Gastrointestinal syndromes preceding a diagnosis of Parkinson’s disease: testing Braak’s hypothesis using a nationwide database for comparison with Alzheimer’s disease and cerebrovascular diseases

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ABSTRACT
Objective Braak’s hypothesis states that Parkinson’s disease (PD) originates in the gastrointestinal (GI) tract, and similar associations have been established for Alzheimer’s disease (AD) and cerebrovascular diseases (CVD). We aimed to determine the incidence of GI syndromes and interventions preceding PD compared with negative controls (NCs), AD and CVD.

Design We performed a combined case-control and cohort study using TriNetX, a US based nationwide medical record network. Firstly, we compared subjects with new onset idiopathic PD with matched NCs and patients with contemporary diagnoses of AD and CVD, to investigate preceding GI syndromes, appendectomy and vagotomy. Secondly, we compared cohorts with these exposures to matched NCs for the development of PD, AD and CVD within 5 years.

Results We identified 24 624 PD patients in the case-control analysis and matched 18 cohorts with each exposure to their NCs. Gastroparesis, dysphagia, irritable bowel syndrome (IBS) without diarrhoea and constipation showed specific associations with PD (vs NCs, AD and CVD) in both the case-control (odds ratios (ORs) vs NCs 4.64, 3.58, 3.53 and 3.32, respectively, all p<0.0001) and cohort analyses (relative risks (RRs) vs NCs 2.43, 2.27, 1.17 and 2.38, respectively, all p<0.0001). gastroparesis, dysphagia, constipation and faecal incontinence were not PD specific, IBS with constipation and intestinal pseudo-obstruction showed PD specificity in the case-control (OR 4.11) and cohort analysis (RR 1.84, respectively). Appendectomy decreased the risk of PD in the cohort analysis (RR 0.48). Neither inflammatory bowel disease nor vagotomy were associated with PD.

Conclusion Dysphagia, gastroparesis, IBS without diarrhoea and constipation might specifically predict Parkinson’s disease.

INTRODUCTION
Parkinsonism is a clinical syndrome characterised by bradykinesia, rest tremor, rigidity and postural instability.1 Its most common cause is Parkinson’s disease (PD), the pathological hallmark of which is thought to be cytoplasmatic eosinophilic Lewy body (LB) depositions. These depositions, mainly consisting of misfolded α-synuclein, have not only been found in the CNS but also in the vagus nerve and enteric nervous system (ENS) of patients with PD.2

These findings led Braak et al to state the neuro-anatomical hypothesis that α-synuclein pathology progresses from peripheral sites such as the ENS to the CNS via vagal or olfactory pathways, thereby introducing the concept that the gastrointestinal (GI) tract might serve as a gateway for environmental factors that induce α-synuclein misfolding and lead to PD.3 A large body of evidence has since then accumulated to support this claim. Even in early untreated stages of the disease, neuropathological studies have found that α-synuclein concentrations in the ENS of patients with PD were higher than those of otherwise healthy individuals, in a
characteristic rostrocaudal gradient following visceromotor projections of the vagus nerve. Complementary studies have shown that various motility disorders and inflammatory bowel disease (IBD) can precede PD and therefore may be risk factors for its development. Moreover, since Gray and colleagues first identified the vermiform appendix as a potential source of misfolded α-synuclein, conflicting observational studies have been published about the impact of appendectomy on the risk of idiopathic PD. Finally, two recent registry based studies have strengthened the concept of retrograde vagal α-synuclein propagation by showing that a truncated vagotomy might be protective against the development of PD. Apart from the bottom-up link formulated by Braak, a top down aetiology in which GI symptoms are present in early phases when neurological manifestations are still unnoticed is also supported by experimental evidence. Even if no causal link exists, GI syndromes might still represent a risk factor through other mechanisms, or both might be related to a yet unknown third factor.

Apart from PD, other neurological disorders have also been hypothesised to have GI precedents, either through similar neuroimmune pathways or translocation of microbiome derived neurotoxins into the CNS. A strong pathological link between neuroimmune pathways or translocation of microbiome derived neurotoxins into the CNS is supported by experimental evidence. Even if no causal link exists, GI syndromes might still represent a risk factor through other mechanisms, or both might be related to a yet unknown third factor.

Methods

Study design and data source

To investigate the association between various GI syndromes and interventions with the subsequent development of new onset PD, we analysed electronic medical records from the TriNetX Analytics Research Network (Cambridge, Massachusetts, USA). At the moment of data collection, the network consisted of more than 80 million patients from 57 predominantly academic medical centres in the USA. Additional information can be found in the online supplemental methods.

