Abstract: Deep Brain Stimulation (DBS) is an invasive therapeutic method involving the implantation of electrodes and the electrical stimulation of specific areas of the brain to modulate their activity. DBS brings therapeutic benefits, but can also have adverse side effects. Recently, neuroethicists have recognized that DBS poses a threat to the very fabric of human existence, namely, to the selves of patients. This article provides a review of the neuroethical literature examining this issue, and identifies the crucial dimensions related to the self which DBS may endanger—personal identity, authenticity, and autonomy. The most influential theories accounting for these dimensions are analyzed herein, and it is argued that most of these theories require further refinement. This paper also demonstrates the interrelation between personal identity, authenticity, and autonomy, and concludes that one can only fully understand the impact of DBS on the self when all of these factors are taken into account.

Keywords: self, personal identity, authenticity, autonomy, deep brain stimulation (DBS), neuroethics.

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Therapeutic potential

Deep Brain Stimulation (DBS) is an invasive therapeutic method involving the implantation of electrodes and the electrical stimulation of specific regions of the brain. Depending on the disease and treatment strategy, different structures are targeted, for example, the subthalamic nucleus (STN), the globus pallidus internus (GPi), or the nucleus accumbens (NAc). The exact action mechanism of DBS has not yet been confirmed. The proposed models, however, can be divided into four categories. Models of the first category reduce the effect of DBS to the inhibition of the neuronal activity of the targeted areas for stimulation; the second to the excitation of these areas; models of the third category postulate that its action is due to a combination of these mechanisms;
and the fourth considers DBS to be interrupting pathological oscillations to regain the synchronized, rhythmic activity of the brain waves.4

The lack of consensus among researchers about these mechanisms has not prevented the systematic use of this technology for medical purposes.5 After the introduction of the first commercial system for clinical use in 1997,6 DBS proved its therapeutic potential in a number of intractable neurological and psychiatric disorders for which previous treatments (e.g. pharmacotherapy, cognitive behavioral therapy) had not provided satisfactory results, such as Parkinson’s disease (PD),7 dystonia,8 essential tremor,9 and obsessive–compulsive disorder (OCD).10 This prompted the American Food and Drug Administration (FDA) to approve DBS as a treatment for these ailments. Thus, DBS should be considered a treatment of last resort for patients with no other viable treatment options.11 The therapeutic range of DBS is constantly expanding, with experimental studies conducted in the treatment of epilepsy,12 Tourette’s syndrome,13 treatment–resistant depression,14 Alzheimer’s disease and other forms of dementia,15 anorexia,16 obesity,17 disorders of consciousness,18 chronic pain,19 Huntington’s disease,20 addictions,21 aggression,22 and schizophrenia.23 In Poland, DBS is mainly used as a treatment for PD, as well as for dystonia and essential tremor.24 Some centers in Poland are also working on the use of DBS for psychiatric disorders: the 10th Military Research Hospital and Polyclinic IPHC in Bydgoszcz, the University Hospital in Wroclaw, and the Wroclaw Medical University have experience in this field. Currently, a multicenter study is planned on the use of DBS in patients with treatment–resistant depression.25

Several companies that produce DBS equipment are operating in the market. In 2017, the leading manufacturer, Medtronic Activia, announced that 150,000 patients had already used their systems for DBS therapies, and this number increases annually by

4 Chiken, Nambu (2015).
7 Hickey, Stacy (2016).
8 Vidailhet, Jutras, Grabli et al. (2013).
9 Ostrem, Racine, Glass et al. (2011).
10 Hamani, Pilitis, Rughani et al. (2014).
11 Many thanks to the anonymous reviewer for pushing me to sufficiently emphasize this crucial feature of DBS therapy as it has important consequences for the neuroethical considerations in this paper.
12 Fisher, Salanova, Witt et al. (2010).
14 Bewernick, Kayser, Sturm et al. (2012).
15 Laxton, Lozano (2013).
17 Whiting, Tomycz, Bailes et al. (2013).
18 Schiff, Giacino, Kalmar et al. (2007).
21 Kuhn, Bührle, Lenartz et al. (2013).
22 Franzini, Broggi, Cordella et al. (2013).
23 Corripio, Roldán, Sarró et al. (2020).
24 Beszlej, Siwicki, Fila-Witecka et al. (2019).
25 Beszlej, Wieczorek, Kobyłko et al. (2019).
approximately 10,000. Currently, the number of patients considerably exceeds 160,000 and is estimated at 175,000. These numbers show that DBS plays an important role in the lives of many people, and as research progresses on the disorders that are currently under study, this impact will include further groups of patients.

**DBS technologies**

DBS systems consist of one or more electrodes inserted into the nervous tissue of the deep brain, connected via a subcutaneous extension wire to a stimulator (“pacemaker”) implanted in the left subclavian area. In order to stimulate the electrodes, the pacemaker generates rectangular electrical pulses. Typical DBS parameter settings for movement disorders range from 2 to 4 V amplitude, with a 60–450 μs pulse width at a frequency of 130–185 Hz. The implantation procedure marks the first stage of the therapy; followed by the device programming process. Until recently, this was performed by painstakingly adjusting the parameters of the stimulation by trial and error. Newer techniques facilitate the programming process by enabling the more optimal adjustment of parameters based on:

1. determining the precise location of the electrodes by Magnetic Resonance Imaging (MRI) or pre–operative MRI in combination with post–operative tomography imaging;
2. determining the desired target of stimulation with the help of previous clinical experience and estimation of current spread within the brain tissue.

This approach is referred to as “open loop” DBS (OL–DBS), and although it is currently the most common technique, it has been recognized as problematic for various reasons. First, it results in a mismatch between the time course of the disease and the timing of adjustments, since clinicians can only adjust the stimulation parameters during medical appointments with patients. Second, it is based on a subjective assessment of the parameters that are beneficial in each treatment. Third, it applies a stimulus pattern established during the programming process throughout the entire lifetime of the device. This operating mode typically results in the battery discharging within 2–5 years; it must then be surgically replaced, which has additional costs, stress, and risks associated with the procedure. Finally, the constant activity of OL–DBS can lead to undesirable permanent changes in neurotransmission in the long run, as well as non–stimulation–dependent neural reorganization.

