Evidence shows that metabolic syndrome (MS) is associated with a greater risk of developing Parkinson’s disease (PD) because of the increase in oxidative stress levels along with other factors such as neuroinflammation and mitochondrial dysfunction. However, because some studies have reported that MS is associated with a lower risk of PD, the relationship between MS and PD should be investigated. This study aimed to investigate the effect of MS on PD. Two authors searched five electronic databases, namely, MEDLINE, PubMed, Scopus, PsycINFO, Web of Science, and Science Direct, for relevant articles between September and October 2020. After screening the title and abstract of all articles, 34 articles were selected for full-text review. Finally, 11 articles meeting the eligibility criteria were included in the study. The quality of articles was critically evaluated using the Joanna Briggs Institute. Overall, we evaluated data from 23,586,349 individuals (including healthy individuals, with MS and PD) aged 30 years or more. In cohort studies, the follow-up period varied between 2 and 30 years. MS contributed considerably to the increase in the incidence of PD. In addition, obesity, a component of MS, alone can increase the probability of developing neurodegenerative diseases. However, despite few studies on MS and PD, changes in cognitive function and more rapid progression of PD disease has been documented in patients with MS using methods commonly used in research.

KEYWORDS: Metabolic Syndrome; Parkinson Disease; Vascular Risk Factors.
this relationship and provide information for the development of preventive strategies and treatments for MS and PD.

**METHODS**

This systematic review was conducted according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines. The study protocol is registered with the International Prospective Register of Systematic Reviews (PROSPERO) (https://www.crd.york.ac.uk/prospero/; registration number: CRD42020210589).

**Eligibility criteria**

**Study design and participants.** Eligibility criteria were defined based on the Population, Intervention, Comparison, Outcome and Study strategy: population, patients with PD and those with MS; comparison, between patients with and without MS; outcome, effects of MS. We included all longitudinal and cross-sectional studies on MS and PD, studies investigating the consequences of MS in patients with PD, and incidence studies investigating the association between MS and PD. Animal studies, studies on other pathologies, studies on some interventions, and revision research were excluded.

**Databases and research strategy**

Two authors (APSS and JMLS) independently searched five electronic databases (Medline/PubMed, Scopus, PsycINFO, Web of Science, and Science Direct) between September and October 2020. Regarding the extraction, selection, and data collection, the two authors initially selected titles and abstracts, according to the eligibility criteria, which were later discussed to resolve any doubts or disagreements. For this stage, the descriptors MeSH “Metabolic syndrome” and “Parkinson disease” were used in all databases.

The full text of selected articles were reviewed to confirm their eligibility. Any disagreements were resolved by discussion with the third author (WMAB). The following data were collected from all articles: author names, year of publication, country, study design, study population (number, sex, and age or average age), goal, analysis methods, data collection instruments, criteria for the diagnosis of MS; biochemical parameters assessed, anthropometric parameters assessed, and general results.

The quality of all articles was assessed using the Joanna Briggs Institute (Institute JB approach available at https://joannabriggs.org/critical-appraisal-tools), an instrument to verify the reliability and relevance of selected articles. Evaluation was performed according to the study design. This step was conducted by two authors (APSS and JMLS) performed the evaluation according to the study design and any disagreements were resolved by consensus.

**RESULTS**

**Selection of studies**

The electronic search of databases yielded 1,053 articles (Medline/PubMed: 278; Scopus: 425; PsycINFO: 81; Web of Science: 214; and Science Direct: 55). After excluding 197 duplicates, the title and abstract of the remaining articles were screened based on inclusion criteria. A total of 34 articles were selected for full text review. Finally, 11 articles meeting the established eligibility criteria were included in the study (Figure 1).

**Study characteristics**

This study included six cohort studies (6,7,9,15–17), four cross-sectional studies (18–21), and one case-control study (22). The locations of the included studies (N=11) were as follows: three studies, South Korea (6,7,15); two studies, the United States of America (19,20); one study, Israel (18); one study, South Africa (22); one study, China (16); one study, Canada (17); one study, Italy (21); and one study, Finland (9) (Table 1).

Overall, were evaluated data from 23,586,349 individuals (including healthy individuals with MS and PD) aged 30 years or more. In cohort studies, the follow-up period varied between 2 and 30 years. Individuals of both sexes were evaluated in all studies, except one study (22) in which only women were evaluated for methodological reasons (longer hair length in women). Three used secondary data from a national database, the National Health Insurance Service (NHIS) of South Korea (6,7,15).

