Trends and Contributing Factors for Prescribing Antipsychotics in Newly Diagnosed Parkinson’s Disease Patients: A Population-Based Study

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Purpose: Psychosis is very common in later stages of Parkinson’s disease (PD) patients, but its prevalence in newly diagnosed patients is not rare. The use of antipsychotics in PD patients is complex given their association with worsening Parkinsonian motor symptoms and increased mortality rate. This study aims to examine factors that affect the use of antipsychotics in newly diagnosed PD patients and to identify changes in prescribing over time.

Patients and Methods: The Secure Anonymized Information Linkage (SAIL) databank was used to identify a cohort of newly diagnosed PD patients aged 40 years and older in Wales. The cohort was longitudinally examined over 17 years to determine the incidence of new antipsychotic use. Logistic regression models were used to analyze the data and were adjusted for several potential confounding variables.

Results: A total of 9142 PD patients were identified after applying inclusion and exclusion criteria, of whom 58.6% were male. During the first year of PD diagnosis, 12% of the patients developed psychosis and were prescribed antipsychotics. Quetiapine was the most commonly prescribed (49%), followed by risperidone (10.7%). The use of antipsychotics in newly diagnosed PD patients was significantly lower in the years 2009–2016 compared to 2000–2008 (OR 0.37, 95% CI 0.32–0.43). Other significant prescribing factors included patient’s age and history of dementia.

Conclusion: A dramatic decline in antipsychotic use was identified across years, showing adherence to warnings against use of antipsychotics for PD patients. Given that psychosis is prevalent in PD patients, the continuous assessment of the safety risks of antipsychotics is a matter of priority.

Keywords: Parkinson’s disease, psychosis, pattern of prescribing, mortality risk

Introduction
Psychosis is very common in Parkinson’s disease (PD) patients. It is estimated that psychosis (visual hallucinations and delusions) affect 40% of PD patients. Although new reports suggest an early emergence of minor hallucinations in PD patients even before the occurrence of motor symptoms, significant psychosis is not common in the first stages of PD; rather, most patients develop psychosis in the advanced stages of PD. Psychotic episodes in PD may occur as a result of the disease itself, or they may be caused by some PD medications, such as dopamine agonists (DAs). Therefore, it is important when treating psychosis to first manage the PD medications and doses, especially in early stages of the disease, and then...
consider antipsychotic medications if needed.\textsuperscript{1} Episodes of hallucinations are more common in the late stages of PD and are the main predictor of nursing home placement in PD patients.\textsuperscript{6, 6}

The use of antipsychotics in PD patients is complex, since antipsychotics block D2 dopaminergic receptors, which may lead to extrapyramidal side effects such as dyskinesia and Parkinson-like motor symptoms.\textsuperscript{7} The second-generation antipsychotics (SGAs), also known as atypical antipsychotics, are safer and cause fewer Parkinson-like symptoms compared to the first generation (FGAs) or typical antipsychotics.\textsuperscript{8} Patients who use typical antipsychotics are more likely to receive PD medications (to control Parkinson-like symptoms) compared to atypical antipsychotic users.\textsuperscript{9} In general, and particularly among atypical antipsychotics, clozapine and quetiapine are recommended for use in PD patients with psychosis.\textsuperscript{10} The efficacy of quetiapine has not been confirmed in randomized clinical trials; nevertheless, it has been suggested that quetiapine should be used first in PD psychosis.\textsuperscript{10} Clozapine has an approved indication for treating psychosis in PD patients and it lacks the Parkinson-like symptoms;\textsuperscript{11} however, it is considered to be a second-line treatment after quetiapine due to its risk of agranulocytosis and its requirement for continuous blood monitoring.\textsuperscript{10}