To resolve these problems, researchers have investigated and developed a new generation of DBS systems, commonly referred to as “closed loop” DBS (CL–DBS). The CL–DBS paradigm is defined by the dynamic adjustment of parameters based on

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26 Lozano, Lipsman, Bergman et al. (2019).
27 Medtronic (2021).
30 Frankemolle, Wu, Noecker et al. (2010).
31 Ondo, Meilak, Vuong (2007).
33 Glannon (2016).
34 Ruge, Cif, Limousin et al. (2011).
the patient’s current clinical condition assessed by measuring brain activity. CL–DBS developed from the field of brain–computer interfaces (BCIs)—algorithms intended to identify the intentions of a person from brain activity to provide control over prosthetic/assistive devices. Implementing BCIs in CL–DBS systems allows classification of brain activity as “healthy” or “adverse.” Thanks to this procedure, CL–DBS can deliver and adjust stimulation in real time according to the particular needs of the patient. For example, when employed in the treatment of epilepsy, it detects early indicators of seizures and applies current to prevent the impending attacks. Compared to OL–DBS, CL–DBS systems result in better adaptation to specific disease dynamics, less invasiveness in neurotransmission, and reduced battery consumption. The CL–DBS paradigm has shown promising results for both neurological diseases like epilepsy or PD, as well as psychiatric disorders such as OCD and major depression.

The emerging trend in DBS–based treatments is not only meant to facilitate the automatic adjustment of the stimulation in response to the abnormal neuronal activation associated with a given disease, but also to provide patients with the kind of active control which allows them to respond to the symptoms and autonomously address therapeutic goals. Systems that aim to facilitate this differ operationally from CL–DBS systems, as they are closed–loop advisory brain devices or volitional CL–DBS (VCL–DBS). In short, when the device detects unwanted neuronal activation, it gives a patient a visual or auditory signal. On this basis, the patient can decide whether stimulation should begin, be increased, and/or rejected. Hence, VCL–DBS enables the patients to stay “in the decisional therapeutic loop”. In that regard, it offers them additional opportunities to choose between experiencing symptoms and suffering side effects. However, it produces other risks which will be considered in the following sections.

**Reports of DBS–induced changes to the self of patients**

As mentioned above, DBS is a last resort treatment for intractable diseases and it often provides beneficial therapeutic effects. Therefore, patients suffering from such diseases who are not candidates for DBS or do not have access to it will continue to suffer from untreated symptoms of their disorder. But DBS itself can also lead to unwanted side effects. Thus, there are adverse effects which may sometimes constitute inevitable tradeoffs for DBS when it is used to control disease-related symptoms. Therefore, one of the crucial questions in the neuroethics of DBS is how we should evaluate the comparative benefits and risks associated with this treatment.
risks and benefits of taking DBS therapy (as well as not receiving it at all). Two main lines of risk analysis associated with DBS have appeared in the neuroethical literature: (more) traditional and non–traditional. The former, more traditional forms of threats resulting from DBS therapy include effects in several domains: physical, cognitive, affective, and communication disorders. Moreover, DBS may also lead to unrealistic expectations for treatment, suicidal ideation, issues associated with informed consent, privacy and data security, and have regulatory implications.

DBS also appears to constitute threats that have not traditionally been discussed in the literature concerning the implications of neurotechnologies. They have become the subject of lively interest in recent years, when many neuroethicists have expressed concerns over the potential negative impact of DBS on the selves of patients. These concerns were initially raised by individual reports of patients treated with DBS.

One such example was the case of a man suffering from Tourette’s syndrome. As soon as the amplitude of DBS increased, he rolled into the corner, covered his face, and repeated in a high, broken voice that he was innocent. The sentences he spoke were grammatically incorrect. When the medical staff tried to help him, he would not allow it, and kept shouting that he was afraid of being locked in the basement. However, when the amplitude of the stimulation was reduced, the patient returned to his previous self, forgetting what had happened, and only reporting that he had a dim recollection of being overwhelmed by bad memories from his childhood.

There have also been cases of patients who, despite functioning seemingly well in most of their everyday activities, nonetheless experienced extreme personality changes and developed uncharacteristic behavior. For example, Bhargava and Doshi reported the cases of a 70–year–old male and a 58–year–old female suffering from advanced PD, who underwent subthalamic nucleus stimulation to treat the motor symptoms of PD. However, DBS induced sexual urges in these patients; urges they were unable to control. At the one–month follow–up visit, the wife of the 70–year–old man reported to the doctors that after DBS “he would insist on sexual gratification every night and would become very aggressive if denied. Once satisfied he would return back to his normal self.” Sim-

47 Chan, Zhu, Yeung et al. (2009).
48 Witt, Daniels, Reiff et al. (2008).
49 Rabins, Appleby, Brandt et al. (2009).
50 Ahlberg, Laakso, Hartelius (2011).
51 Montel, Bungener (2009).
52 Gilbert (2013).
53 Beeker, Schlaeper, Coenen (2017); Glannon (2010); Nyholm, Campbell (2016).
54 Denning, Matsuoka, Kohno (2009).
56 More recently, the potential of another neurotechnology, i.e., the memory-modifying potential of optogenetics, has prompted neuroethicists to discuss non-traditional risks also in the context of possible future use of optogenetics (see: Adamczyk, Zawadzki, 2020; Zawadzki, Adamczyk, 2021).
57 Baylis (2013); Clausen (2009); Gilbert (2018); Glannon (2014b); Hildt (2006); Merkel, Boer, Fegert et al. (2007); Nyholm, O’Neill (2016); Schechtman (2010); Schermer (2011); Synofzik, Schlaeper (2008); Witt, Kuhn, Timmermann et al. (2013).
58 Goethals, Jacobs, Van der Linden et al. (2008).
59 Bhargava, Doshi (2008).
60 Ibidem.
ilarly, the 58-year-old woman preoperatively had almost no sexual relationship with her husband for years, but after DBS she often forced herself into her husband’s room for sexual purposes. More strikingly, with time she also started to expose herself to other males in her family, demanding sex. Surprisingly, despite experiencing these changes, whenever doctors evaluated her, she acted as “an extremely mature person who showed no indications of her abnormal behavior.” This behavior continued for five years, until it was controlled with the introduction of an antipsychotic drug.

Another example of DBS-induced changes to the self of a person is the case of a patient suffering from OCD, known in the literature under the cryptonym “Mr. B.” His treatment with DBS was effective, but simultaneously caused an unexpected reaction. Under the influence of DBS therapy, Mr. B. became a fan of a very narrow subset of music: Johnny Cash’s songs from the late stage of his career, that is, when the musician’s voice became “harsher.” His new appreciation was so intense that he bought all of the performer’s records and assigned selected works to specific occasions and moods. Moreover, he claimed that listening to Cash’s music gave him confidence. When the device was turned off, however, Mr. B. lost interest in the artist’s music, along with his newly acquired self-confidence. Mr. B. applied two separate names for his self-transformations: “Mr. B. I” to the self when DBS was turned off, and “Mr. B. II” to the self with the activated device. He also behaved in a way that made the medical staff believe that whenever DBS was active, he felt like a new, improved version of himself.