Study population and variables of interest

In the case control analysis, we examined the incidence of exposures retrospectively (ie, before an initial diagnosis of PD compared with matched controls). Patients with PD were captured using a previously validated method. Patients were queried using the International Statistical Classification of Diseases and Related Health Problems, 10th revision (ICD-10) diagnosis of PD (G20), if documented between 1 January 2005 and 1 July 2021; the first ever diagnosis was used as the index event. Only those with at least two prescriptions of an antiparkinsonian drug and a documented ambulatory visit at least 2 years before the first diagnosis of PD were included; secondary causes of PD were excluded. To determine PD specific exposures, control subjects comprised three groups: NCs, and patients with a diagnosis of AD and CVD. NCs consisted of patients without a recorded ICD-10 diagnosis of PD, with at least two documented ambulatory visits between the ages of 50 and 90 years, at least 2 years apart, recorded between 1 January 2005 and 1 July 2021. A minimum of 2 years of retrospective follow-up was ensured by using the second of these visits as the index event. Similarly, 2 years of follow-up was ensured for the AD and CVD groups, and the first ever documented respective ICD-10 diagnosis in the medical records after 1 January 2005 was chosen as the index event. In a pairwise fashion, these groups where then matched to the PD group for age, sex, race and ethnicity using a propensity score matching algorithm.

To cover the entirety of the GI tract, 18 exposures were investigated: achalasia, dysphagia, gastro-oesophageal reflux disease (GORD), gastroparesis (GP), functional dyspepsia (FD), paralytic ileus (PI), diarrhoea, irritable bowel syndrome (IBS) with constipation (IBS-C), IBS with diarrhoea (IBS-D), IBS without diarrhoea, intestinal pseudo-obstruction (approximate synonym of K59.8: other specified functional GI disorders), faecal incontinence (FI), Crohn’s disease (CD), ulcerative colitis (UC), microscopic colitis (MC), appendectomy and vagotomy. We conducted additional sensitivity analyses in the network, which included stratified analyses based on sex and age at the diagnosis of the index event. A detailed breakdown of the query, inclusion and exclusion criteria, stratified analyses and coding can be found in the online supplemental methods.

To validate the results from the case-control analyses, we set up a complementary cohort study design. Eighteen cohorts, each diagnosed with one of the investigated exposures in the case-control analysis, were queried and compared with a respective NC cohort (ie, without the exposure) for the prospective risk of developing PD, AD or CVD. Only those with at least 5 years of prospective follow-up were included, and were propensity score matched for age, sex, race and ethnicity, and additionally for a set of potential risk factors and risk modifiers for the development of PD, AD and CVD: arterial hypertension, diabetes mellitus, atrial fibrillation and flutter, and nicotine dependence.

Statistical analysis

In the case-control analyses, patients were counted as positive for an exposure if the respective ICD-10 code was documented any time before the first diagnosis of PD or the control health event. To approximate the diagnostic interval between each exposure and the first PD diagnosis, a yearly cross sectional prevalence for each exposure was calculated for the PD and NC groups, up to 6 years before the index event. To detect and quantify potential surveillance bias in our case-control analyses, we collected an agnostic set of negative exposures (Charlson comorbidities). This allowed us to determine the OR that should be considered as indicative of no association. Additionally, we collected positive exposures based on a previous case-control study that identified prodromal motor and non-motor symptoms of PD. This enabled us to assess the ability of our dataset to reproduce existing associations. The coding can be found in the online supplemental methods.

In the cohort analyses, patients diagnosed with the exposure of interest and their NCs were counted as positive for an outcome
(PD, AD or CVD) if the respective new onset ICD-10 diagnosis occurred within a 5 year follow-up. Subjects who already had the outcome of interest before the index event were excluded after propensity score matching.

Exposures and outcomes were collected as absolute numbers; ORs and RRs were calculated with 95% CIs. Standardised mean differences (SMDs) were used to compare baseline characteristics; an SMD of <0.2 was considered well balanced. A Pearson \( \chi^2 \) test was calculated to compare outcomes, and a two sided P value of <0.05 was used to indicate statistical significance. Correction for false discovery rate (FDR) was performed using the step up procedure by Benjamini and Yekutieli, with the Stats package in R (V.4.3.0).25

**RESULTS**

For the case-control study, 24 624 patients with PD met all of the criteria and were matched with 8 267 744 NCS, and 36 187 AD and 528 207 CVD patients, giving 24 624 patients in the comparison with NCS, 19 046 with AD and 23 942 with CVD. Baseline characteristics after pairwise matching are presented in table 1; minimal differences in age at index persisted. SMDs and p values before and after matching can be found in online supplemental tables 1,2.