Parkinson-like symptoms are not the only safety issue related to the use of antipsychotics. Based on several reports, the US Food and Drug Administration (FDA) issued a warning in 2005 regarding the risk of death associated with the use of atypical antipsychotics in patients with dementia.\textsuperscript{12} This warning was extended to include the typical antipsychotics in 2008.\textsuperscript{13} Based on these reports, the Banerjee report was published in the UK in 2009 and called for immediate action to reduce the amount of antipsychotics prescribed to dementia patients and to identify ways to manage this issue.\textsuperscript{14} Recent studies have extended this warning of higher mortality rate to use of antipsychotics in PD patients.\textsuperscript{15, 16} In these studies, typical antipsychotics were more likely to cause death compared to atypical antipsychotics.\textsuperscript{15, 16} It is important to recognize that the high mortality among PD patients who used antipsychotics is not necessarily caused by the use of antipsychotics. Other factors can contribute to increased mortality rates such as greater age, being male, and psychotic episodes themselves,\textsuperscript{17} which makes attributing the higher mortality rate to antipsychotics alone a matter of controversy.

Some drugs used in the treatment of dementia, such as rivastigmine (a cholinesterase inhibitor), show some evidence of hallucination reduction and improved psychotic symptoms in PD patients, although rivastigmine has a slower response compared to atypical antipsychotics.\textsuperscript{18} The recent National Institute for Health and Care Excellence (NICE) guidelines did not recommend rivastigmine as part of therapy; however, further research was recommended to compare rivastigmine with atypical antipsychotics in managing psychosis in PD patients.\textsuperscript{19}

The prevalence of antipsychotic use in PD patients has been measured in multiple studies.\textsuperscript{20–22} Many of these studies focused on psychosis in the later stages of PD, and concluded that antipsychotic use was high and should be decreased.\textsuperscript{20–22} However, psychosis in early stages of PD stages is not rare,\textsuperscript{3} and the use of antipsychotics during this stage is under researched and needs more attention from researchers. If reducing antipsychotic use in the later stages of PD is desirable,\textsuperscript{2} then reducing use in the early stages of PD is a priority.

The primary objective of this study was to examine the trend in the use of antipsychotics after PD diagnosis in early PD patients who had psychosis and to determine whether the FDA safety warnings regarding the association of antipsychotics with a higher mortality rate decreased antipsychotic use in these patients. The secondary objective was to assess to what extent rivastigmine and other anti-dementia drugs were used to treat psychosis in PD patients.

Patients and Methods
Design and Data Source
This is a repeated cross-sectional study that examined the trend of antipsychotic use during the first year after PD diagnosis between 2000 and 2016. The Secure Anonymized Information Linkage (SAIL) databank was used to obtain the data for this study. SAIL contains the Welsh Longitudinal General Practice (WLGP) primary care dataset for at least 80% of the patients in Wales. SAIL is a central repository that was created by the Health Information Research Unit (HIRU) in the College of Medicine at Swansea University.\textsuperscript{23} The individual-level data contained in SAIL include but are not limited to GPs, hospital episodes, outpatient visits, demographics, and social deprivation data.\textsuperscript{23} This study was approved by SAIL’s independent Information Governance Review Panel (ref 0507). The data accessed in the SAIL databank
Identifying Early (Newly Diagnosed) PD Patients

Eligibility criteria included individuals with a first Read Code of a definitive PD diagnosis in GP data (Appendix 1). The Read Code system is a hierarchical system that covers most clinical terms, including diagnoses and drugs.24 The Read Codes indicating a definitive PD diagnosis were used and validated in a previous study.25 Patients were excluded for the following reasons: less than 40 years of age, taking any antiparkinsonian drug before PD diagnosis, no prescription of any antiparkinsonian drug within one year after PD diagnosis, a first code of PD within 6 months from SAIL registration date, death within a year of PD diagnosis, and previous antipsychotic use within the year prior to PD diagnosis, as antipsychotics can cause Parkinson-like symptoms which could be mistakenly diagnosed as PD.26

Identifying Psychosis Diagnosis and Antipsychotics

The purpose of this study was to identify the incidence of new cases of antipsychotic use to treat psychosis in the first year after PD diagnosis. To be considered an antipsychotic user, first, a patient was required to have a new Read Code of psychosis diagnosis (Appendix 1) that appeared in the data after the diagnosis of PD diagnosis. Additionally, the Read Code for antipsychotics must have appeared after the diagnosis of psychosis and within the first year after PD diagnosis. Appendix 2 lists drugs considered antipsychotics and their drug classes (FGA or SGA).