Lipsman and Glannon, inspired by an actual clinical case, proposed an example of a 70-year-old professor suffering from PD, who, unlike Mr. B, did not feel that DBS treatment had radical self-transformative effects, but his relatives reported that it was the case. Along with a reduction in symptoms of the disease, the patient acquired a tendency to impulsiveness and mood swings, involving rapid changes from a depressed to a euphoric state. Due to these changes, his daughter reported that she no longer perceived her father to be the same person than before the therapy: “It’s as if he’s someone else entirely, […] he’s there, and he’s our dad, but not like we’ve always known him.”

More reports of cases in which patients experienced similar changes due to DBS treatment can be found in various qualitative studies, as well as the first systematic study addressing these unwanted effects in a semiquantitative way which has been

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61 Ibidem.
63 Some details of the actual case that inspired Lipsman and Glannon’s vignette was changed to preserve confidentiality and to emphasize some aspects of the case that the authors wanted to highlight. Thus, it is important to emphasize that although cases discussed by Lipsman and Glannon are not “uncommon” in the current DBS practice, there are fictional components in this particular vignette – in contrast to the all previously discussed cases.
64 Lipsman, Glannon (2013).
65 Agid, Schüpbach, Gargiulo et al. (2006); de Haan, Rietveld, Stokhof et al. (2013); de Haan, Rietveld, Stokhof et al. (2015); Gilbert, Goddard, Viaña et al. (2017); Gilbert, Viaña (2018); Gilbert (2018); Haahr, Kirkevold, Hall et al. (2013); Hariz, Limousin, Tisch et al. (2011); Houeto (2002); Lewis, Maier, Horstkötter et al. (2015); Liddle, Phillips, Gustafsson et al. (2018); Mathers, Rick, Jenkinson et al. (2016); Pham, Solbak, Skogseid et al. (2015); Scaratti, Zorzi, Guastafierro et al. (2020); Schüpbach, Gargiulo, Welter et al. (2006); Smeets, Duits, Horstkötter et al. (2018); Thomson, Segrave, Carter (2019); Thomson, Segrave, Racine et al. (2020).
Based on these results, there is now a broad consensus in the neuroethical literature that DBS may fundamentally alter the selves of patients. However, how a neuroethicist captures DBS-induced changes depends on a conceptual scheme she applies. Over the years, scholars have proposed various different understandings of the self, this elusive, yet familiar phenomenon.

**Theories of the self and neuroethical considerations of DBS**

One of the first theories of the self proposed in the neuroethical literature in an attempt to understand DBS effects was that of Synofzik and Schlaepfer. In opposition to the subjective tradition of theories of the self, such as the Cartesian concept, they propose a naturalistic model, describing the self as an objective, biological–cognitive representative system with the ability of self–representation. This self–representation, however, is not built by a homogeneous, monolithic entity. In contrast to how it phenomenally appears, the self is in fact constructed by various modules operating on different levels of representational and functional complexity. The most fundamental are levels consisting of sensomotor processes, while the most complex are those covering conceptual and meta–representational processes. This system of self–representation is able to produce subjective representations of the self of a person based on her actions, perceptions, emotions, and beliefs. Witt et al. criticize Synofzik and Schlaepfer’s model; they point out that if one was to consistently maintain it, the self would become an ephemeral, ever–changing phenomenon. As such, this approach is not helpful for understanding the actual clinical cases of patients that experienced DBS–induced changes.

Witt and colleagues propose a foundational–function model, in which only changes in central attitudes of the person, i.e., beliefs, expectations, desires, ideals, or plans, could pose a real threat to the self. From this hierarchical perspective, only DBS–induced changes in the most central attitudes imply self–disruption. In order to apply this theory to neuroethical considerations, one must identify which attitudes count as “core” or “central” for a person. Moreover, one should also ask why changes in core attitudes should matter from a moral perspective, as well as how one could distinguish between changes that matter morally from those that do not. The authors offer some guidance for the future empirical research that might allow clarification of these issues.

In contrast to both of the theories described above, Schechtman argues that the self should be understood in narrative terms. She claims that only narrative identity,
with its historical dimension allowing the introduction of patterns of coherence and meaning between individual experiences, meets a commonsensical expectation for our concept of the self. We create who we are through the construction of an autobiographical narrative. The relation of the narrative and the self can be conceptualized in two ways in this approach: narrative can play a strong role (i.e., create the self), or a weak role (i.e., be created by the self). However, we are probably dealing here with something similar to the hermeneutic circle, i.e., the self of the person is constructed by the narrative. On the other hand, this narrative is narrated by that person’s self. Not all narratives can be identified as constituting the self. For this to be the case, they must meet the constraints of articulation and reality. The first constraint states that a person must be able to present her story, life situation and motivations. It does not have to be an accurate and completely coherent story, but the narrator should be able to explain why she does what she does, believes what she believes, and feels what she feels. The second constraint indicates that the personal narrative must be coherent with the basic facts about the person and her life. The narrative cannot be delusional, rest on evident factual errors, be resistant to revision in light of contrary evidence, or be entirely inconsistent with how others view ourselves. Baylis considers Schechtman’s claim regarding the narrative and autobiographical nature of the self to be true. However, she embraces an account of the self inspired by feminist thought, in which the self is constituted in and through relationships. In her relational approach, Baylis recognizes embodied nature of the self that is situated in particular social, cultural, political, and historical contexts. She claims that the self resides not only in the brain, but in the negotiated spaces between one’s body and brain as well as the brains and bodies of others. In Baylis’s approach, the narrative constitutive for the self effectively balances between how the person perceives and understands herself, and how others perceive and understand her. In this model, the potentially threatening effects of DBS on the selves of patients are those that can disturb this balance in a way that would infringe on the person’s autonomy.

Problems of proposed models of the self and some suggestions for their resolution

It seems that the disagreement in the literature regarding the concept that adequately characterizes the self in the context of the neuroethical considerations concerning the consequences of DBS is profound. Furthermore, most approaches postulate relatively narrow concepts of the self, focusing on one or two of its aspects, and emphasize the representational, functional, narrative, or relational dimensions. As Dings and de Bruin note, the narrow scope of the proposed models of the self is problematic, as

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76 Baylis (2012); Baylis (2013).
77 Synofzik, Schlaepfer (2008).
78 Witt, Daniels, Reiff et al. (2008).
79 Schechtman (2009); Schechtman (2010).
80 Baylis (2012); Baylis (2013).
81 There is also important affordance–based account of de Haan, Rietveld, Stokhof et al. (2013) focusing on enactive dimension.
most of them aim to provide a complete understanding of the potential consequences of DBS on the self. Finally, the influential theories described above discuss DBS–induced changes in general, which renders it difficult to use them to account for changes specific to the disease treated with DBS, as well as changes specific to the type of DBS system utilized in a given therapy (i.e. OL–DBS, CL–DBS, or VCL–DBS).