The results of the case-control analyses are presented in figure 1 and online supplemental table 3. All GI syndromes were significantly increased in the PD group compared with NCSs (OR >1; p<0.05). However, only dysphagia (OR 3.58), GP (OR 4.64), FD (OR 3.39), intestinal pseudo-obstruction (OR 3.01), diarrhoea (OR 2.85), constipation (OR 3.32), IBS-C (OR 4.11), IBS-D (OR 4.31), IBS without diarrhoea (OR 3.53) and FI (OR 3.76) gave ORs that were numerically greater than the upper limit of what is expected by surveillance bias (OR range 1.20–2.79). Only for GORD and appendectomy we observed significant differences compared with PD with NCSs, but gave ORs below the upper limit of what is expected by surveillance bias (OR range 1.20–2.79). Only for GORD and appendectomy we observed significant differences compared with AD (OR 1.14, p<0.0001) and CVD (OR 0.57, p=0.03), respectively. The latter did not remain significant after correction for FDR (p=0.21). Prior vagotomy did not impact the risk of PD.

For GP, women were approximately twice as likely as men to develop PD (OR 7.3 for women, 3.05 for men, both p<0.0001 (online supplemental tables 7–9 and online supplemental figure 4), and the OR for GP was especially high for early onset PD (online supplemental tables 10–11). Exclusion of previous anti-dopaminergic drug use did not significantly alter any associations with PD compared with NCSs (online supplemental table 12 and online supplemental figure 5). The ORs of all PD specific exposures were positioned well within the range of those of established motor and non-motor prodromes of PD (ie, positive exposures; OR PD vs NCSs 2.04–7.41). An approximation of the diagnostic interval for each exposure can be found in online supplemental figure 6.

**Table 1 Baseline characteristics for subjects with PD and controls in the case-control analyses after pairwise matching.**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Patients (n (%))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parkinson’s disease</td>
<td>Negative controls</td>
</tr>
<tr>
<td>(n=24 624)</td>
<td>(n=24 624)</td>
</tr>
<tr>
<td>Parkinson’s disease</td>
<td>Alzheimer’s disease</td>
</tr>
<tr>
<td>(n=19 046)</td>
<td>(n=19 046)</td>
</tr>
<tr>
<td>Age</td>
<td>70.8 (8.49)</td>
</tr>
<tr>
<td>Age (years) (mean (SD))</td>
<td>70.4 (8.57)</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
</tr>
<tr>
<td>Men</td>
<td>14254 (57.89)</td>
</tr>
<tr>
<td>Women</td>
<td>10103 (41.03)</td>
</tr>
<tr>
<td>Unknown</td>
<td>267 (1.08)</td>
</tr>
<tr>
<td>Race</td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>20476 (83.15)</td>
</tr>
<tr>
<td>Black or African American</td>
<td>1294 (5.26)</td>
</tr>
<tr>
<td>Other and unknown†</td>
<td>2854 (11.59)</td>
</tr>
<tr>
<td>Ethnicity</td>
<td></td>
</tr>
<tr>
<td>Hispanic or Latino</td>
<td>939 (3.81)</td>
</tr>
<tr>
<td>Not Hispanic of Latino</td>
<td>19746 (80.19)</td>
</tr>
<tr>
<td>Unknown</td>
<td>3939 (16)</td>
</tr>
</tbody>
</table>
| *Characteristics were identified using electronic medical health record data from the TriNetX Research Network. Baseline characteristics before matching can be found in online supplemental table 1, and p values and standardised mean differences between groups in online supplemental table 2. †Includes Asian, American Indian, Alaska Native, Native Hawaiian or other Pacific Islander, or unknown.
Figure 1 Case-control analyses. Odds ratios (ORs) of previous exposures in patients with Parkinson’s disease (PD) compared with matched negative controls (NCs), or patients with Alzheimer’s disease (AD) and cerebrovascular diseases (CVD). In the case-control analysis, the incidence of exposures was examined retrospectively (ie, before an initial diagnosis of PD compared with matched NCs, and AD and CVD patients). Patients were diagnosed with PD or the respective control health event between 1 January 2005 and 1 July 2021, and NCs were queried using two ambulatory visits during the same time window. Patients and exposures were identified with the International Statistical Classification of Diseases and Related Health Problems, 10th revision (ICD-10) codes, and Current Procedural Terminology (CPT) codes, using electronic medical health record data from the TriNetX research network (online supplemental methods). Exposures were included if they were documented any time before the diagnosis of PD or the control health event, in the available medical records. ORs were calculated as follows: odds of documented exposure in the PD cohort/odds of documented exposure in the control cohort. Absolute rates can be found in online supplemental table 3. \(^{P}\) values were calculated using a Pearson \(\chi^2\) test, after matching for baseline characteristics (ie, age, sex, race and ethnicity). Correction for false discovery rate was performed using the step up procedure by Benjamini and Yekutieli, with the Stats package in R. \(^{*}\) The term intestinal pseudo-obstruction was used as an approximate synonym for the ICD-10 code K59.8 other specified functional intestinal disorders. \(^{**}\) The term IBS not otherwise specified is commonly used as an approximate synonym for the ICD-10 code K58.9 IBS without diarrhoea. IBS, irritable bowel syndrome.
Gut brain axis