Identifying Anti-Dementia Drugs Used to Treat Psychosis in Patients without Dementia

Dementia is highly associated with both psychotic episodes and PD, especially in its later stages.27 To examine the use of anti-dementia drugs to treat psychosis in PD patients only, this analysis was restricted to PD patients without a previous history of dementia diagnosis and/or antipsychotic drug and/or anti-dementia diagnosis within 1 year after PD diagnosis (Appendix 1). The first prescription of anti-dementia drug within 1 year after PD diagnosis was reported. Anti-dementia drugs examined were donepezil, galantamine, memantine, and rivastigmine.

Factors Affecting Use of Antipsychotics in PD Patients

Patient demographics and social variables were obtained from SAIL. These included sex, age at PD diagnosis, year of PD diagnosis, status of social deprivation, and health board geographical location. Other variables included history of comorbidities (before PD diagnosis), the first PD medications prescribed, and previous use of drugs that induce psychosis. Age was categorized into three stages: 40–60 (early onset of PD), 61–80, and 81 years or older. Year of PD diagnosis was grouped to pre-2008 (ie, 2000–2008), and post-2008 (ie, 2009–2016). The year of 2008 was chosen to examine the effect of the FDA warning issued in that year.13 Year of PD diagnosis was treated as a continuous variable when examining use of anti-dementia drugs to treat psychosis. The status of social deprivation was classified into “most deprived” (quintiles 1 and 2), “middle deprived” (quintile 3), and “least deprived” (quintiles 4 and 5), according to the scale of the Welsh Index of Multiple Deprivation (WIMD). The seven health boards in Wales were listed in Appendix 3. Comorbidities were extracted from hospital data using the Charlson comorbidity index.28 The comorbidities consisted of acute myocardial infarction, cancer, cerebral vascular accident, congestive heart failure, connective tissue disorder, dementia, diabetes, diabetes complications, liver disease, metastatic cancer, paraplegia, peptic ulcer, peripheral vascular disease, pulmonary disease, renal disease, and severe liver disease. Each comorbidity was treated as a dichotomous variable. The first PD medication prescribed was defined as the first one prescribed to a patient after the PD diagnosis. For antipsychotic users, the first PD medication was required to be initiated before the first antipsychotic prescription. Drugs that induce psychosis included muscle relaxants, antihistamines, antidepressants, cardiovascular medications, antihypertensive medications, analgesics, anticonvulsants, and corticosteroids.29

Statistical Methods

The data were exported and analyzed using the Statistical Package for Social Sciences (SPSS) version 25 for Windows. Both descriptive and inferential statistics were used. Multivariate logistic regression was used to test the association between the independent predictors (eg, sex,
age, year of PD diagnosis and others) and the outcome
(Use of antipsychotic or anti-dementia drug to treat psy-
chosis). The Wald test was used to assess the significant
overall effect of each independent predictor in the model.
A particular predictor was included in the multivariate
analysis if the association had a p-value of <0.20. Odds
ratios and 95% confidence intervals were obtained from
the regression results. Statistical significance was set at p <
0.05. Abiding by the rules of SAIL to preserve patient
anonymity, aggregate numbers less than five were
excluded from the analysis.