This theoretical pluralism, combined with the lack of consensus regarding an adequate understanding of the concept of the self, has become a particularly pressing issue in recent years in the context of the ongoing neuroethical debate. These circumstances preclude the provision of a consensual explanation of the effects of DBS on the selves of patients, and consequently, prevent the development of a uniform procedure for evaluating the implications of DBS in individual clinical cases. In turn, this makes it impossible to calculate the benefits and risks related to decisions to initiate, continue, or discontinue the treatment.

At the same time, however, there can be a tendency in the neuroethical literature to regard considerations on the effects of DBS exclusively through the lens of a specific model of the self as an approach that is too coarse–grained. Realizing that theoretical disputes regarding the most adequate model of the self are far from reaching a resolution, neuroethicists have engaged in more fine–grained considerations of the effects of DBS on the selves of patients. For instance, some applied the concept of burden of normality to explain postoperative experiences of DBS patients. According to this notion, DBS patients may experience difficulties in adjusting to becoming “normal,” meaning that they cannot cope with the fact of being symptom-free. Although the notion of burden of normality is central in explaining some of the difficulties experienced by successfully “treated” DBS patients, the majority of the discussion of DBS postoperative changes to the self is focused on abnormal side effects. As the extensive literature review undertaken in this article reflects, neuroethicists employ various concepts related to the notion of the self in this debate such as personal identity, authenticity, and autonomy. The engagement of researchers and the constantly growing number of theoretical works analyzing the effects of DBS in terms of the concepts mentioned above show that these concepts are regarded as crucial to understanding a broader scope of DBS threats to the selves of patients. Thus, this paper attempts to present the most relevant potential consequences of DBS in these crucial “dimensions” of the self.

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82 Dings, de Bruin (2016).
83 Many thanks to the anonymous reviewer for pointing out that this important notion of the neuroethics of DBS is worth mentioning in this context.
84 Baertschi, Favez, Radomska et al. (2019); Gilbert (2012).
85 Baylis (2013); Bluhm, Cabrera, McKenzie (2019); Focquaert, DeRidder (2009); Lipsman, Glannon (2013); Mathews (2011); Schechtman (2010); Witt, Kuhn, Timmermann et al. (2013).
86 Gisquet (2008); Johansson, Garwicz, Kanje et al. (2011); Johansson, Garwicz, Kanje et al. (2014); Kraemer (2013a); Kraemer (2013b); Mackenzie, Walker (2015); Mackenzie (2014); Maslen, Pugh, Savulescu (2015); Mosley, Hall, Forlini et al. (2014); Nyholm, O’Neill (2016); Pugh, Maslen, Savulescu (2017a); Pugh, Maslen, Savulescu (2017b).
87 Brown, Thompson, Herron et al. (2016); Clausen (2010); Douglas (2014); Gilbert (2015); Gilbert, O’Brien, Cook (2018); Glannon (2014a); Goddard (2017); Goering (2015); Goering, Klein, Dougherty et al. (2017); Kellmeyer, Cochrane, Müller et al. (2016); Klein (2015); Müller, Walter (2010); Pugh, Pycroft, Sandberg et al. (2018); Unterrainer, Oduncu (2015); Wardrope (2014).
Personal identity and DBS

In Gilbert’s study, 8 out of 17 patients (47%) reported changes in self-perception after DBS. Some seemed to experience alterations to their sense of the self.\footnote{Gilbert (2018).} For example, Patient 4 said: “I feel like I am who I am now. But it’s not the me that went into the surgery […] No I can’t be the real me anymore—I can’t pretend. […] Well, I think that I felt that the person that I have been was somehow observing somebody else but it wasn’t me.” Patient 7 reported: “Oh God, I wasn’t me and I knew I wasn’t me and there was nothing I could do about it […] I knew what it was! I knew it had been turned up that day. Unlike the drugs which creep up on you and you don’t know what’s happening. With this, I knew what it was so I knew it was fixable.” Patient 13 revealed: “I would revert to a state of hysteric or something like that much more easily than I would normally have done […] I felt like I had lost my true self, it was way behind me.” Others reported changes in activities, mood, or socio-familial dynamics; for example, Patient 14 recounted: “I lost all interest in painting.” When interviewer asked: “Would you say that it was voluntary or involuntary loss of interest?,” the subject replied, “Involuntary. I tried sitting down and paint something, but I just didn’t want to do it.” Schüpbach reported that among 29 patients with PD (15 male, 14 female) examined during a 3-year study, 66% expressed feeling of strangeness and unfamiliarity with themselves after DBS.\footnote{Schüpbach, Gargiulo, Welter et al. (2006).} For example, Patient 1, a 38-year-old female journalist, married with one child, reported “Now I feel like a machine, I’ve lost my passion. I don’t recognize myself anymore.” Other patients in this study reported: “I don’t feel like myself anymore,” or “I haven’t found myself again after the operation.”

Schechtman’s narrative model of self,\footnote{Other versions of narrative concepts of personal identity can be found, for example, in the works of Ricoeur (1991), Mackenzie, Poltera (2010), or Atkins (2008).} in which narratives constitute the self if they meet articulation and reality constraints, has been most influential in the analyses of such cases.\footnote{Schechtman (1996); Schechtman (2009); Schechtman (2010).} If a person cannot construct a narrative fulfilling these conditions, or given dispositions, thoughts, or actions are recognized as resulting from DBS’s action, not from the individual’s plans, desires, or needs that are part of the narrative of the self, the personal identity of the patient is threatened. However, when considering the influence of DBS on the patients’ identities within the narrative account, Schechtman concludes that, although DBS can lead to disruption of personal narrative, the narrative approach itself suggests “a way to bring potential discontinuities back into one’s life narrative, rather than allowing them to disrupt it.”\footnote{Schechtman (2010).} In her narrative account, a person might reconstruct and reconstitute her identity in the face of randomness and temporal change, as well as contingencies, such as illness, accidents, traumas, or neurointerventions, by incorporating them into her self-understanding and life story. As the narrative is a dynamic concept, preservation of personal identity could be compatible even if the changes that the patients experience are substantial. Schechtman provides an example

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89 Schüpbach, Gargiulo, Welter et al. (2006).
90 Other versions of narrative concepts of personal identity can be found, for example, in the works of Ricoeur (1991), Mackenzie, Poltera (2010), or Atkins (2008).
91 Schechtman (1996); Schechtman (2009); Schechtman (2010).
92 Schechtman (2010).
of the resolution of this sort by recalling the case of an initially alienated woman studied by Schüpbach et al., who managed to cope with the fact that she had an electronic device in her brain by creating an artwork depicting her chest X-ray with the stimulator. Schechtman interprets this case as an example of a patient precluding the threat of DBS to her identity by engaging in constructing and maintaining a coherent narrative of change, and as a result, managing to post-operatively “find herself again”.