The included studies used different criteria and guidelines for the diagnosis of MS, which were as follows: four studies (6,16,20,21), the National Cholesterol Education Program Adult Treatment Panel III; two studies (7,22), the Joint Interim Statement (JIS); one study, the harmonized definition of metabolic syndrome (9); and in 4 studies (15,17–19), the guideline used for the diagnosis of MS was not mentioned, but only the criteria used.

Six studies (16–18,20–22) used the same scale (the Unified Parkinson’s Disease Rating Scale [UPDRS]) to assess the disease progression of PD in participants. A study (20) evaluating the progression of PD reported high UPDRS scores in individuals with MS, i.e., faster progression of PD. However, a study evaluating the association of MS with the occurrence of falls and PD progression (21) revealed that MS is associated with a reduction in the number of falls.

In the present review, the effects of MS on incidence of PD were investigated. Two studies evaluated the cognitive capacity of individuals (16,17). However, other studies evaluated this variable using common scales such as the Montreal Cognitive Assessment (17,18), the Symbol Digit Modalities Test (SDMT) (17,20), and the Mini-Mental State Examination (16,19,21).

There is little evidence in the literature regarding the effect of MS on the incidence of PD. Four studies included in this review investigated the effect of MS on the incidence of PD (6,7,9,15). Only one study (9) reported that the presence of MS indicated a 50% lower risk of developing PD. In two studies (6,7), this trend increased with the number of MS components. A study evaluating prodromal characteristics (18) did not indicate the increased risk of developing PD in individuals with MS.

**Quality evaluation of articles**

No study was excluded according to the quality assessment carried out for each study design (Tables 2, 3, and 4).

**DISCUSSION**

In this systematic review, we investigated the effect of MS on PD. The main finding of this study is that the incidence of PD is high among individuals with MS, confirming the research hypothesis that MS has promotive effects on PD.
In addition, abdominal obesity, overweight, and obesity increase the risk of developing PD among individuals with MS (6,9,15).

Some studies, including a cohort study, investigated the incidence of PD using secondary data of 314,737 individuals aged >40 years (7). In an average 7.3-year follow-up period, 48% of the volunteers developed PD. The incidence of PD was higher in the group with MS. Similar trend was observed among patients with hypertension of both sexes. Another cohort study evaluated 6,641 individuals aged 30–79 years for a period of 30 years (9). A total of 89 individuals developed PD and overweight patients had a suggestively greater risk of PD pathology; however, presence of MS indicated a 50% lower risk of developing a neurodegenerative disease. However, although the follow-up period was long, the sample size of this study was relatively small and may have interfered with the results in terms of incidence.

In this review, a study evaluating secondary data of 17,163,560 individuals aged >40 years found that abdominal obesity increased the risk of PD (6). Moreover, with an average follow-up of 5.3 years, the study found that patients with MS are more likely to develop PD than those without MS. In addition, the study reported a positive association between PD and the number of components of MS in question.

The components of MS include increased waist circumference, a characteristic of abdominal obesity, and waist circumference can be related to dietary patterns. In this sense, a cohort study highlighted the importance of a healthy diet in association with a lower risk of PD development (23). The study evaluated 3,653 individuals of both the sexes (average age: 81.5 years) who were followed up for an average period of 6.94 years, verified the quality of the diet of the participants using a dietary screening tool, and reported 47 incident cases of PD. However, although the follow-up period was short, the number PD of cases was high. This may be because of the high average age of the participants; hence, studies investigating participants with a lower average age for a longer period if time should be conducted. In addition, the authors indicated some limitations such as the small diversity of races in the study population, lack of evaluation of energy consumption, and few food options in the questionnaire used. However, the same study meta-analyzed four more studies on the topic, including 140,617 individuals, and

Figure 1 - Flowchart of study selection.
Table 1 - Main characteristics of the included studies.