Results
A total of 9142 PD patients were identified after applying
inclusion and exclusion criteria, of whom 58.6% were
male. Table 1 depicts the demographic profiles of the
newly diagnosed PD patients investigated in the study.
The majority (62%) were 61–80 years, with a mean age
of 73.7 and a standard deviation of 9.8. The years of PD
diagnoses were similar pre- and post-2008 (49.8% and
50.2%, respectively). Most PD patients (82%) used
Levodopa as an initial therapy. During the first year of
PD diagnosis, 21.8% of the patients developed psychosis,
and 55% of them were subsequently prescribed antipsy-
chotics. In general, SGAs were more commonly pre-
scribed (9.3%) to treat psychosis compared to FGAs
(2.7%), although the use of both notably decreased across
years (Figure 1). Among antipsychotic users (n = 1095),
quetiapine was the most commonly prescribed (49%),
followed by risperidone (10.7%), olanzapine (9.3%), and
haloperidol (7.1%) (Figure 2). After excluding PD patients
with a previous or current diagnosis of dementia (n = 255),
2.8% of PD patients with a diagnosis of psychosis were
prescribed anti-dementia drugs to treat psychosis
(Table 1). The prescribed anti-dementia drugs were done-
pezil, galantamine, memantine, and rivastigmine; rivastig-
mine was most commonly prescribed (76%).

The first logistic regression model (Table 2) showed
that the use of antipsychotics in newly diagnosed PD
patients was significantly lower in the years of
2009–2016 compared to 2000–2008 (OR 0.37, 95% CI
0.32–0.43). Age had a significant impact on antipsychotic
prescribing; patients aged 61–80 were highly likely to use
antipsychotics compared to those aged 40–60 (OR 1.44,
95% CI 1.11–1.86). Patients with a history of dementia
were 3.16 times more likely to use antipsychotics after PD
diagnosis (95% CI 2.35–4.24). Comorbidities such as

| Table 1 Patient Demographics |
|-------------------------------|
| Demographics                  | Mean (SD) or n (%) |
| N=9142                        |                  |
| Age, years (at PD incidence)  | 73.7 (9.8)       |
| Age group                     |                  |
| 40–60 years                   | 845 (9.2)        |
| 61–80 years                   | 5670 (62)        |
| >80 years                     | 2627 (28.8)      |
| Sex, male                     | 5358 (58.6)      |
| Health board (geographical health board location) |         |
| Abertawe Bro Morgannwg        | 2128 (23.28)     |
| Aneurin Bevan                 | 1408 (15.4)      |
| Betsi Cadwaladr               | 2005 (21.93)     |
| Cardiff and Vale              | 1275 (13.95)     |
| Cwm Taf                       | 845 (9.24)       |
| Hywel Dda                     | 1158 (12.67)     |
| Powys                         | 323 (3.53)       |
| Year of PD diagnosis          |                  |
| 2000–2008                     | 4551 (49.8)      |
| 2009–2016                     | 4591 (50.2)      |
| PD medications*              |                  |
| Amantadine                    | 22 (0.2)         |
| Anticholinergics              | 325 (3.6)        |
| COMT inhibitors               | 58 (0.6)         |
| Dopamine agonists             | 1291 (14.1)      |
| Levodopa                      | 7366 (80.6)      |
| MAO-B inhibitors              | 719 (7.9)        |
| PD patients with a diagnosis of psychosis without antipsychotics and/or anti dementia drugs (within 1 year after PD diagnosis) | 1994 (21.8) |
| PD patients with a diagnosis of psychosis and received antipsychotics (within 1 year after PD diagnosis) | 1095 (12) |
| Antipsychotics medications    |                  |
| First generation antipsychotics | 245 (2.7)      |
| Second generation antipsychotics | 850 (9.3)       |
| Anti-dementia drugs likely used to treat psychosis (PD patients with a dementia diagnosis were excluded, n = 255, so the total PD patients, n = 8887) |       |
| Cholinesterase inhibitors anti dementia (Donepezil, Galantamine, Rivastigmine) | 247 (2.7) |
| NMDA antagonist anti dementia (Memantine) | 12 (0.1) |

Note: *It is possible that a patient can take more than one of the following medications.

Abbreviations: COMT inhibitors, catechol-O-methyltransferase inhibitors; MAO-B inhibitors, monoamine oxidase B inhibitors; NMDA antagonist, N-methyl-D-aspartate receptor antagonist; PD, Parkinson’s disease.
myocardial infarction and peptic ulcer increased the odds of antipsychotic use (OR 1.59, 95% CI 1.15–2.2 and OR 2.27, 95% CI 1.06–4.83, respectively). There was no significant effect of social deprivation status (WIMD scale) on antipsychotic use (see Table 2 for associated ORs and 95% CIs).