Narrative concepts are one of the most frequently used models of the self in the contemporary neuroethical debate around DBS; however, the ideas behind these concepts suffer from various problems. Mackenzie and Walker point out the following weaknesses of the narrative approach: narratives accounts suggest more authorial control over people’s lives and identities than they in fact have; imply that we must constantly self-consciously reflect on them; and involve interpretation rather than representation of our life stories, which raises the question of how one could distinguish truthful from confabulated ones. Because advocates of narrative accounts have already responded to these critiques, here, I focus on important arguments for why narrative accounts of personal identity are questionable, coming from recent research on the psychology of memory and reflections on cases of patients with neurological disorders.

Patients suffering from episodic amnesia whose episodic memory system has been disrupted or damaged seem to retain access to facts about their selves; that is, if a system of semantic self-knowledge of one’s own traits remains intact, it can potentially form the foundation for a sense of personal identity. In fact, some researchers have reached similar conclusions in recent years. This result suggests that narrative approaches do not adequately capture conditions for the sense of identity, since, due to the non-functional episodic memory system, a person not only cannot create a narrative that meets the articulation and reality constraints imposed by Schechtman, but may not

93 Ibidem.
95 Christman (2004).
96 Strawson (2004).
97 Matthews, Kennett (2012).
99 Episodic memory system records events as being experienced by the self from a specific point of view in time and space. Remembrance of these events takes place by re-experiencing them in a quasi-perceptual way, i.e., with the awareness that “this happened to me” (see: Tulving, 1985; Tulving, 1993; Wheeler, Stuss, Tulving 1997).
100 Semantic memory contains relatively general, contextless knowledge (e.g., apples are round). It usually “lacks memory” of the source of its origin, i.e., it is experienced as knowledge without reference to circumstances of where and when it was acquired. Semantic memory system may also contain knowledge about facts related to the self (e.g., I was born in Warsaw). However, one does not have to reproduce the context of acquiring this knowledge in order to retrieve information, even when the knowledge regards the very self of a person.
101 Knowledge of one’s own traits is a kind of semantic self-knowledge system (see: Tulving, 1985; Tulving, 1993; Wheeler, Stuss, Tulving, 1997). It has a form of generalizations and summaries of one’s dispositions. These generalized summaries are a form of database that allows decision-making processes to quickly evaluate one’s characteristics (see, e.g., Klein, Gangi, 2010; Klein, Cosmides, Costabile et al., 2002).
102 See: Craver (2012); Haslam, Jetten, Haslam et al. (2011); Illman, Rathbone, Kemp et al. (2011); Klein (2013).
be able to create any narrative at all about her life. Nevertheless, a person still retains relevant sense of the self. Such a conclusion seems to hold in several clinical cases of various neurological disorders. Obviously, the issue of the empirical adequacy of narrative models, as well as a potential falsification of the narrative as the necessary condition for personal identity, require further detailed discussion which is beyond the scope of this paper.

However, there are also issues associated with narrative theories that are more apparent and pressing in the context of neuroethical considerations. Narrative theories can be undermined not only from the position that they do not adequately show the foundation upon which the sense (or experience) of personal identity stands (necessary condition), but also from the argument that they do not give justice to the multidimensionality of the analyzed entity, namely, the self. This is because narrative theories fall into the category of “deflationary” conceptions of the self—they claim that the self is one thing and nothing more. This, in turn, does not allow for a sufficiently rich account of the self in the framework in which one could understand all relevant threats that DBS may pose to personal identity.

Two issues require resolution in this context. First, neuroethical considerations should make use of a theory of the self that finds the features of the self that are necessary to preserve personal identity. Second, this theory should embrace more aspects of the self. Responding to these concerns would help in constructing an account of personal identity which is capable of explaining the diachronic constitution and reconstitution of the identities of DBS patients.

Authenticity and DBS

In recent neuroethical literature, the concept of authenticity has become increasingly influential in addressing the threats to the self posed by DBS. For some, authenticity is in strict interrelation with narrative identity; if the desires of a person would have been fundamentally altered by DBS in a manner that does not fit within her narrative identity, then these desires are not authentically owned by this person. Others argue that: “The appeal to authenticity is redundant and blurs the distinction between narrative identity and autonomy.”

The most basic idea of authenticity was well expressed by William: “Some things are in some real sense really you, or express what you are, and others aren’t.” Such a characterization implies that humans have various elements of the self; some of these

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103 A disorder known as dysnarrativia.
104 See, e.g., Klein (2014); Klein (2012); Klein (2013); Klein, Nichols (2012); Klein, Lax (2010).
105 For example, there is a need for a conceptual discussion on the notions associated with personal identity, such as, sense of personal identity, self-continuity, sense of the self, as well as the issue of how they relate to one another.
107 For such a multidimensional proposition see: e.g. Dings, de Bruin (2016); Gallagher (2018); Zawadzki (2020).
108 For example, Christman (2009); Sharp, Wasserman (2016).
110 As cited in Johansson, Garwicz, Kanje et al. (2011).
elements are parts of the true selves, and others are not. In this approach, being authentic assumes we are living in accordance with elements that are part of our true selves, and alienated from those that are not. Kraemer claims that appealing to the concepts of “authenticity” and “alienation” in explaining the experience of patients after DBS could allow us to better understand these patients, and, in some cases, could even lead to the re-evaluation of the consequences of DBS for the self.111

In the context of DBS treatment, Kraemer interprets reports like “I feel like myself” as expressions of authenticity. Alienation, on the other hand, as the opposition to authenticity, is indicated by the following statements: “I felt like I had lost my true self, it was way behind me.”,112 or “I don’t feel like myself anymore.”113 Kraemer regards both authenticity and alienation as mental states. In her view, they are not neutral experiences, however; they involve normative component: authenticity is something that a person should strive for, alienation is something one should avoid. She believes that this normative component stems from the way we experience these states. Counterintuitively, however, authenticity does not necessarily have to be a pleasant experience for the person. Feeling authentic can be difficult, yet valuable. Similarly, alienation, despite being something one ought to avoid, may be experienced positively. Kraemer cites Frederick Neuhouser in explaining what alienation can “mean” for the person experiencing it: “a sense of meaninglessness or estrangement, a loss of power in relation to self and world, and a subjugation to the products of one’s own activity.” Both of these experiences seem to be present in patients receiving DBS treatment.

