Parkinson's disease subtypes: critical appraisal and recommendations

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Systematic Review

Parkinson’s Disease Subtypes: Critical Appraisal and Recommendations

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Abstract. 
Background: In Parkinson’s disease (PD), there is heterogeneity in the clinical presentation and underlying biology. Research on PD subtypes aims to understand this heterogeneity with potential contribution for the knowledge of disease pathophysiology, natural history and therapeutic development. There have been many studies of PD subtypes but their impact remains unclear with limited application in research or clinical practice.

Objective: To critically evaluate PD subtyping systems.

Methods: We conducted a systematic review of PD subtypes, assessing the characteristics of the studies reporting a subtyping system for the first time. We completed a critical appraisal of their methodologic quality and clinical applicability using standardized checklists.

Results: We included 38 studies. The majority were cross-sectional (n = 26, 68.4%), used a data-driven approach (n = 25, 65.8%), and non-clinical biomarkers were rarely used (n = 5, 13.1%). Motor characteristics were the domain most commonly reported to differentiate PD subtypes. Most of the studies did not achieve the top rating across items of a Methodologic Quality checklist. In a Clinical Applicability Checklist, the clinical importance of differences between subtypes, potential treatment implications and applicability to the general population were rated poorly, and subtype stability over time and prognostic value were largely unknown.

Conclusion: Subtyping studies undertaken to date have significant methodologic shortcomings and most have questionable clinical applicability and unknown biological relevance. The clinical and biological signature of PD may be unique to the individual, rendering PD resistant to meaningful cluster solutions. New approaches that acknowledge the individual-level heterogeneity and that are more aligned with personalized medicine are needed.

Keywords: Parkinson’s disease, heterogeneity, subtypes

INTRODUCTION

Parkinson’s disease (PD) is known for its heterogeneity in terms of both clinical practice and underlying biology [1]. One approach to understanding the variability present in PD is to identify subtypes; that is, to define groups of patients with a set of differentiating features at a clinical, biological, genetic or pathological level.

Research on PD subtypes has garnered considerable attention for its potential to elucidate the disease pathophysiology and natural history and ultimately, guide therapeutic development. However, the impact of PD subtype identification on our understanding of PD pathogenesis or clinical treatment remains unclear with limited application in research or clinical practice. Methodologic heterogeneity, lack of reproducibility, poor prognostic value, and limited clinical applicability are among the possible reasons for the reduced impact of PD subtypes [2].

The Task Force for PD Subtypes was convened in 2018 by the International Parkinson’s disease and Movement Disorders Society (MDS) to critically appraise the available PD subtyping studies and to provide guidance for the design and conduct of future studies on PD subtypes. In this systematic review of PD subtypes, we evaluate both the methodologic quality and clinical applicability of studies proposing a new subtyping system. Based on these findings, we provide our assessment of the future of subtyping research.

METHODS

Overview

We searched for PD subtyping studies in PubMed/MEDLINE using the following search terms: ‘Parkinson Disease’[Mesh] AND (‘Subtype’ OR ‘Phenotype’)[Mesh] OR ‘Phenotyp*’ OR ‘Biomarkers’
We identified 51 studies after the initial screening, of which 13 studies were excluded after full-text review (Supplementary Figure 1). Of the 38 included studies, 68.4% were cross-sectional (n = 26) and 84.2% were conducted in a tertiary care center (n = 32). Across all studies, the mean disease duration was 59.8 months. Nine studies (23.7%) included untreated participants exclusively. Commonly used descriptors of PD patients (e.g., Unified Parkinson’s Disease Rating Scale scores) were not reported in a significant number of studies (Table 1).