| Study, year, Country, Study design | Sample (number, sex, age or mean age) | Objective | Methods | Data collection instruments |
|-----------------------------------|---------------------------------------|-----------|---------|---------------------------|
| **Roh et al. (7)** | N= 314,737 | To investigate the effect of MS on the incidence of PD | Analysis of data from the National health service database. (NHIS-HealS) (2002-2015) | Secondary data |
| South Korea, Cohort study | 56.1% M | | | |
| | 85,530 individuals with MS and 85,530 individuals without MS | | | |
| | 40 years old | | | |
| | N=562 | To evaluate the effect of MS on the phenotype of LRRK2 and PD GBA and on the prevalence of prodromal characteristics among individuals at risk for PD | Genetic tests for the G2019S mutation in the LRRK2 gene and for the AJ GBA mutations; laboratory tests; Information; cognitive, motor, olfactory and affective functions. | MDS-UPDRS; MoCA, BDI; NMSQ; SCOPA-AUT; RBDO; UPSIT. Demographic data; PAN; Blood analysis of HbA1c, TG and HDL levels. |
| **Thaler et al. (18)** | N=562 | To evaluate the effect of MS on the phenotype of LRRK2 and PD GBA and on the prevalence of prodromal characteristics among individuals at risk for PD | Genetic tests for the G2019S mutation in the LRRK2 gene and for the AJ GBA mutations; laboratory tests; Information; cognitive, motor, olfactory and affective functions. | MDS-UPDRS; MoCA, BDI; NMSQ; SCOPA-AUT; RBDO; UPSIT. Demographic data; PAN; Blood analysis of HbA1c, TG and HDL levels. |
| South Africa, Cross-sectional study | M and F | | | |
| | 104 individuals with idiopathic PD, 40 individuals with LRRK2-PD, 70 individuals with GBA-PD, 196 healthy carriers, 55 individuals with LRRK2-PNM and 97 individuals with GBA-PNM | | | |
| **van den Heuvel et al. (22)** | 56 F (25 pcts F with PD and 31 controls) aged 45 to 78 years | To examine the association of PD diagnosis, clinical characteristics, and risk of PD-CVD (as defined by the MS) with the levels of GC (cortisol and cortisone) in hair. | Analysis of hair samples, representing a 3-month retrospective window of GC levels using tandem mass chromatography liquid chromatography. | Glycemia, TG, HDL, HbA1c; LDL, total blood cholesterol. Sociodemographic data, MINI, UPDRS, H&I, NMSS; PSS-10; WHO STEPS; GPAQ. |
| South Africa, Case-control study | F and M | | | |
| | > 40 years old | | | |
| | N=6,098,405 | To assess the association between the serum GGT level and PD risk. Analyze the possible interaction between the GGT level and obesity or MS. | National Health Insurance Service (NHIS) database | Secondary data |
| **Nam et al. (6)** | N=17,163,560 | To investigate the association of MS and its components with the development of PD in the South Korean population using large cohort data. | Analysis of NHIS data from January 1, 2009, to December 31, 2012 and follow-up of pcts up to 31 December 2015. | Secondary data |
| South Korea, Cohort study | age > 40 years | | | |
| | Individuals with MS: 58.1 years Individuals without MS: 51.9 years | | | |
| **Peng et al. (16)** | N=787 | To investigate the effect of MS on cognitive impairment in PD. | Physical and neurological examinations, blood analysis and neuroimaging. MS incidence assessed based on medical records or doctors’ assessment. Annual monitoring for 5 years. | UPDRS, MMSE, RAVLT, Doppler ultrasonography, computed tomography angiography, magnetic resonance imaging |
| China, Cohort study | 60 years | | | |
| | Individuals with PD divided into three groups: PD-CN, PD-CCL and PDD. 249 individuals with PD and MS at the beginning of the study divided into 3 subgroups according to the treatment of each component of MS: untreated pcts, pcts who received treatment for some components, and pcts who received treatments for all components. | | | |
| **Oxenkrug et al. (19)** | 7 F pcts with PD and 11 M pcts with PD (age range: 50 to 74 years). 12 F pcts with probable AD and 8 M pcts with probable AD (age range: 60 to 75 years). 24 healthy individuals matched for age and gender (12 F and 12 M). | To compare peripheral quinueneines in AD and PD with an emphasis on quinueneines associated with MS (i.e., KYNA, ANA, 3-HK, and XA) | Analysis of blood samples collected after overnight fasting. | Analysis of Trp, Kyn, ANA, KYNA, and 3-HK using high performance liquid chromatography coupled with mass spectrometry. MMSE |
| United States of America (USA), Cross-sectional study | | | | |
| Study, year, Country, Study design | Sample (number, sex, age or mean age) | Objective | Methods | Data collection instruments |
|-----------------------------------|--------------------------------------|-----------|---------|----------------------------|
| Leehey et al. (20) USA Cross-sectional study | N=1022 M and F 2 groups: Individual with MS (N=396, mean age 63.9 years) over the 3 years of the study and individuals without evidence of MS (N=626, mean age 59.9 years) over the 3 years. | To compare the progression of PD between individuals with MS during the first 3 years of the trial and individuals with no evidence of MS. | Secondary analysis of patient data from the NET-DP LS study 1. The changes in the UPDRS and SDMT scores from randomization to 3 years were compared. | UPDRS, SDMT |
| Doiron et al. (17) Canada Cohort study | 367 pts (&gt; 50 years old) with PD at baseline and 310 pts at 24-month follow-up. | To investigate hypertension, dyslipidemia, DM, and BMI as possible risk factors for cognitive impairment in a large cohort of untreated patients with PD and without dementia. | Clinical and neurological examinations, biological sampling, neuropsychological assessments, and neuroimaging. Participants reassessed approximately after 24 months. | UPDRS, GDS, GDS, LNS, SDMT, HVLT-R, BJLO, total blood cholesterol, LDL, HDL, and TG. |
| Lauidisio et al. (21) Italy Cross-sectional study | N=194 Elderly individuals with PD (IM 73 years old) treated at a geriatric hospital at the Catholic University of Rome. | To evaluate the association between MS and occurrence of falls in patients with PD. | Recorded the history and number of falls in the last year. | ADL, IADLs, GDS, MMSE, UPDRS, MNA, dual energy X-rays Absorptiometry, Portable dynamometer, Tinetti, 24-h BP recording. |
| Sääksjärvi et al. (9) Finland Cohort study | 6641 aged 30-79 years and without PD at baseline (1978-1980) | To investigate whether MS or its components, or serum total cholesterol, can predict the incidence of PD in a prospective cohort. | Based on the Mini-Finland Health Survey conducted from 1978 to 1980. Anthropometric, biochemical blood analyses were performed. | HDL, TG, Glycemia, and total cholesterol in the blood. 25-hydroxy vit D serum by radioimmunoassay |
Table 1 - Columns Continued.

