez-Oroz et al., 2001, 2009; Hamani et al., 2004; Temel et al., 2005; Obeso et al., 2008; Krack et al., 2010).

However, the assumption of a tripartite STN was at odds with our initial impression of the empirical literature. To test the empirical support for a tripartite STN, we therefore, conducted an independent and comprehensive literature review. In the first section below we summarize the results on the number of subdivisions in the STN, in the second section we also summarize the results on the presumed anatomical connectivity and localization of these subdivisions.

NUMBER OF SUBDIVISIONS IN THE STN

Following an extensive literature review we initially identified 43 studies, published between 1925 and 2010, that investigated anatomical connections via the use of tracers or lesions in the STN for human and nonhuman primates or used cytoarchitectonic studies of the STN for human and nonhuman primates (Foix and Nicolesco, 1925; Kodama, 1928; Whittier and Mettler, 1949; Nauta and Mehler, 1966; Carpenter and Strominger, 1967; Fussenich, 1967; Carpenter et al., 1968, 1981a,b; Petras, 1968; Carpenter and Peter, 1972; Kim et al., 1976; Kuzemensky, 1976; Rafols and Fox, 1976; Monakow et al., 1978; Nauta and Cole, 1978; DeVito et al., 1980; DeVito and Anderson, 1982; Parent et al., 1984, 1989; Smith and Parent, 1986; Parent and Smith, 1987; Mizuno et al., 1988; Nakano et al., 1990, 1994; Hazrati and Parent, 1991, 1992a,b; Sadikot et al., 1992; Lavoie and Parent, 1994; Nambu et al., 1996, 1997; Shink et al., 1996; Takada et al., 1996, 2001; Inase et al., 1999; Sato et al., 2000; Karachi et al., 2004; Kelly and Strick, 2004; Miyachi et al., 2006; Bostan et al., 2010; Rico et al., 2010).
With respect to the animal studies, we included studies only on nonhuman primates: just as in humans, the STN in nonhuman primates is a closed nucleus (i.e., dendrites are restricted to the nucleus), in contrast for example to the rat STN (Smith et al., 1990; Marani et al., 2008). With respect to the human studies, we chose only to include studies using cytoarchitectonics since this is still the golden standard in anatomy (Amunts et al., 2007) and seems to be more sensitive than cytochemical stainings in detecting different borders generally (Uylings et al., 2010; van de Werd et al., 2010).

From the initial group of 43 studies, we focused on a subset of 33 studies that explicitly consider the number of subdivisions in the STN (Foix and Nicolesco, 1925; Kodama, 1928; Whittier and Mettler, 1949; Nauta and Mehler, 1966; Fussenich, 1967; Kim et al., 1976; Kuzemenský, 1976; Rafols and Fox, 1976; Monakow et al., 1978; Nauta and Cole, 1978; Carpenter et al., 1981a,b; DeVito and Anderson, 1982; Parent et al., 1984, 1989; Smith and Parent, 1986; Parent and Smith, 1987; Nakano et al., 1990; Smith et al., 1990; Sadikot et al., 1992; Lavoie and Parent, 1994; Nambu et al., 1996, 1997; Shink et al., 1996; Inase et al., 1999; Sato et al., 2000; Takada et al., 2001; Karachi et al., 2004; Kelly and Strick, 2004; Miyachi et al., 2006; Bostan et al., 2010; Rico et al., 2010). These 33 studies were assigned to one of three categories, (A) 25 studies used anterograde and retrograde tracers to investigate possible STN subdivisions and their anatomical location in old-world and new-world monkeys (Carpenter et al., 1968; Kim et al., 1976; Monakow et al., 1978; Nauta and Cole, 1978; Carpenter et al., 1981a,b; DeVito and Anderson, 1982; Parent et al., 1984, 1989; Smith and Parent, 1986; Parent and Smith, 1987; Nakano et al., 1990; Smith et al., 1990; Sadikot et al., 1992; Lavoie and Parent, 1994; Nambu et al., 1996, 1997; Shink et al., 1996; Inase et al., 1999; Sato et al., 2000; Takada et al., 2001; Karachi et al., 2004; Kelly and Strick, 2004; Miyachi et al., 2006; Bostan et al., 2010; Rico et al., 2010), (B) two studies used electrical lesions in old-world monkeys (Nauta and Mehler, 1966; Carpenter et al., 1968), (C) six studies used classical cytoarchitectonics to study STN subdivisions by examining STN cell type, cell size, and cell density in human and nonhuman primates (Foix and Nicolesco, 1925; Kodama, 1928; Whittier and Mettler, 1949; Fussenich, 1967; Kuzemenský, 1976; Rafols and Fox, 1976). For each of these 33 studies, we summarized the data as supporting none, two, three, or four subdivisions in the STN. This classification reflected the opinion of the original authors of the studies we reviewed, and we did not attempt to reinterpret data or alter their conclusions in any way.