Figure 2  Cohort analyses. Relative risks (RRs) of developing Parkinson’s disease (PD), Alzheimer’s disease (AD) or cerebrovascular diseases (CVD) within 5 years of the diagnosis of a given exposure, compared with negative control (NCs) without the respective exposure. *For each analysis, a cohort of patients identified by the diagnosis of a given exposure was compared with their respective NCs for the prospective risk of PD, AD and CVD within 5 years of the index event (ie, diagnosis of the given exposure, or a visit for NCs). After propensity score matching, patients that already had the investigated outcome (ie, PD, AD or CVD) documented before the index event were excluded from the analysis. Exposures and outcomes were identified using the International Statistical Classification of Diseases and Related Health Problems, 10th revision (ICD-10,) and Current Procedural Terminology (CPT) codes. Electronic medical health record data were collected from the TriNetX research network. Diagnostic coding can be found in the online supplemental methods. **RRs were calculated as follows: risk of outcome in the exposure cohort/risk of outcome in the control cohort. Absolute rates can be found in online supplemental table 13. *P values were calculated with a Pearson $\chi^2$ test, after matching for baseline characteristics and risk factors. Baseline characteristics included age, sex, race and ethnicity; risk factors included arterial hypertension, diabetes mellitus, atrial fibrillation and flutter, and nicotine dependence. Correction for false discovery rate was performed with the step up procedure by Benjamini and Yekutieli, using the Stats package in R. The term intestinal pseudo-obstruction was used as an approximate synonym for the ICD-10 code K59.8 other specified functional intestinal disorders. **The term IBS not otherwise specified is commonly used as an approximate synonym for the ICD-10 code K58.9 IBS without diarrhoea. IBS, Irritable bowel syndrome.
CVD, and statistical significance (p<0.05) was achieved only for constipation and dysphagia (online supplemental table 15). In line with the case control analyses, FI equally increased the risk of PD and AD (PD: RR 1.74; AD: RR 1.76, both p<0.0001), and in the GORD cohort the RR of developing CVD was higher than that of PD (CVD: RR 1.38; PD: RR 1.13, both p<0.0001). FD and diarrhoea were associated with all three disorders (p<0.05), and PI, CD and MC only increased the risk of CVD (RR 1.22, p<0.0001; RR 1.20, p<0.0001; RR 1.26, p=0.02, respectively). Although no specific association was found in the case-control analysis, appendectomy significantly reduced the risk of developing PD (RR 0.48, p=0.05), which did not remain significant after correction for FDR. For all other exposures (ie, achalasia, UC, IBS-C, IBS-D and vagotomy), no significant associations were found.

**DISCUSSION**

We used a nationwide EHR network to comprehensively investigate disorders across the entire GI tract before a diagnosis of PD. We used two complementary study designs to establish that dysphagia, gastroparesis, constipation and IBS without diarrhoea specifically increase the risk of a subsequent new onset diagnosis of idiopathic PD, even compared with other neurological diseases, such as AD and CVD.