| Study, year, Country, Study design | Diagnosis of MS and related component | Results: biochemical parameters | Results: anthropometric variables | Main Results |
|-----------------------------------|--------------------------------------|-------------------------------|---------------------------------|--------------|
| **Roh et al. (7)** South Korea Cohort study | Joint Interim Statement (JIS) | HDL↑ | WC was not associated with the incidence of PD in individuals in general. | Average follow-up period: 7.23 years. 819 (0.48%) individuals developed PD. Incidence among individuals with MS Hypertensive 1.34 incidence of PD in both sexes Incidence of PD increased with the number of components of MS Groups with PD and PNM did not differ in the number of metabolic components. 70 PD patients (32.7%) with MS. This group was older and had a longer diagnosis time. 94 (27.2%) PNM with MS. Hypertension was associated with age and sex M was responsible for 35.7% of the variance. Individuals with MS did not have a high risk of developing PD. |
| **Thaler et al. (18)** Israel Cross-sectional study | Change in 3 of the 5 components: fasting glucose levels, BP, BMI, HDL, and TG. | LRRK2 -PD↑ TG levels and pre-diabetes rates. LRRK2 -PNM TG levels PNM with probability rates for prodromal PD. ↑ 50% had frequencies of hypertriglyceridemia and pre-diabetes. Pre-diabetes associated with age and UPDRS-III scores accounting for 38.8% of the variance. The probability ratio for conversion to PD in 10 (3%) individuals was >80% (non-carriers: 4 individuals, GBA-PNM: 3 individuals, and LRRK2-NMC: 3 individuals). | Pre-diabetes associated with age and sex M was responsible for 35.7% of the variance. Individuals with MS did not have a high risk of developing PD. |
| **van den Heuvel et al. (22)** South Africa Case–control study | Joint Interim Statement (JIS) | Capillary cortisone in PD patients than in controls. Levels of cortisone in hair in unadjusted analysis were associated with the number of criteria for MS, use of antihypertensive medication, WHR, known CVD and use of aspirin, and a tendency to demonstrate significance for HDL-C. | Prevalence of MS in Pcts with PD was 56.0% and that in control individuals was 25.8%. Non-motor symptoms of PD (mood anhedonia and anxiety) were associated with capillary cortisone levels. No significant interactions were found between PD and MS based on the GC levels in hair. Average follow-up period was 6.4 years. DP developed in 20,895 (0.34%) individuals (M: 9,512 and F: 11,383). PD incidence was low in M, whereas ↑ high in F according to GGT activity Obesity and MS are the risk factors for PD in both sexes. A sub additive interaction was found between the serum GGT level and obesity in women. |
| **Yoo et al. (15)** South Korea Cohort study | Change in 3 of the 5 components or use of antihypertensive, hypoglycemic, or hypolipemic medication. | - | Factors associated with the serum GGT level were age, low income, BMI, current smoking, light to moderate alcohol consumption, exercise, DM, hypertension, dyslipidemia, chronic kidney disease, and MS |