Table 1 summarizes the results. Overall, more than half of the studies we reviewed concluded that the STN has two subdivisions (61%), 21% concluded that the STN has no subdivisions, 12% concluded that the STN has three subdivisions, and 6% concluded that the STN has four subdivisions. These results contrast sharply with the popular academic opinion that the STN is a tripartite system. In addition, although the data most support the hypothesis that the STN has two subdivisions, the extent of this preference (i.e., 61% vs. 39% based on 33 studies) is not conclusive and the results appear to vary substantially from one study to the next.

The variability across studies may be partly due to methodological differences. In particular, it could be that results vary as a result of the specific tracer injection sites. It is also possible, however, that the general summary of results in Table 1 hides additional variability due to more detailed information regarding location and connectivity. For example, when researcher X argues that the STN has two subdivisions, one lateral and one medial, this claim is still inconsistent with that of researcher Y, who may also argue that the STN has two subdivisions, but believes that one is dorsal and the other ventral. The next section examines the connection profiles and STN subdivisions in relative detail.

Table 1 | Summary of 33 studies that examined the number of subdivisions in the STN for human and nonhuman primates.

<table>
<thead>
<tr>
<th>NUMBER OF STN SUBDIVISIONS</th>
<th>Tracer studies</th>
<th>Lesion studies</th>
<th>Cytoarchitectonic studies</th>
<th>All studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>5 (all nh)</td>
<td>1 (nh)</td>
<td>1 (h)</td>
<td>7 (1 h; 6 nh)</td>
</tr>
<tr>
<td>Two</td>
<td>15 (all nh)</td>
<td>0</td>
<td>5 (3 h; 2 nh)</td>
<td>20 (3 h; 17 nh)</td>
</tr>
<tr>
<td>Three</td>
<td>3 (all nh)</td>
<td>1 (nh)</td>
<td>0</td>
<td>4 (all nh)</td>
</tr>
<tr>
<td>Four</td>
<td>2 (all nh)</td>
<td>0</td>
<td>0</td>
<td>2 (all nh)</td>
</tr>
</tbody>
</table>

Between brackets indicates if the study used nonhuman primate (nh) or human (h) tissue.

ANATOMICAL CONNECTIVITY AND LOCALIZATION OF SUBDIVISIONS IN THE STN

We separately analyzed the results from each of 27 studies that used either tracing or electrical lesions to construct connection profiles and localize subdivisions in the STN. See mckitchen.files.wordpress.com/2010/01/supplementary-table-stn.pdf for a detailed overview for the methods, the number of animals and species used in the studies included in this review.

The results of these studies are summarized in Figure 1 and ordered by the number of subdivisions reported and the localization of the subdivisions within the STN. The subdivisions are represented on three schematic slices of the STN in rostrocaudal direction. This was done because individual studies could not be registered in one common anatomical space. Note that subdivisions were determined based both on statements derived from the individual studies and on figures contained in the articles.

To exemplify the kind of information visualized in Figure 1, consider the work by DeVito and Anderson (1982), who state: “Silver grains were distributed diffusely over the STN, with no apparent pattern to reflect an underlying anatomical organization (p. 110);” as another example, Karachi et al. (2004) mention: “In summary, the pallidosubthalamic projection appears to be topographically arranged, with the sensorimotor part of the STN occupying its dorsolateral half, and the limbic part being restricted to its most anterior and medioventral portion (p. 178);” finally, Miyachi et al. (2006) provide a figure (i.e., their Figure 6B) that presents the layout of subdivisions (see Table 2 for all the statements used to visualize all the included studies).