Identification and description of reported PD subtypes

In the 38 included studies, 65.8% (n = 25) used exclusively a data-driven approach and 21.1% (n = 8) used exclusively a hypothesis-driven approach, evaluating differences between pre-determined groups. Five studies either used a combined approach (n = 2) or alternative analytical approaches such as regression analyses or post-hoc grouping based on results of the study (n = 1). Most of the data-driven studies (n = 16/25, 64.0%) used at least three phenotypic domains to identify PD subtypes, while 7/8 hypothesis-driven studies used a single domain. Overall, the motor domain was most frequently used.
Phenotypic domains and statistical methods used in the included studies

<table>
<thead>
<tr>
<th>Phenotypic domain(s) - N (%)</th>
<th>Hypothesis-driven studies</th>
<th>Data-driven studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demographic</td>
<td>2 (25.0)</td>
<td>8 (32.0)</td>
</tr>
<tr>
<td>Motor</td>
<td>3 (37.5)</td>
<td>18 (72.0)</td>
</tr>
<tr>
<td>Cognitive</td>
<td>1 (12.5)</td>
<td>17 (68.0)</td>
</tr>
<tr>
<td>Emotional</td>
<td>1 (12.5)</td>
<td>17 (68.0)</td>
</tr>
<tr>
<td>Autonomic</td>
<td>1 (12.5)</td>
<td>11 (44.0)</td>
</tr>
<tr>
<td>Treatment</td>
<td>-</td>
<td>3 (12.0)</td>
</tr>
<tr>
<td>Non-clinical Biomarkers*</td>
<td>-</td>
<td>5 (20.0)</td>
</tr>
</tbody>
</table>

(*Imaging (n = 4), biochemical (n = 1)

Statistical approaches

<table>
<thead>
<tr>
<th>Pre-determined groups</th>
<th>Hierarchical cluster analysis</th>
<th>Non-hierarchical cluster analysis</th>
<th>Hierarchical and Non-hierarchical cluster analysis</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>8 (100)</td>
<td>-</td>
<td>7 (28.0)</td>
<td>16 (64.0)</td>
<td>1 (4.0)</td>
</tr>
</tbody>
</table>

Five studies were not included in this analysis for the following reasons: combined hypothesis- and data-driven approach (n = 2) and other methods (regression analyses, n = 2; subtype criteria defined post-hoc, n = 1).

for subtyping in both types of studies, followed by non-motor domains, such as cognitive, emotional or autonomic, in studies using a data-driven approach, and the demographic domain (age or sex) in hypothesis-driven studies. Only five studies included non-clinical biomarkers, all of which adopted a data-driven approach (Table 2).

From the descriptions of the resulting subtypes, reviewers identified which specific features distinguished the groups in a statistically significant manner. The reviewed PD subtypes were found to have on average three statistically significant distinctive features between subtypes (mean = 3.3, range: 0–9). Features within the motor domain were the most commonly reported to differentiate PD subtypes. Imaging biomarkers were found to be a differentiating feature in three out of four studies that included this domain (Table 3).

Methodologic quality (Table 4, Supplementary Table 2)

Most of the studies did not achieve the top rating across methodologic quality items, with the exception of the items 'diagnostic methods' and 'variables compared between subtypes'. Statistical methods were difficult to evaluate. Emerging themes were recruitment from clinics rather than community sources, the lack of reporting of specific methods of clustering, lack of specification of how the number of clusters was determined, lack of justification of sample size, lack of adjustment for multiple comparisons when comparing features of the clusters, and lack of adjustment for or exploration in the report of the impact of fundamental baseline characteristics such as disease duration [3].

The four studies with the best quality ratings [4–7] were multi-center, longitudinal, data-driven studies published after 2016. These studies used more than one clinical domain (motor and non-motor) and only one study used additional CSF and neuroimaging biomarker data [7]. Two of the four studies used a homogeneous PD population in terms of stage/disease duration and conducted validation of the identified PD subtypes [4, 5]. In spite of incorporating similar clinical domains represented in the cluster analyses, there were no clear similarities among subtypes described in these two studies [4, 5]. Only one study developed an algorithm to classify individual participants into a PD subtype [4] and only one of these studies assessed the temporal stability of PD subtypes [6].