(Continued)
Table 1 - Continued.

| Study, year, Country, Study design | Diagnosis of MS and related component | Results: biochemical parameters | Results: anthropometric variables | Main Results |
|-----------------------------------|--------------------------------------|--------------------------------|---------------------------------|--------------|
| Nam et al. (6) South Korea Cohort study | National Cholesterol Education Program Adult Treatment Panel III. | Pcts with 3 MS components at 31% PD risk, and those with all 5 components at 66% risk compared with those without any component | MS group had higher mean BMI, WC, BP, glucose level, serum total cholesterol level, TG level, and LDL-C level. Individuals with abdominal obesity or hypertriglyceridemia had approximately 13% PD risk. | Average follow-up period was 5.3 years. 44,205 incident cases of PD. Individuals with MS (n=5,848,508) had a higher risk of PD than individuals without MS (n=11,315,052). Each component of MS was positively associated with the risk of PD. Positive correlation was found between the incidence of PD and the number of components of MS. At 5 years, 255 (32.4%) individuals were diagnosed with PD-CCL and 105 (13.3%) individuals were diagnosed with PDD. MS was associated with PD-CCL. MS was associated with the main cognitive domains in Pcts with PDD. Incidence of PD-CCL and PDD was higher in individuals with MS than in those without MS. Pcts treated for some or all of the major components of MS had a lower risk of PDD than patients with untreated MS. |
| Peng et al. (16) China Cohort study | National Cholesterol Education Program’s Adult Treatment Panel III. | Individuals with PD-CCL and those with PDD had high levels of hypertension, glucose, and hypertriglyceridemia than individuals with PD-CN. Associations between BP, glucose, TG, and PDD. BP and glucose levels were associated with the main cognitive domains; TG level was associated with executive function, language, memory, and visuospatial function; HDL-C level was associated with memory and visuospatial functions. | Individuals with PD-CCL and those with PDD were significantly older and had higher BMI, UPDRS 3 scores, H&Y stage, systolic and diastolic BP, glucose level, total cholesterol level, and TG level as well as longer duration of PD. Still had lower education level (< 6 years), lacunar infarction, white matter lesions, and MS than those with PD-CN. |
| Oxenkrug et al. (19) United States of America (USA) Cross-sectional study | N/S differences in plasma concentrations of Trp, Kyn, and all Kyn metabolites were found in untreated patients and patients treated with L-DOPA. | | In Pcts with PD, Trp concentrations and the reason Kyn: Trp, Kyn, ANA and KYNA ↑ than in the control. Concentrations of 3-HK in Pcts with PD were below the sensitivity limit of the method. In Pcts with AD, the serum ANA concentrations were approximately ↑ 3 times higher and KYNA concentrations were approximately ↓ 40% more than those in control individuals. In Pcts with PD, Trp concentration is ↑ but Kyn concentration is ↓. In Pcts with AD, Trp concentration is ↑ but Kyn concentration is ↓. |
| Leehey et al. (20) USA Cross-sectional study | National Cholesterol Education Program Adult Treatment Panel III (modified). | - | - | Pcts with older MS: propensity to be of the sex M (75.3% versus 57.0%), and ↑ an average level of uric acid. |