One might argue that a temporal bias may be present, i.e., a lower number of subdivisions could have been found due to older methods and that more recent studies using advanced methodologies converge to a specific number of subdivisions. Figure 2A
FIGURE 1 | Continued
FIGURE 1 | Connection profile and subdivisions of the STN show highly variable results. (Top of figure) Color-coded lesion, anterograde, and retrograde tracer injection sites including cortical, subcortical, and brainstem regions. In the middle, the anatomical orientation of the STN template is displayed. Three schematic slices from caudal to rostral are shown. Anatomical abbreviations: BA, Brodmann Area; CMAd, Cingulate Motor Area dorsal; CMAr, Cingulate Motor Area rostral; CMAv, Cingulate Motor Area ventral; GPe, Globus Pallidus externa segment; GPi, Globus Pallidus interna segment; HVIIB, Gracil Lobule; M1, Primary Motor cortex, PMd, Pre-Motor dorsal; PMv, Pre-Motor ventral; PPN, Pedunculopontine Nucleus; Pre-SMA, Pre-Supplementary Motor Area; S1, Primary Sensory cortex; SMA, Supplementary Motor Area; SN, Substantia Nigra compacta; SNr, Substantia Nigra reticulata; VA, Ventral Anterior Nucleus; VL, Ventral Lateral Nucleus. (A–F) Six studies arguing for zero subdivisions. The author and publication year is shown in the left upper corner. (G–U) Fifteen studies arguing for two subdivisions. The author and publication year is shown in the left upper corner. (V–Y) Four studies arguing for three subdivisions. The author and publication year is shown in the left upper corner. (Z–AA) Two studies arguing for four subdivisions. The author and publication year is shown in the left upper corner. Black arrows denote the direction of connection. The red arrow denotes the location of a lesion. Localization of subdivisions within the STN are color coded according to the injection or lesion site.
Table 2 | The exact statement or figure number referring to the number of subdivisions derived from all tracings and lesions studies displayed in Figure 1.