Compared with hypothesis-driven studies, data-driven studies more frequently used multi-center data collection (p = 0.04) and more than one clinical domain or biomarker (p = 0.003). We did not identify significant differences between the groups of studies published before or after 2015 (Supplementary Table 4).
Table 4
Item score distributions in the 13-item Methodologic Quality tool of included subtyping studies using a data-driven or hypothesis-driven approach

<table>
<thead>
<tr>
<th>Item</th>
<th>Score rating</th>
<th>Hypothesis-driven (n = 8)</th>
<th>Data-driven (n = 25)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disease stages/duration (study population)</td>
<td>0 = mixture of stages/disease duration at baseline or not reported</td>
<td>5 (62.5)</td>
<td>18 (72)</td>
<td>0.67</td>
</tr>
<tr>
<td></td>
<td>1 = homogeneous disease stage/duration</td>
<td>3 (37.5)</td>
<td>7 (28)</td>
<td></td>
</tr>
<tr>
<td>Study setting (representativeness)</td>
<td>0 = single-center or not reported</td>
<td>7 (87.5)</td>
<td>10 (40)</td>
<td>0.04</td>
</tr>
<tr>
<td></td>
<td>1 = multi-center</td>
<td>1 (12.5)</td>
<td>15 (60)</td>
<td></td>
</tr>
<tr>
<td>Recruitment source (generalizability)</td>
<td>0 = clinic-based or not reported</td>
<td>8 (100)</td>
<td>25 (100)</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>1 = community or population-based</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td>Diagnostic methods</td>
<td>0 = not described or 1 or 2 not applicable</td>
<td>2 (25)</td>
<td>1 (4)</td>
<td>0.04</td>
</tr>
<tr>
<td></td>
<td>1 = Use of formal diagnostic criteria or diagnosis by an expert neurologist</td>
<td>6 (75)</td>
<td>24 (96)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 = postmortem diagnosis</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td>Sampling method</td>
<td>0 = convenience or not reported</td>
<td>5 (62.5)</td>
<td>21 (84)</td>
<td>0.32</td>
</tr>
<tr>
<td></td>
<td>1 = consecutive or random</td>
<td>3 (37.5)</td>
<td>4 (16)</td>
<td></td>
</tr>
<tr>
<td>Comprehensiveness of data used for subtyping (subtype definition)</td>
<td>0 = single clinical or biomarker domain</td>
<td>8 (100)</td>
<td>9 (36)</td>
<td>0.003</td>
</tr>
<tr>
<td></td>
<td>1 = &gt; 1 clinical domains or biomarkers</td>
<td>0 (0)</td>
<td>16 (64)</td>
<td></td>
</tr>
<tr>
<td>Variables compared between subtypes (post hoc)</td>
<td>0 = not done</td>
<td>0 (0)</td>
<td>2 (8)</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>1 = single clinical domain or biomarker</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 = &gt; 1 clinical domains or biomarkers</td>
<td>8 (100)</td>
<td>23 (92)</td>
<td></td>
</tr>
<tr>
<td>Statistical methods used for subtyping</td>
<td>0 = low quality</td>
<td>4 (50)</td>
<td>4 (16)</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>1 = intermediate quality</td>
<td>3 (37.5)</td>
<td>10 (40)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 = high quality</td>
<td>1 (12.5)</td>
<td>11 (44)</td>
<td></td>
</tr>
<tr>
<td>Longitudinal follow-up</td>
<td>0 = none (cross-sectional) or longitudinal &lt; 1 year</td>
<td>6 (75)</td>
<td>17 (68)</td>
<td>0.68</td>
</tr>
<tr>
<td></td>
<td>1 = short-term (1–3 years) OR longer-term but &lt; 3 time-points</td>
<td>2 (25)</td>
<td>4 (16)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 = longer-term (&gt; 3 year) AND &gt; = 3 time-points</td>
<td>0 (0)</td>
<td>4 (16)</td>
<td></td>
</tr>
<tr>
<td>Completeness of follow-up</td>
<td>0 = cross-sectional or ≤ 50% complete or not reported</td>
<td>7 (87.5)</td>
<td>20 (80)</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>1 = 50–75% complete</td>
<td>1 (12.5)</td>
<td>2 (8)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 = &gt; 75 % complete</td>
<td>0 (0)</td>
<td>3 (12)</td>
<td></td>
</tr>
<tr>
<td>Subtype stability</td>
<td>0 = not assessed</td>
<td>8 (100)</td>
<td>23 (92)</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>1 = assessed</td>
<td>0 (0)</td>
<td>2 (8)</td>
<td></td>
</tr>
<tr>
<td>Algorithm for classifying individual patients</td>
<td>0 = not provided</td>
<td>0 (0)</td>
<td>24 (96)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td></td>
<td>1 = provided</td>
<td>8 (100)</td>
<td>1 (4)</td>
<td></td>
</tr>
<tr>
<td>Validation (internal or external)</td>
<td>0 = not assessed</td>
<td>8 (100)</td>
<td>19 (76)</td>
<td>0.57</td>
</tr>
<tr>
<td></td>
<td>1 = use of a test set from the same population</td>
<td>0 (0)</td>
<td>4 (16)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 = validation in an external population</td>
<td>0 (0)</td>
<td>2 (8)</td>
<td></td>
</tr>
</tbody>
</table>