For example, in the case of Patient 1 in Schüpbach et al. mentioned above, researchers reported that she had been a dynamic person before DBS despite her motor handicap.114 However, after 18 months of treatment, despite a major improvement of her motor symptoms due to STN stimulation, she reported that she no longer felt like herself; she was no longer able to conduct any professional activity, constantly felt exhausted, and had lost her vitality, interest in her family, inspiration, and taste for life in general. Kraemer claims that the experiences of this patient can be explained when considered alongside the concept of alienation: “After treatment, not only does her work–life seem alienating, but also she feels alienated from life–goals she identified with, as well as from her new, improved body.”115

However, DBS can also have the opposite effect on the self; namely, patients may feel that the treatment allows them to find their true selves. Kraemer analyzed the case of Patient 2 from the study of Schüpbach et al., a 48–year–old male accountant who worked part–time for 7 years because of PD. Before DBS, he required daily care, with his wife helping him with all of his daily tasks. After receiving treatment, he felt empowered and decided to regain control over his life. He reported “During all these years I was asleep, now I am going to take my life in hand, my life before PD.” Kraemer interprets this case as reintroducing the connection of the patient with his own true self. She suggests that

111 Kraemer (2013b).
113 Schüpbach, Gargiulo, Welter et al. (2006).
114 Ibidem.
115 Kraemer (2013b).
the claim about “taking life in his own hands” indicates that, before DBS, the patient felt that he was inauthentic, and alienated from his true self; the treatment, however, made him ready to become the “master of his own destiny,” with the means to live a life aligned with his true self.

Kraemer postulates that both Patient 1 and Patient 2 seem to distinguish two modes in their lives: “mode one,” that is, the mode experienced when they regard themselves as authentic; and “mode two,” the mode they experience when alienated from their true selves. For Patient 2, the “authentic mode” was realized when under treatment; then, he felt ready to reclaim his real life (“I want to regain my life, my life before PD”). For him, PD was the cause of being in “mode two”. He felt that his illness put his true self to sleep; he felt that his life was not really his own during PD, and thus he reported being alienated. Strikingly, the experiences of Patient 1 are in contradiction with those of Patient 2. Patient 1 had felt that her “mode one” was present when she was diseased. Her daily struggle with PD gave her goal in life. She reports: “Before stimulation, I wanted to be like everybody else, I didn’t want to be considered only as sick. I fought for that every day! Now I find myself less impassioned, I regret the period where I did battle. Now it’s the warrior’s repose, I no longer have something to struggle against, my life is empty. I get up every day, but have no goal, no horizon.”

Kraemer notes that Patient 2 identifies authenticity with autonomy. For him, in order to become himself, he must regain autonomous functioning by freeing himself from the care of his wife. It is worth pointing out that the case of Patient 1 also shows the crucial role of autonomy in experiencing authenticity. Her constant struggle with her disease gave Patient 1 purpose; its realization gave her a sense of agency, which she could not find after the successful treatment. Both of these interpretations are consistent in that prominent philosophical accounts of autonomy take authenticity as partially constitutive of autonomy. For example, Christman explicitly states “Put most simply, to be autonomous is to be one’s own person, to be directed by considerations, desires, conditions, and characteristics that are not simply imposed externally upon one, but are part of what can somehow be considered one’s authentic self.” For this reason, in the following sections I focus on the latter of those dimensions potentially endangered by the effects of DBS—autonomy.

**Autonomy and various DBS systems**

The reference point in the neuroethical literature concerning the possible effects of DBS on autonomy is a hierarchical model proposed by Frankfurt. Frankfurt argues that autonomy requires a person’s ability to possess second–order volitions with which she identifies—that is, regards them as authentic elements of herself. Crucially, second–or-

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116 For a systematic approach to the role of such narrative factors in the sense of agency, see: Zawadzki (2021).
117 It is worth noting that this case can also be interpreted as an example of the “burden of normality”.
118 Wardrope (2014).
120 Frankfurt (1969); Frankfurt (1971).
der volitions also enable a person to distance herself from spontaneous, often elusive, first–order desires. In this view, autonomy emerges when there is a congruency between a person’s second–order volitions and her first–order desires; an autonomous person endorses her first–order desires. In contrast, autonomy is disrupted when there is a discrepancy between second–order volitions and first–order desires, but the person still behaves in accordance with the dictates of the latter.

One example, where Frankfurt’s model of autonomy has been applied from the ethical point of view of DBS treatments concerns patients with anorexia nervosa. Müller et al. suggest that people suffering from anorexia become caught in a vicious circle from which they cannot escape: they want to starve themselves despite knowing that such behavior is unhealthy and threatens their lives.121 Interpreting this scheme against the background of Frankfurt’s theory of autonomy, the first–order desires dominate over the second–order volitions of these patients.

Maslen, Pugh, and Savulescu discuss the potential of DBS as a means to treat the symptoms of anorexia.122 They debate various strategies to achieve this goal; one, inspired by Frankfurt’s theory, is termed “promotion of comparative cognitive control.” There are two ways in which comparative cognitive control can be enhanced with DBS: either by reducing the compulsive need to diet and avoid eating, or by increasing top–down control over this compulsion. If the patient experiences an irresistible first–order desire that does not align with her autonomously chosen goals, reducing the strength of such a compulsive urge with the use of DBS can help the patient to regain control and act in a manner aligned with her higher–order desires. Conversely, in cases in which the patient feels that the motivation to engage in weight loss behaviors expresses her authentic self, she may not welcome an attempt to reduce her first–order desire that leads to a positively valued end. However, Maslen, Pugh, and Savulescu claim that, even in such situations, DBS could still promote the person’s autonomy even if it only serves as a means to give her resources to be more successful in self–governing by remedying a failure in top–down cortical control over compulsive drives to diet. Maslen and colleagues compare this strategy to the situation of a person, who can resist drinking, but nonetheless chooses to drink on some occasions. This strategy, however, seems riskier than the former, as it assumes that the person will choose (due to the additional cognitive resources) to behave in a manner aligned with other second–order desires, like desire to survive, sufficiently often.