Surveillance bias is an inherent problem in observational studies. When not addressed appropriately, it can compromise the validity of causal inference and lead to irreproducible results. A broader implementation of empirical approaches to evaluate and correct for the presence of systematic error in observational studies is needed. Therefore, we set up an approach to understand the true extent of surveillance bias in our study and its potential contribution to implicating premorbid factors for PD. Hence we collected data on all diagnoses included in the Charlson comorbidity index. These premorbid conditions were considered as a comprehensive set of agnostic negative exposures. In our case-control study, we observed statistically significant increases in most of these exposures in PD cases compared with NCs, but not compared with AD and CVD (online supplemental figure 1). To establish whether these increases represented surveillance bias or true associations (although unlikely based on the current literature), we investigated the same exposures for other neurological disorders (AD and CVD) compared with their NCs (online supplemental figure 3). Since the same significant correlations emerged, surveillance bias was likely. Subsequently, we determined the range of ORs that should be expected if they are a result of surveillance bias alone. These ORs ranged between 1.20 and 2.79 in the analysis of PD with NCs. To determine whether existing associations with PD could be replicated with ORs greater than those of negative exposures, we also collected prodromal motor and non-motor symptoms of PD (ie, positive exposures). These resulted in ORs ranging between 2.04 and 7.41 (online supplemental figure 1).

Having established a measure of surveillance bias in the case-control study, we then determined ORs for the putative GI pre-/comorbidities of PD. Relative to the upper limit of the negative exposures, GI exposures fell into two categories. First were those for which the ORs overlapped with the ORs expected for surveillance bias (OR 1.20–2.79). For these exposures, including achalasia, GORD, PI, IBD (ie, CD, UC, and MC), appendectomy and vagotomy, we cannot be confident that these were true associations, although within the constraints of our study we cannot categorically state that they were not. Second were those for which the ORs were clearly higher than the ORs expected for surveillance bias (OR >2.79). For these exposures, including dysphagia, GP, FD, intestinal pseudo-obstruction, diarrhoea, constipation, IBS-C, IBS-D, IBS without diarrhoea and FI, we can confidently state to have established significant associations with new onset PD. To determine the specificity of the identified significant exposures for PD in the case-control analyses, we subsequently compared subjects with PD with subjects with AD and CVD. Only dysphagia, GP, constipation, IBS without diarrhoea and IBS-C remained specific for PD compared with both neurological diseases (table 2). Importantly, we cannot exclude the possibility that these factors might still be associated with these diseases, although at a smaller scale. Similarly, while other exposures were not specific for PD (ie, FD, intestinal pseudo-obstruction, diarrhoea, IBS-D and FI), we cannot strictly exclude the possibility that these conditions might still be risk factors for PD.

Finally, to validate these findings both in terms of their significance and specificity and establish one RR estimate of developing PD after the diagnosis of each exposure, we set up a complementary cohort study. Here, five exposures (ie, dysphagia, GP, IBS without diarrhoea, intestinal pseudo-obstruction and constipation) significantly increased the risk of PD and resulted in RRs that were numerically greater than those of AD and CVD.
The positive predictive value of ICD-high (80–95%), their accuracy in indicating specific subtypes was 1.05 to 1.3). Specifically increased in the case-control analysis, only if intestinal LB pathology becomes important to determine the presence of LB pathology in PD.\(^3^2\)\(^3^3\) Interestingly, the highest RR was observed for GP (RR \(2.38, 95\%\) CI 1.18 to 2.87). This disorder for evaluating enteric involvement in PD. Although anorectal symptoms are among the most frequent GI symptoms in PD, our data suggest that the presence of FI might not distinguish between the development of PD and other neurodegenerative diseases.\(^4^3\)\(^4^4\) Even if not prodromal to AD, our findings support the fact that the progression of cognitive decline in AD is frequently unmasked by FI.\(^4^5\) While diarrhoea and PD increased the risk of all three diseases, intestinal pseudo-obstruction showed specificity for PD in the cohort analyses (RR 1.84, 95% CI 1.18 to 2.87). This disorder, characterised by impaired peristalsis and presumably caused by a neuropathy or myopathy, has been described in various neurological disorders, including PD.\(^4^5\)