Clinical applicability (Supplementary Table 3, Supplementary Figure 2)

Overall, items reflecting the clinical importance of differences between subtypes, potential treatment implications and applicability to the general population of PD patients were rated poorly for most studies. Subtype stability over time and prognostic value were largely unknown due to the paucity of longitudinal studies. Compared with hypothesis-driven subtypes, data-driven subtyping was seen as burdensome and time-consuming (p = 0.01). Both data-driven and hypothesis-driven subtyping were usually rated as inexpensive. There were no clear differences between newer and older studies for the different items of the Clinical Applicability tool (Supplementary Figure 3).

DISCUSSION

Our systematic review has revealed gaps in the field and highlighted the limitations of current approaches to PD subtyping. It thereby informs recommendations on the methodology of future subtyping studies and suggests alternative directions. Recommendations are highlighted in italics.
Research methodology: Can we do better?

Quality ratings for subtyping studies revealed clear areas for improvement. More extensive use of longitudinal data is critical for an understanding of the stability of proposed subtypes, and their prognostic value. The use of longitudinal data to define or evaluate subtypes does appear to be more common in the last 5 years [4–8], making use of large, publicly available cohorts [4, 9]. To our knowledge, only one study has used longitudinal profiling as the basis for defining subtypes [7], incorporating data on the evolution of clinical or biological features across time into the definition of subtypes. Alternatively, serial cluster analyses could provide data about the stability of proposed subtypes and the influence of disease duration on their characteristics. Such approaches could provide additional prognostic value, using information about the early evolution of disease to inform later prognosis or underlying biology.

An area for future study is incorporating the prodromal phase of PD, which presents the opportunity to start defining subtypes of the disease earlier in the pathological process, which may provide new insights. A better understanding of the starting point and initial progression may better predict the course and subsequent clinical progression and offer the opportunity for an early target-specific therapeutic development.

A recurring reporting flaw was failure to describe statistical methods in detail. This issue may become more frequent as machine learning will probably play an important role in complex (hypothesis-free) analyses, particularly as the scope of available data expands [10]. It is important that the clustering method that is used is explicitly described in sufficient detail to facilitate independent replication and pursuit of subsequent hypothesis-driven studies [11].

Data-driven analyses were rarely replicated in a separate cohort. This is likely related to the fact that similar data are not always available on different cohorts and cohorts differ in their eligibility criteria (and assessments), resulting in fundamentally different characteristics of patients. However, it was also rare for studies to use internal replication techniques such as resampling. Internal or external replication attempts are important to assess the general applicability of suggested subtyping methods.

It was noted that the domains covered in subtyping studies and the instruments used were variable, limiting the value of comparisons across studies. Consensus moving forward as to the methodologic approach and the key core set of domains would be helpful for the field.

Subtyping by design

Possible uses of subtyping include clinical prognostication or identifying biological subtypes that predict therapeutic response to symptomatic (targeting convergent mechanisms, such as dopamine deficiency) or disease-modifying interventions (targeting divergent biological mechanisms, such as mitochondrial dysfunction). Different study designs may serve different purposes.