Frankfurt’s autonomy model can help examine the ethics of DBS treatment, and it even illuminates the therapeutic potential of DBS in cases of patients suffering from certain diseases, such as anorexia. However, the future of the ethics of DBS will require a theory of autonomy that takes into account not only challenges related to the specific ailments DBS aims to treat, but also the new kinds of threats arising from the technology of DBS itself—that is, emerging forms of treatments, like advisory brain devices123 or VCL–DBS.124 It might seem that patients’ autonomy will only be enhanced by these

121 Müller, Riedmüller, Walter et al. (2015).
122 Maslen, Pugh, Savulescu (2015).
124 Brown, Moore, Herron et al. (2016).
technologies, as patients play a decisive role in applying stimulation with such systems. While some authors argue that this is the case, others claim that such a result is far from obvious. Below, these issues are analyzed in the context of widely debated case of “the Dutch Patient” (DP); I discuss it in a different context than it is usually the case in neuroethical literature. Most authors consider the ethics of this case from the perspective of taking the most adequate measures to respect DP’s autonomy in his complex institutional situation. Below, the hypothetical situation in which DP uses one of the emerging forms of DBS treatments is presented as an illustration of a potential threat to his autonomy that could arise solely from the technology used.

DP was treated for PD with the help of OL–DBS. The treatment was effective, but it caused multiple side effects: mania, megalomania, and impulsiveness. Given that the treatment appeared to be leading the patient to compulsive gambling, falling into debt, conflict with the police, and ultimately, forced hospitalization in a psychiatric hospital, doctors faced a dilemma about whether the device should be permanently disabled. The solution that seemed appropriate to the medical staff was to temporarily turn off the device and then ask the patient about his preferences concerning the continuation of DBS therapy. With his DBS disabled, DP decided that it was more important to him to stop the symptoms of PD, even given the symptoms caused by DBS, and signed an advance directive agreeing to remain under psychiatric care while the device was activated.125

There is much to consider from the ethical point of view of DP. For example, whether DP’s behavior under DBS was the result of realizing authentic desires of his true self; whether it was autonomous;126 and even whether DP was autonomous when the device was turned off, or if DBS irreversibly changed him.127 Here, however, as mentioned above, even more complicated hypothetical scenario building upon the case of DP, in which is assumed to have used a DBS system of the new generation, will be considered.

New DBS systems allow patients to stay in “the decisional loop.” However, there is no data concerning the impact of such systems on the sense of self of the patients. Recently, Gilbert, O’Brien, and Cook addressed this gap by examining how the first–in–humans experimental advisory brain devices capable of predicting epilepsy seizures influence the sense of autonomy of patients.128 To this end, researchers conducted in–depth semi structured interviews to obtain first–person narratives of the subjective experiences of patients. Gilbert and colleagues regrouped subjects into five phenomenological clusters reflecting patients’ autonomy and decision–making processes: (1) insecurities and risks attached to living with epilepsy; (2) how patients integrate device predictions into their deliberative processes leading up to their relevant decisions; (3) patients’ distrust of the device; (4) device–induced sense of control and empowerment; and (5) device–induced lack of confidence and sense of control. The authors’ analysis of the data led them to the following conclusions: 1) being in the loop may partly increase a sense of autonomy over decision–making; 2) being in the loop may partly decrease a

125 Smeding (2008).
126 Kraemer (2013a).
127 For a discussion of the reversibility of DBS see: Pugh (2019).
sense of autonomy over decision-making; 3) being in the loop may not impact a sense of autonomy over decision-making.

Here, I would like to focus on one seemingly paradoxical consequence of Conclusion 1, namely that increasing a sense of autonomy over decision-making by staying in the decisional loop can actually lead to a decrease in a patient’s overall autonomy. Klein et al., discussing the ethics of VCL–DBS in the context of psychiatric disorders, raises this issue by indicating that there is a possibility that “if a patient is given control over device settings, the temptation to increase stimulation settings to feel better and better may be difficult to resist, and patients may fear the introduction of a new kind of addiction.”\textsuperscript{129}

It is worth noting here that VCL–DBS systems may not only threaten patient autonomy when used to treat addiction; it seems to be an inherent risk of the use of DBS systems that give control to the patients (regardless of the disorder being treated).

Even in the case of PD patients, such as DP, there is a threat to autonomy if the patient is treated with the use of a VCL–DBS system. If experiencing DBS–related euphoria, a common symptom in the manic state, DP could experience an insurmountable temptation to turn on and amplify the stimulation, in order to achieve this phenomenologically pleasant mental state. As he would have total control over the device, he could enter a self–destructive cycle, leading him to engage in riskier, or even dangerous actions. As Brown, Moore, et al. point out in discussing the case of DP: “Mania is often accompanied by feelings of grandiosity or invulnerability, and so a patient may refuse to acknowledge the negative consequences of their stimulation.”\textsuperscript{130} They further comment that for this reason, DP’s autonomy could be “hijacked” or could become “addicted” to the system through a feedback loop that causes him to decide to leave the device constantly turned on. In this type of case, what is particularly problematic is that, if medical staff working with OL–DBS could easily detect dangerous changes in behavior at the time of programming, with VCL–DBS, patients could potentially trigger and experience changes over time, in a manner that preclude early detection and re–programming of the device. Moreover, patients “addicted” from achieving a desired psychological state may even intentionally hide dangerous symptoms caused by the stimulation by turning it off during medical appointments.

Interestingly, Pugh et al. argue that one may still understand DP’s choice as enhancing his autonomy, if the stimulation allows him to achieve the goals that he himself believes that he has most reasons to achieve.\textsuperscript{131} They quote Unterrainer and Oduncu in using the notion of Ulysses’ contract to analyze the case of DP in this context.\textsuperscript{132} Just as Ulysses tied himself to the mast as a means to protect himself from being lured to his death by the Sirens’ song, so DP had to reduce his autonomy to achieve his desired end; DP chose to live in a state of mania, in order to avoid remaining bed–ridden due to his motor disability. Pugh and colleagues argue that this means DP’s decision can be interpreted as actually enhancing autonomy. His actions facilitated the pursuit of the desired end he felt was most important for him, even if the effects of his decision took away his freedom to achieve other competing goals.