Finally, some exposures in the case-control analyses gave ORs in the range of those expected from surveillance bias. These exposures included achalasia, GORD, PI, IBD (ie, CD, UC, and MC), appendectomy and vagotomy. Other than for GORD, the cohort analyses also did not show any significant associations with PD for these exposures. While IBD has been linked to PD in various observational\(^4^6\)\(^4^7\) and genetic studies,\(^4^8\) neither of our study designs supported this link. However, we cannot strictly dismiss the possibility of an association based on this empirical surveillance bias cut-off and a relatively limited follow-up. In addition, we were unable to assess the impact of anti-tumour necrosis factor (anti-TNF) therapy exposure, which has been hypothesised to decrease the risk of PD.\(^4^9\) Prospective studies are necessary to investigate this association and to establish whether anti-TNF therapy can effectively protect against PD. Notably, concordant with an earlier study that linked reflux oesophagitis to an increased risk of stroke and transient ischaemic attack in patients with atrial fibrillation,\(^5^0\) the risk of CVD in our study was significantly greater after a diagnosis of GORD, CD and MC. These findings suggest that a better understanding of the link between GI inflammation and cerebrovascular events may lead to improved risk stratification and identification of new preventive strategies.\(^5^1\)

After Grey et al first discovered that α-synuclein was most abundant in the appendicular mucosa,\(^5^2\) conflicting evidence has emerged about the impact of an appendectomy on PD risk. While three studies did not find any association,\(^1^1\)\(^1^4\)\(^3^0\) one abstract reported an increased risk of PD\(^1^3\) and two large observational studies supported a protective effect.\(^1^0\)\(^1^2\) Despite our limited follow-up and sample size compared with the aforementioned studies, we observed a relative risk reduction of 52% in our cohort analysis, while the case-control analysis was likely underpowered to detect any consistent association for appendectomy. Multiple studies suggest that the appendix constitutes a prominent source of seeding competent pathologically folded α-synuclein,\(^1^0\) and houses bacteria capable of releasing inflammatory mediators.\(^1^0\) The subsequent migration of α-synuclein to the CNS has been substantiated by studies showing a protective effect of a truncal vagotomy on PD development.\(^1^5\)\(^1^6\) Compared with these reports, our study was underpowered to detect any consistent associations for vagotomy.\(^1^5\)\(^1^6\)

Finally, we attempted to assess the proximity of each diagnosis to the diagnosis of PD in the case-control study (online supplemental figure 5). We found that the OR for dysphagia and constipation decreased considerably as the distance from the diagnosis of PD increased, while the OR for GP and IBS without diarrhoea remained relatively constant. This suggests that differences in lead time exist, but future longitudinal population-based studies will be crucial to determine whether these PD specific GI syndromes are part of the early manifestation of PD or truly precede the disease. Importantly, the combination of two complementary study designs reduced the potential for selection bias. The case-control analysis consisted of patients with PD, AD or CVD without the requirement of any previous exposure, while the cohort analyses consisted of patients with newly diagnosed GI exposures without the requirement of a subsequent PD, AD or CVD diagnosis. This study is subject to intrinsic limitations of EHR data, including unknown completeness of records and absent validation of diagnoses. The multi-centre character and inclusion of racially and ethnically diverse subjects ensured that these results are generalisable to patients at academic medical centres across the USA.
CONCLUSION
This study is the first to establish substantial observational evidence that the clinical diagnosis of not only constipation, but also dysphagia, GP and IBS without diarrhoea might specifically predict the development of PD, whereas other exposures were less specific. An appendectomy appeared protective, leading to further speculation about its role in PD pathophysiology. These findings warrant alertness for GI syndromes in patients at higher risk for PD and highlight the need for further investigation of GI precedents in AD and CVD. To establish a stronger body of clinicopathological evidence, we advocate for future studies to assess the sensitivity and specificity of these disorders and their clinicopathological correlates for the early detection of neuropathology.

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Contributors
BK: conceptualisation, methodology, formal analysis, data curation, validation, visualisation, writing-original draft, and writing-review and editing. LV: conceptualisation, resources, and investigation. JV: conceptualisation, methodology, and visualisation. GB: conceptualisation, resources, and investigation. RB: conceptualisation, methodology, validation. Y: funding agency in the public, commercial or not-for-profit sector. MM: conceptualisation.

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Patient and public involvement
Patients and/or the public were not involved in the design, conduct, or reporting, or dissemination plans of this research.

Patient consent for publication
Not applicable.

Provenance and peer review
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Data availability statement
Data are available upon reasonable request. Data exported from TriNetX were saved in Excel files and archived. Every co-author affiliated to Johns Hopkins University was granted access to the TriNetX Research network by the Institute of Clinical and Translational Research (ICTR). The online supplemental materials contain an extensive list of tables representing the original data; researchers will be granted access to the original aggregated data upon reasonable request, with agreement of the corresponding author (PJP).

Supplemental material
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