Design of subtyping studies fell into two main categories: 1) data-driven analyses and 2) hypothesis-driven analyses based on pre-determined groups. Hypothesis-driven studies were usually single domain analyses of a clinical or demographic characteristic such as age of onset, or tremor-dominant vs. akinetic-rigid (‘motor subtyping’), and this was felt to be an inferior approach compared with considering multiple domains of data in a hypothesis-free subtyping exercise. However, it may be that focused hypothesis-driven evaluation of pre-defined groups is important for answering specific research questions, and this approach may have more potential for direct neurobiological mapping with concomitant biomarker measurement. For example, this approach has successfully identified biological differences between individuals based on motor phenotype [12, 13]. Findings from data-driven studies should be similarly used for hypothesis-driven studies to define the clinical applicability and/or biological underpinnings of the described subtypes. Importantly, clinical applicability may not be a relevant objective of some subtyping efforts.

Subtyping studies have usually enrolled individuals at various points in the disease course and analyzed the cohort without stratification on disease duration or stage. This approach makes it more difficult to describe phenotypic variability at specific phases of the disease. In addition, patients may be misclassified if individuals at different disease durations are included without taking the phenotypic changes with disease duration into account. Certain aspects such as motor phenotype tend to be more heterogeneous early in the disease process, converging to a common phenotype toward the end [14, 15]. Thus, subtyping exercises may produce very different results depending on the distribution of disease duration or stage at which the subtyping is performed. We would recommend subtyping based on
longitudinal data starting from a defined disease duration or milestone in order to maximize interpretability and minimize these confounding issues.

The included studies were all clinic-based, rather than recruiting from community sources. This may be a more or less important source of bias depending on the purpose of the subtyping; individuals from these two sources are unlikely to differ in the underlying biology of the disease but may well differ in clinical severity and prognosis. A community-based sample would be more generalizable.

What is the clinical and biological relevance of subtypes?

The clinical importance of the differences between the defined subtypes was deemed moderate or less for the vast majority of studies. Whether or not the proposed subtypes are stable, or become more or less different over time, could not be evaluated in this review for most classification systems since longitudinal differences were not assessed in the original descriptions of the subtypes. These limitations raise important questions about the clinical relevance of subtypes defined to date, with the caveat that we have not evaluated the literature following up the initial descriptions of the subtypes. We are aware of many studies evaluating subtypes defined by motor features contrasting tremor-dominant (TD) with other phenotypes. Some have looked at the prognostic implications of these subtypes [16, 17] and several studies have demonstrated that a high proportion of individuals with the TD phenotype will switch to a PIGD phenotype over time [14, 15, 18]. An adverse cognitive prognosis and prominent cortical Lewy body involvement associated with the non-tremor dominant phenotypes confirms the prognostic relevance of those subtypes [19]. Recently, there has also been an independent examination of the clinical and pathological evolution of data-driven subtypes as defined by an earlier cohort study [4]. Important differences in time to important clinical milestones between the subgroups were shown in a separate cohort, supporting the clinical importance of those subtypes [20]. Unfortunately, such studies are rare and we encourage such prognostic studies to clarify the clinical relevance of any described subtypes.

The relationship between clinically-based subtypes and the underlying biology of disease has rarely been assessed to date. Several studies have evaluated the biological characteristics of TD and PIGD phenotypes, finding differences in CSF composition (alpha-synuclein [21], 5-hydroxyindoleacetic acid, glycine [12]) and degree of cardiac sympathetic denervation, for example [21]. We are aware of only two studies evaluating the biomarker profile of data-driven subtypes, identifying a possibly pro-inflammatory profile [9] and an “Alzheimer’s-like” profile of low CSF amyloid-β and high tau [4] associated with a severe motor and non-motor disease subtype. In the absence of validated non-clinical biomarkers of PD progression, the value of clinically-based subtyping to help direct disease-modifying therapies targeting specific biological processes is unknown. Establishing the relationship between clinically-based subtypes and biomarkers of the underlying disease biology is an important goal, as it may inform the development of subtype-targeted therapies.

We also found that there was little inclusion of biomarkers in studies defining PD subtypes. This has the same implications as mentioned above, limiting our understanding of the biological relevance of subtypes. We further recommend biologically-based subtyping studies.

Can subtyping be incorporated into clinical practice?