Regarding the use of anti-dementia drugs to treat psychosis in PD patients within the first year of their PD diagnosis, the second logistic regression model (Table 3) showed different factors compared to the previous model (Table 2). For example, although Figure 3 showed a decrease in anti-dementia drug use over years, this decline was not significant in the adjusted regression model (OR 0.78, 95% CI 0.52–1.57). Patients aged >80 years were more likely to use anti-dementia drugs (OR 0.53, 95% CI 0.28–0.98). Further, patients who resided in the least
Table 2 Multiple Logistic Regression Examining Potential Factors Affecting the Trend Antipsychotics Use in PD Patients Within the First Year of Their PD Diagnosis

| Variables                              | OR (95% CI) | P-value |
|----------------------------------------|-------------|---------|
| Year of PD diagnosis (ref = 2000–2008) | 0.37 (0.32–0.43) | 0.000   |
| Age groups (years)                     |             |         |
| 40–60                                  | Reference   |         |
| 61–80                                  | 1.44 (1.11–1.86) | 0.005   |
| >80                                    | 1.21 (0.91–1.6)  | 0.186   |
| Sex, (ref = male)                      | 0.99 (0.87–1.13) | 0.894   |
| Comorbidities                          |             |         |
| Myocardial infarction                  | 1.59 (1.15–2.2)  | 0.005   |
| Dementia                               | 3.16 (2.35–4.24) | 0.000   |
| Peptic ulcer                           | 2.27 (1.06–4.83) | 0.034   |
| WIMD (Welsh index of multiple deprivation) |         |         |
| Most deprived (WIMD 1 and 2)          | Reference   |         |
| Middle deprived (WIMD 3)               | 0.87 (0.73–1.04) | 0.130   |
| Least deprived (WIMD 4 and 5)          | 0.93 (0.81–1.08) | 0.366   |
| PD medications                         |             |         |
| Levodopa                               | 0.93 (0.77–1.13) | 0.484   |
| MAO-B inhibitors                       | 0.56 (0.4–0.79)  | 0.001   |
| Anticholinergics                       | 3.19 (2.43–4.2)  | 0.000   |
| Previous use of psychosis-induced medications |         |         |
| Beta blockers                          | 1.19 (1.02–1.39) | 0.031   |
| Calcium channel blockers               | 0.88 (0.74–1.06) | 0.176   |
| ACE inhibitors                         | 0.9 (0.76–1.06)  | 0.205   |
| ARB blockers                           | 0.63 (0.46–0.87) | 0.005   |
| TCA                                    | 1.43 (1.17–1.75) | 0.001   |
| SSRI antidepressant                    | 1.64 (1.39–1.92) | 0.000   |
| Other antidepressants                  | 2.17 (1.69–2.77) | 0.000   |

Note: Bold numbers indicate statistically significant differences.

Abbreviations: MAO-B inhibitors, monoamine oxidase B inhibitors; PD, Parkinson’s disease; ACE, angiotensin-converting enzyme; ARB, angiotensin receptors blockers; TCA, tricyclic antidepressants; SSRI, selective serotonin reuptake inhibitors.

deprieved areas (WIMD 4 and 5) were more likely to be prescribed anti-dementia drugs (OR 1.62, 95% CI 1.17–2.24) (see Table 3 for associated ORs and 95% CIs).

Discussion
This study examined antipsychotic use in a large, national sample of patients in Wales who received a diagnosis of PD and who had been followed up for one year after PD diagnosis between 2000 and 2016. A substantial decrease in antipsychotic prescribing patterns was found after the FDA issued its warnings against the use of antipsychotics in dementia patients. Although the FDA warning was directed to dementia patients, the prevalence of dementia in PD patients can reach 80%, especially in the later stages of PD. Additionally, dementia is strongly associated with psychosis in PD. Therefore, it was expected that the FDA warning would decrease the use of antipsychotics in PD patients. The current study provides data to show this decrease, demonstrating a reasonable adherence to national and international guidance.