<table>
<thead>
<tr>
<th>Subdivision</th>
<th>Reference</th>
<th>Statement/Description</th>
</tr>
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<tbody>
<tr>
<td>ZERO SUBDIVISIONS</td>
<td>Nauta and Mehler (1966)</td>
<td>p. 32: “The material available was, however, too restricted to afford an adequate picture of further details of the topological organization, particularly because no lesions were placed in the far rostral and caudal quarters of the globus pallidus.”</td>
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<td></td>
<td>Kim et al. (1976)</td>
<td>p. 273: “Radioactive labeling was especially prominent in ventral regions of the lateral part of the nucleus. Only one section through caudal parts of the subthalamic nucleus revealed label in terminal distributed to all parts of the nucleus.”</td>
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<td></td>
<td>DeVito and Anderson (1982)</td>
<td>p. 110: “Silver grains were distributed diffusely over the subthalamic nucleus, with no apparent pattern to reflect an underlying anatomical organization.”</td>
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<td>Lavoie and Parent (1994)</td>
<td>p. 227: “In the squirrel monkey, afferent fibers from the PPN are uniformly distributed in the subthalamic nucleus and these fibers do not form close pericellular contacts with subthalamic neurons.”</td>
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<td></td>
<td>Sato et al. (2000)</td>
<td>p. 145: “The plotting of each type of STN projection neurons of a series of equally spaced (280 µm apart) parasagittal sections (Figure 6) did not reveal any correlation between the precise location of the labeled cells within the nucleus and their axonal branching pattern.”</td>
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<td>Rico et al. (2010)</td>
<td>p. 387: “Most importantly, the delivery of CTB into the VA/VL thalamic nuclei led to the appearance of a moderate number of retrograde-labeled neurons in the ipsilateral STN nucleus. CTB-labeled neurons were distributed throughout the STN in all 6 monkeys single-injected with CTB, although labeled neurons appeared to be discretely and preferentially located within medial STN territories.”</td>
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<tr>
<td>TWO SUBDIVISIONS</td>
<td>Monakow et al. (1978)</td>
<td>p. 402: &quot;While the precentral connections occupy the lateral and dorsal moiety of N. Subthalamicus, it was found that the remaining frontal lobe regions may project more to the medial and ventral districts.”</td>
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<td></td>
<td>Nauta and Cole (1978)</td>
<td>p. 3: “A comparison of the two cases MS-7 and MS-8 suggest a topographic organization in which the lateral part of the subthalamic nucleus (injected in case MS-7) projects to the main sector of the pallidal complex while the rostro-medial part of the nucleus (injected in case MS-8) projects to the rostral and medial parts of the pallidal complex.”</td>
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<td>Carpenter et al. (1981a,b)</td>
<td>p. 22: “(1) cells in the medial half of the middle third of the STN projected to the rostral division of the LPS and (2) cells in central regions of the rostral two-thirds of the STN projected to the central division of the LPS.”</td>
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<td></td>
<td>Parent et al. (1984)</td>
<td>p. 389: “Also noteworthy is the situation in the subthalamic nucleus where GP-labeled cells occupy the entire dorsolateral two-thirds of the structure whereas SN-labeled cells are mostly confined to the ventromedial third, with again only a small number of double-labeled neurons.”</td>
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<td>Smith and Parent (1986)</td>
<td>p. 357: “The subthalamic nucleus also contains numerous positive cells which abound in the dorsolateral two-third portion of the structure on the putamen-injected side, and in the ventromedial third of the nucleus on the caudate-injected side.”</td>
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<td></td>
<td>Parent and Smith (1987)</td>
<td>p. 307: “(1) a large (80%) dorsolateral ‘sensorimotor’ zone where most neurons project to the lenticular nucleus and terminate in the globus pallidus and/or the putamen, (2) a small (20%) ventromedial ‘associative’ zone whose neurons give rise to either ascending projections to the caudate nucleus or descending projections to the substantia nigra, and (3) a smaller (10%) overlapping zone whose neurons send axon collaterals to both the lenticular nucleus and the substantia nigra.”</td>
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<td>Nakano et al. (1990)</td>
<td>p. 65: “The medial (horizontally shaded area) or lateral (solid star area) part of the STN sends projection fibers to the caudate nucleus (horizontally shaded area in CN) or the putamen (solid star area in Putamen), respectively.”</td>
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<td>Smith et al. (1990)</td>
<td>p. 321: “Instead, much like the striatum, it appears to be composed of separate subsystems that originate from different regions of the nucleus and modulate the various target structures.”</td>
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<td>Sadikot et al. (1992)</td>
<td>p. 149: “The present study shows that CM projects lightly to the dorsal lateral part of the subthalamic nucleus, whereas PF projects mainly to its medial and rostral parts.”</td>
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<td></td>
<td>Shink et al. (1996)</td>
<td>See Figure 13 for the layout.</td>
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<td></td>
<td>Nambu et al. (1997)</td>
<td>p. 15: “The terminal zones from the PMd and PMv were located mostly in the medial aspect of the STN, whereas the terminal zone from the M1 was in its lateral aspect.”</td>
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<td>Inase et al. (1999)</td>
<td>See Figure 4 for the layout.</td>
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<td></td>
<td>Takada et al. (2001)</td>
<td>See Figure 13 for the layout.</td>
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(Continued)
shows that this is not the case. Another possible bias is the species, i.e., new- or old-world monkey, and the type of tracer used. Again, no clear evidence is found for such a bias (Figure 2B).

Two important conclusions can be drawn from this overview: (1) claims about the number of subdivisions in the STN do not depend on injection site in a clear and systematic way; (2) claims about the localization of subdivisions within the STN are highly variable. The variability of the results is clear from a visual inspection of Figure 1, but it is also apparent from other considerations; for instance, studies using the same subcortical injection sites such as the Globus Pallidus (Parent et al., 1989; Karachi et al., 2004) yield different localizations of subdivisions within the STN.

Based on this overview, there is a discrepancy between the empirical work and the general conclusions of the reviews published since 1985, as 75% of those reviews argue that there are three subdivisions, 6% conclude that there are four subdivisions, 13% state that there are five subdivisions, and finally 6% conclude that there are subdivisions in the STN but do not specify the number. Notably, during the past 26 years, no review paper has been published arguing in favor of none or two subdivisions in the STN despite the empirical studies that show, if at all, more evidence for two than three subdivisions.

It is also noteworthy, that the misconception of a tripartite STN could have been caused by misquotations. For instance, two review articles (Alexander et al., 1986, 1990) argue for five subdivisions, however, they have frequently been cited by others in favor of three subdivisions (e.g., Parent and Hazrati, 1995a,b). Such quotations are prone to yield a scientific consensus despite a yet to be answered question about the number of subdivisions in the STN.