Clinical applicability may not be the goal of all subtyping efforts, such as those seeking to understand heterogeneity in the underlying biology of the disease. Therefore, clinical applicability may not be a relevant criterion on which to rate some studies. Nonetheless, it is relevant to note that our expert group had reservations about the feasibility of subtyping as part of routine clinical practice with the tools currently available. That conclusion was driven by 1) the extra time required to assign individuals to subtypes using multi-domain data, beyond that allocated to usual clinic visits and 2) the lack of an algorithm to classify individual patients. An advantage of hypothesis-based single-domain subtyping systems is the availability of a clear and simple algorithm to assign individuals to groups, facilitating both replication of findings in different cohorts [20] and application in a clinical setting. It is rare that data-driven studies use their results to derive an algorithm for assigning individuals to subtypes, although there is at least one exception [4, 20]. Even if feasible, the applicability of the results to individuals is unclear given that subtyping has so far been studied only at the group level. Studies examining the prognostic or biological relevance at the individual level are needed to
establish clinical applicability. Furthermore, simple classification systems based on a manageable number of variables would be a prerequisite for adoption by the clinical community; however, simplicity is at odds with the fact that PD is etiologically and phenomenologically complex, and being a disease associated with aging, subject to multiple intersecting biological processes [22] and co-existing medical comorbidities [23] that can intervene to influence the phenotype at any point in the disease process.

Is subtyping possible?

Subtyping studies undertaken to date have significant methodologic shortcomings and most have questionable clinical applicability. Many studies were limited in terms of the clinical domains considered to define subtypes, and measures used in data-driven analyses were highly heterogeneous, resulting in highly variable findings. Even the two data-driven studies receiving the highest quality ratings and incorporating variables from comparable domains showed limited similarity of the subtypes they found. Despite decades of subtyping research, there has been minimal incorporation of subtype information into research or formal incorporation in clinical practice. These observations call into question the feasibility of clinical subtyping and suggest that alternative approaches to describing and understanding the heterogeneity of PD are needed.

As described above, PD is biologically complex, a result of many intersecting processes that vary from person to person. Blood and cerebrospinal fluid parameters associated with PD show high inter-individual heterogeneity, requiring combinations of multiple markers to optimize diagnostic accuracy [24]. Variable co-pathology adds to the heterogeneity [25]. Several ‘causative’ genes, none completely penetrant, and a vast array of susceptibility genes governing widely-varying metabolic processes have been elucidated [26]. RNA expression studies demonstrate the complex genetic and biologic heterogeneity of PD [27]. It is a testament to this complexity that metabolomics are now being used to measure the downstream effects of a vast array of contributory genetic, environmental and physiological processes [28]. As a result, the phenotype of PD is unique to an individual. Given these challenges, it seems unlikely that a purely clinically-based subtyping system measuring early disease features and seeking to place patients into a small number of categories will be able to adequately describe PD heterogeneity in order to provide accurate pathophysiological or prognostic insights. It remains to be seen whether this can be improved by the discovery of better biomarkers but underlying biological heterogeneity may also render biomarker-based subtyping resistant to meaningful cluster solutions. This idea is not new; it has been previously proposed that PD is not one disease but rather a syndrome representing the manifestations of multiple or even an infinite number of underlying diseases [29].

Contemporary medicine is increasingly promoting personalized treatment, including in Parkinson’s disease, with a focus on the individual [30]. To date, subtyping places individuals in groups with similar but not identical features. This may represent an important step toward identifying individuals that can respond preferentially to certain treatments but by virtue of placing individuals within a group will inevitably fall short of the truly ‘personal’ goal. Although perhaps more challenging, modern computational techniques are increasingly allowing us to manage vast amounts of data and may soon allow us to take full advantage of the information available in the heterogeneity to describe individuals without the need for group-level subtyping. Many of the recommendations outlined above could apply to future studies where the unit of measure is the individual’s disease fingerprint rather than the group phenotype. Granted, such an individual approach poses financial and logistical challenges when it comes to clinical trials which will have to be overcome. Nonetheless, having reviewed the existing literature on subtyping and explored the methodologic pitfalls and challenges associated with performing the optimal subtyping studies described above, it is time to re-evaluate our approach to understanding and describing PD heterogeneity.

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CONFLICT OF INTEREST

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SUPPLEMENTARY MATERIAL

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REFERENCES