\textsuperscript{129} Klein, Goering, Gagne et al. (2016).
\textsuperscript{130} Brown, Moore, Herron et al. (2016).
\textsuperscript{131} Pugh, Pycroft, Sandberg et al. (2018).
\textsuperscript{132} Unterrainer, Oduncu (2015).
While this perspective may seem convincing in the particular case of DP, it can
be ethically problematic as a general rule; DP’s decision not only takes away his liberty
in pursuing competing goals, but also undermines his autonomy in regard to all possible
decisions afterwards. Although making decision with the DBS device disabled allowed
him to exercise autonomy in a given moment in time, it led him to sacrifice his autonomy
in the future. One could understand this dilemma better as the choice between local and
global autonomy. In a local understanding of autonomy, one may ask whether a person
is autonomous with regard to a particular decision in a given situation at a given time,
and whether she has the necessary resources to act in a manner aligned with her will.
In a global sense of autonomy, on the other hand, one may ask whether a person is able
to autonomously pursue chosen goals over an extended period. From this perspective,
one could argue that DP enhanced his local autonomy at the cost of reducing, or in his
extreme case, erasing his global autonomy (as DBS diminished DP’s autonomy both in
terms of permanent volitional impairments, e.g., compulsive gambling and mania, and
the physical restriction of his freedom in the form of compulsory lifelong hospitaliza-
tion). Although local autonomy, the autonomy of a particular decision, is relevant, there
are situations in which we should give it up in order to make paternalistic decisions to
preserve the global autonomy of the patient. This is particularly important if this kind
of decision has severe life-altering consequences. DP’s case could seemingly serve as
an example of such an approach; however, in his case, the relevant choice is not only
between different kinds of autonomy, but also between global autonomy and movement
abilities required for normal everyday functioning.

Taking into account the fact that DBS is a last resort therapy, patients such as DP
have failed to respond to all other treatments and thus it is imperative to emphasize that
DP would continue to suffer from untreated symptoms of the disease without DBS. It
therefore seems that DP has to face a dramatic choice between prisons of different sizes
and types—in the case of being treated, a ward and a “mental prison,” and if left un-
treated, a bed. This makes DP’s choice almost impossible to assess from a neuroethical
standpoint and devastating for the patient and his family. At this point, we should also
note that PD that could make DP bedridden poses a threat not only to his autonomy, but
also to his identity and authenticity. Thus, symptoms of the disease that DBS can control
are as much of a threat to these dimensions of the self as the adverse side effects of the
treatment. It should therefore be emphasized once again that there is no choice between
DBS and the “neutral state” of the patient. The choice concerning taking DBS therapy
is always entangled in a situational context in which the patient has a very serious and
intractable disease. To conclude, the case of DP is extremely complicated from the neu-
roethical perspective because it is profoundly difficult to assess whether the symptoms
of the disease or the institutional and psychiatric side effects of treating it should be
considered more beneficial (or harmful). Moreover, it is impossible to simply transfer the
responsibility for the decision to the patient himself (which might seem the best option
in such a difficult situation) because it is not obvious at which moment the patient is ac-

133 Dworkin (1988).
134 After all, what is worse – the symptoms of the disease or the institutional and psychiatric side
effects of treating it?
tually given an autonomous choice (whether it is when his local (synchronous) or global (diachronic) autonomy is respected). As discussed above, both DP’s decision-making process as well as his final decision depend on what kind of autonomy decision-makers (i.e., legal guardians, clinicians, ethical boards) choose to respect.

The utilization of OL–DBS in the actual case of DP led him to be considered non-autonomous, and effectively prompted the medical staff to temporarily turn off the device. Only under this condition did doctors ask him about his preferences concerning further therapy. In this decision, the local autonomy of the patient was the most important factor. The utilization of a VCL–DBS system could further complicate cases such as DP’s, as its use can be intertwined with a sense of autonomy and self-narrative of the patient. Unlike OL–DBS systems, where there seems to be clarity concerning the forces distinct from the self which influence the behavior of a person, in VCL–DBS, a patient may experience the sense of agency stemming from the control over the stimulation. This could lead the patient to feel more responsible for his actions during treatment, as he might not attribute the changes in his behavior to the external force of DBS, but rather to his own weakness of will that prevented him from turning off the stimulation.

Such considerations lead to the conclusion that OL–DBS and VCL–DBS systems can have both positive and negative consequences for the autonomy of patients. The influence of each of these types of DBS, however, differs in substantial ways. Being in the decisional loop can further complicate the impact of DBS on patient autonomy, instead of straightforwardly enhancing it. Moreover, local (or synchronous) autonomy is not necessarily the same as the global (or diachronic) autonomy of the patient. Finally, discussion of the case of DP indicates one more relationship that must be addressed regarding the ethics of DBS; namely, that autonomy of a person is interwoven with her own sense of responsibility. However, as there is currently a gap in the knowledge concerning how remaining in the decisional loop influences patients’ sense of responsibility, and there is only one theoretical work concerning hypothetical cases that has sought to prepare the stage for conceptual considerations regarding this issue, more work in neuroethics will be required to understand this complex relationship.

Conclusions

The article examined potential threats posed by various DBS treatments to the selves of patients and various dimensions related to these threats: personal identity, authenticity, and autonomy. The most influential theoretical models and practical considerations explored in the neuroethical literature have been critically analyzed in this paper. I argued that various proposed models may be theoretically, empirically, or ethically questionable, and thus their further refinement seems to be required.

I have pointed out why narrative models of personal identity seem to be empirically questionable, and why their deflationary character poses a problem for accounting for DBS threats to the identities of patients. Focusing on different aspects of DBS threats to the self, I applied concepts of authenticity and alienation to better understand patients’ reports after DBS. The analysis of clinical cases undertaken here suggests that the issue of

135 Brown, Moore, Herron et al. (2016).
authenticity is interwoven with the issue of autonomy of the patient. Frankfurt’s account of autonomy, with its embedded authenticity reference, proved helpful in examining the ethics of DBS treatment, as well as in illuminating therapeutic potential of DBS in cases of patients with anorexia. However, as pointed out, modern DBS systems (e.g., VCL–DBS) require a theory of autonomy that can account for new kinds of threats arising from technological innovation in DBS treatments. In particular, the issue of “being inside the decisional loop” was analyzed against the background context of various related theories. I have adopted the notions of local (synchronous) and global (diachronic) autonomy to account for the potential consequences of utilizing such systems in DBS therapies, and concluded that these two concepts may sometimes be at odds with each other. Finally, I have pointed out the relationship between autonomy and the sense of responsibility in DBS patients and indicated the need for neuroethicists to further study this relationship.

This paper sought to demonstrate that the effects of DBS on the selves of patients cannot be accounted for solely through the lens of any specific extant model of the self; more fine-grained concepts are required. The extensive review of the literature undertaken in this article shows that neuroethicists should employ notions of personal identity, authenticity, and autonomy in the ongoing debate, as DBS poses tangible threats to the selves of patients in these dimensions. The considerations described and examined in this paper show that these dimensions are intertwined, and that DBS’s potential impact on the selves of patients can only be properly understood when all of these factors are taken into account. Hence, novel approaches that carefully scrutinize the complex interrelations between these dimensions are needed to gain new insights into the ethics of neuromodulatory treatments such as DBS.

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