**CLASSICAL CYTOARCHITECTONIC STUDIES IN HUMAN AND NON-HUMAN PRIMATES**

The classical cytoarchitectonic studies (i.e., studies depending on measures such as cell type, cell size, and cell density) are also inconsistent with the tripartite STN (Figure 3). We found six articles, including four with human specimens and two with old-world monkeys, published between 1925 and 1976. Five of these six studies reported two subdivisions in the STN (Foix and Nicolesco, 1925; Kodama, 1928; Whittier and Mettler, 1949; Fussenich, 1967; Rafols and Fox, 1976), clearly arguing for a heterogeneous neuronal distribution and one study with a human specimen arguing for no subdivisions (Kuzemenský, 1976). Although these results are less variable than the results of the tracer studies (80% of the cytoarchitectonic studies report the same number of subdivisions compared to 55% in the tracer and lesion studies), considerable variability regarding the localization of the subdivisions remains, as only three cytoarchitectonic studies show a comparable localization of the subdivisions within the STN.

**CONCLUSION**

Our review of the literature sharply contrasts with the popular belief that the STN is a tripartite system (see Figure 1). In addition, our review shows that the variability across studies is surprisingly large—studies that support the same number of subdivisions almost always disagree on the localization of these

<table>
<thead>
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<th>Table 2</th>
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<td>Kelly and Strick (2004)</td>
<td>See Figure 3 for the layout.</td>
</tr>
</tbody>
</table>

**THREE SUBDIVISIONS**

| Carpenter et al. (1968) | p. 558: “(a) the most medial and lateral regions of the lateral pallidal segment project fibers respectively to medial and lateral parts of the subthalamic nucleus, and (b) the medial segment of the globus pallidus projects a small number of fibers to the subthalamic nucleus which terminate in regions caudal and medial to those that receive fibers from the lateral pallidal segment.” |
| Parent et al. (1989) | p. 143: “in the rostral third of the subthalamic nucleus, GPe labeled cells were slightly more numerous than GPi-positive neurons and each population occupied a distinct territory, GPe-labeled cells being confined to the lateral half and GPi-positive neurons to the medial half of the nucleus. In the middle and caudal thirds of the subthalamic nucleus, the two cell populations remained well segregated but appeared progressively displaced medially leaving the lateral third of the nucleus mostly devoid of labeled cells.” |
| Nambu et al. (1996) | p. 2680: “Each of the two mediolaterally separated portions of the STN is characterized with somatotopically arranged hyperdirect cortical inputs form the M1 and the SMA. The first set of body parts representations is transformed mainly from the M1 to the lateral STN, whereas the second set is transformed primarily from the SMA to the medial STN.” |
| Karachi et al. (2004) | p. 178: “In summary, the pallidosubthalamic projection appears to be topographically arranged, with the sensori-motor part of the STN occupying its dorsolateral half, and the limbic part being restricted to its most anterior and medioventral portion.” |

**FOUR SUBDIVISIONS**

| Carpenter et al. (1981a,b) | See Figure 8 and figure text caption for layout. |
| Miyachi et al. (2006) | See Figure 6B for the layout. |

All studies are based on non-human primates.
subdivisions. A finding that cannot be explained by a potential bias regarding the publication year of the studies, methodological differences between studies, and the species under investigation (see Figure 2).

In the face of these results, it is difficult to explain why the belief in a tripartite STN continues to be as popular as it is. Moreover, Alexander et al. (1990) considered 5 parallel loop circuits in the basal ganglia and the present view considers 10 or more parallel circuits (e.g., Middleton and Strick, 2001) involving the STN. Therefore, one might speculate that the STN consist of more than three subdivisions. In sum, our review suggests that to better understand the anatomical structure of the STN further systematic studies are needed. We hope that future post-mortem and in vivo studies using ultra-high resolution magnetic resonance imaging and histology will provide more definitive evidence regarding STN subdivisions and will help answer some important questions (Cho et al., 2010; Forstmann et al., 2010; Turner, 2011). Based on the empirical data that are presently available, firm conclusions about the number and localization of the STN are presently not warranted.

**OUTSTANDING QUESTIONS**

- Are there subdivisions in the STN? If so, how many?
- What are the clinical implications of subdivisions in the STN?
- To what extent are nonhuman primate STN subdivisions comparable to in vivo human STN subdivisions?
- Are there individual differences in STN structural connectivity that predict differences in cognition?
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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.