Several previous studies estimated that the prevalence of antipsychotic use in PD patients ranged from 0.45% to 18.3%. The cumulative probability of using antipsychotics after PD diagnosis was found to be 35% after 7 years in a Canadian study, and 51% after 6 years in a Taiwanese study. However, these studies examined all stages of PD, and none examined antipsychotic use in newly diagnosed PD patients. Psychosis in the early stages of PD is not rare, and symptoms such as minor hallucination could affect...
Table 3  Multiple Logistic Regression Examining Potential Factors Affecting the Trend of Anti-Dementia Use to Treat Psychosis in PD Patients Within the First Year of Their PD Diagnosis

| Variables                                      | OR (95% CI) | P-value |
|------------------------------------------------|-------------|---------|
| Year of PD diagnosis (continuous)              | 0.78 (0.52–1.07) | 0.422 |
| Age groups (years)                             |             |         |
| 40–60                                          | Reference   |         |
| 61–80                                          | 1.12 (0.67–1.88) | 0.657 |
| >80                                            | 0.53 (0.28–0.98) | 0.043 |
| Sex, (ref = male)                              | 0.74 (0.56–1) | 0.047 |
| Comorbidities                                  |             |         |
| Cerebrovascular accidents                      | 0.71 (0.21–2.47) | 0.595 |
| Congestive heart failure                       | 0.47 (0.11–2.1) | 0.323 |
| Diabetic type 2                                | 0.53 (0.26–1.06) | 0.072 |
| Pulmonary diseases                             | 0.8 (0.38–1.69) | 0.558 |
| WIMD (Welsh index of multiple deprivation)     |             |         |
| Most deprived (WIMD 1 and 2)                   | Reference   |         |
| Middle deprived (WIMD 3)                       | 1.16 (0.78–1.72) | 0.477 |
| Least deprived (WIMD 4 and 5)                  | 1.62 (1.17–2.24) | 0.004 |
| PD medications                                 |             |         |
| Levodopa                                       | 1.15 (0.76–1.74) | 0.505 |
| Dopamine agonists                              | 1.67 (1.05–2.64) | 0.029 |
| Current use of antipsychotics                  | 104.66 (68.07–160.93) | 0.000 |
| Previous use of psychosis induced medications  |             |         |
| Beta blockers                                   | 1.22 (0.89–1.67) | 0.219 |
| NSAIDs/COX2 inhibitors                         | 1.7 (1.22–2.35) | 0.002 |
| SSRIs antidepressant                           | 0.84 (0.59–1.2) | 0.330 |

Note: Bold numbers indicate statistically significant differences.
Abbreviations: PD, Parkinson’s disease; NSAIDs, non-steroidal anti-inflammatory; COX2, cyclooxygenase-2; SSRIs, selective serotonin reuptake inhibitors.

42% of newly diagnosed PD patients. Therefore, a novel finding from the current study suggests that 12% of newly diagnosed PD patients were prescribed antipsychotics after being diagnosed with psychosis.

The current study found that more than half of PD patients with psychosis (55%) were prescribed antipsychotics, leaving about 45% of the patients without treatment. This is understandable since the first recommended approach to treating psychosis in PD patients is to reduce the doses of PD medications and then to consider antipsychotic medications if needed, however, the data on prescribed doses are not available in the SAIL databank to confirm this explanation. Similar to previous international studies, the current study found that the majority of antipsychotics prescribed to PD patients were SGAs. Quetiapine in particular constituted almost half of the prescribed antipsychotics, the same pattern found in two previous US studies. The tendency to prescribe quetiapine and to avoid FGAs in the current study is in line with medical literature that linked FGAs to worsening Parkinson-like motor symptoms, which it is not desirable in PD patients. Additionally, among SGAs, quetiapine is more tolerated in PD patients compared to other SGAs. Although no clinical trial has confirmed its efficacy and superiority over other SGAs, clinical experience suggests that quetiapine is possibly useful in treating psychosis in PD patients without noticeable side effects that need continuous monitoring. It is of note that clozapine—although it is recommended for use in PD psychosis was not prescribed to PD patients in the study sample. Two possible reasons for this are the risk of agranulocytosis that accompanies clozapine use, and its requirement for continuous blood monitoring. Another reason might be that the UK guidance (NICE guidance) considers clozapine to be the second-line treatment after quetiapine. Furthermore, the current study examined only the first use...
of antipsychotics during the first year after PD diagnosis, which increased the likelihood that the therapy would start with a treatment that requires less monitoring, ie, quetiapine. It is possible that clozapine was initiated in some PD patients in the later stages of PD but due to the lack of data later than 1 year after the initial PD diagnosis, this cannot be confirmed. Among antipsychotics, the current study did not include newer SGAs such as pimavanserin. Pimavanserin was approved by the USFDA in 2016 for treating psychosis in PD patients and it shows good efficacy to treat psychosis without worsening Parkinsonian symptoms in PD patients; however, it is not yet approved in the UK.

Regarding the comorbidities associated with antipsychotic use, the current study found that myocardial infarction and peptic ulcer increased the odds of antipsychotic use in PD patients. Although it has been reported that antipsychotic use could cause myocardial infarction and peptic ulcer, no previous study has suggested that the two comorbidities increased the likelihood of antipsychotic use. Whether this is due to a bidirectional relationship between antipsychotics and the two comorbidities, or due to a previous use of antipsychotics that caused comorbidities before PD diagnosis, is not clear. However, the current study excluded from the analysis all newly diagnosed PD patients who used antipsychotics within 1 year prior to PD diagnosis. Nevertheless, more research is needed to determine the nature of such an association.

In contrast to previous literature that linked residing in more deprived areas to an increase in antipsychotic use in the UK general population, the current study found no difference between areas ranked as the most deprived relative to areas ranked as deprived in terms of antipsychotic use in PD patients. A possible explanation may be that patients in deprived areas are more likely to present the first symptoms of PD to a primary care provider in later stages of the disease; therefore, the provider may avoid prescribing antipsychotics since they worsen PD motor symptoms.

No previous study has examined the prescribing pattern of anti-dementia drugs to treat psychosis in PD patients. The current study found that the use of anti-dementia drugs to treat psychosis in newly diagnosed PD patients was very low (2.8%), and this trend declined across the years of the study. Although there is some evidence that rivastigmine, for example, can reduce hallucination and improve behavioral symptoms in PD patients, recent NICE guidelines do not recommend rivastigmine as part of therapy. The low prescribing rate of anti-dementia drugs in newly diagnosed patients shown in this study cannot be extrapolated to the later stages of PD due to the high dementia rate in advanced PD patients, which may lead to increased prescribing of rivastigmine to manage both dementia and psychosis.

In contrast to antipsychotic use, which showed no difference among areas of different deprivation status, anti-dementia drugs were more likely to be used by PD patients who resided in the least deprived areas. This social inequality in access to anti-dementia drugs was seen previously in dementia patients in the UK. The
current study extended this inequality to PD patients who use these drugs to treat psychosis. Anti-dementia drugs may be less accessible to patients in more deprived areas because these patients have a higher profile of morbidity and mortality, may attend primary care clinics that lack resources, and may not attend their appointments. Efforts must be made to address this inequality.

Strengths of the study include the large sample size, robust inclusion and exclusion criteria, and the use of multiple covariates linked to primary care data from different sources in Wales. Limitations include the inability to verify the PD diagnosis, such as confirming the diagnosis by a movement disorders neurologist; however, the diagnosis of other disease was validated in the SAIL data. Another limitation was that the SAIL data did not indicate the severity of PD.

Conclusion
This study found a substantial decrease in antipsychotic prescribing patterns after the FDA issued warnings in 2005 and 2008 against the use of antipsychotics, showing a reasonable adherence to national and international prescribing guidance. Continuous and closer monitoring of antipsychotic use is warranted, which must be balanced with the effective and safe use of antipsychotics in PD patients.

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