(Continued)
| Study, year, Country, Study design | Diagnosis of MS and related component | Results: biochemical parameters | Results: anthropometric variables | Main Results |
|-----------------------------------|---------------------------------------|-------------------------------|-----------------------------------|--------------|
| Doiron et al. (17) Canada Cohort study | Hypertension, DM, Obesity | HDL was correlated with BJLO performance at baseline | BMI was associated only with the verbal fluency Z score at 24 months. | In Pcts with MS there was an additional 0.6- (0.2) units per year in the total UPDRS and 0.5- (0.2) units in the motor UPDRS scores compared with participants without MS. N/S changes in SDMT scores. Hypertension was the most prevalent comorbidity in 35.4% of the individuals at baseline and 41.1% of the individuals at follow-up. Time of hypertension and pulse pressure were associated with worse cognitive outcomes. Longer history of hypertension and higher pulse pressure were associated with impaired verbal episodic memory and semantic verbal fluency. Other vascular risk factors were not associated with cognitive outcomes. |
| Laudisio et al. (21) Italy Cross-sectional study | National Cholesterol Education Program Adult Treatment Panel III. | N/S associations were noted between the occurrence of falls and any of the components of MS. | - | 91 (47%) participants reported falls. MS was diagnosed in 44 (23%) participants. Severity of PD according to the UPDRS did not differ significantly according to the diagnosis of MS. MS was associated with less falls. |
| Sääksjärvi et al. (9) Finland Cohort study | Harmonized definition of the metabolic syndrome | ↑ Serum TG and fasting plasma glucose predicted ↑ risk of PD, even after excluding the first 10 years of follow-up. PA, HDL, or total cholesterol do not predict PD risk. | ↑ Risk suggestive of PD observed in overweight individuals | During 30-year follow-up (1978-2007), 89 incident cases of PD were recorded. MS predicted the risk of PD up to 50%. |

Abbreviations: 3-HK, 3-hydroxyquinurenine; AD, Alzheimer's disease; ADLs, Katz's activities of daily living; ANA, anthranilic acid; BDI, Beck Depression Inventory; BJLO, Benton Judgment of Line Orientation; BMI, body mass index; BP, blood pressure; CC, waist circumference; CVD, cardiovascular disease; F, female; GBA, glucocerebrosidase genes; GC, glucocorticoid; GDS, Geriatric Depression Scale; GGT, serum gamma-glutamyltransferase; GPAQ, global physical activity questionnaire; HbA1c, Glycated hemoglobin; HDL, high-density lipoprotein; H HVLT-R, Hopkins Verbal Learning Test Revised; IADL, Lawton and Brody scale for instrumental activities of daily living; KYNA, quinurenic acid; LDL, low-density lipoprotein; LNS, Letter–Number Sequencing; LRRK2, repeat of leucine-rich kinase 2; M, male; MINI, Mini International Neuropsychiatric Interview; MMSE, Mini-Mental State Examination; MNA, Mini Nutritional Assessment; MoCA, Montreal Cognitive Assessment; MS, Metabolic syndrome; NET-PD LS 1, National Institute of Neurological Diseases and Stroke Exploratory Trials in the Long Term Study 1 of PD; NHS-HEALS National Health Insurance Service-National Health Screening Cohort; NNSQ, Non-Motor Symptoms Questionnaire; NMSS, Non-Motor Symptoms Scale; N/S, not significant; Pcts- patients; PD, Parkinson's disease; PD-CN, Parkinson's disease with normal cognitive function; PDD, Parkinson's disease and dementia; PD-CCL, mild cognitive impairment in PD; PNIM, Carrier not manifest; PSS10, Perceived Stress Scale; RAVLT, Rey Auditory Verbal Learning Test; RBDO-REM, Sleep Behavior Disorder Questionnaire; SCA, Scale of Atomatic Function in PD; SDMT, Symbol Digit Modalities Test; TG, triglycerides; Trp, Tryptophan; UPDRS, Unified Parkinson's Disease Rating Scale; UPSIT, University of Pennsylvania Smell Identification Test; WHR, waist to hip ratio; XA, xanthurenic acid.
Table 2 - Results of the critical assessment of the included studies (Cohort) using the Joanna Briggs Institute approach.

| Studies | Q1 | Q2 | Q3 | Q4 | Q5 | Q6 | Q7 | Q8 | Q9 | Q10 | Q11 |
|---------|----|----|----|----|----|----|----|----|----|-----|-----|
| (7)     | Y  | Y  | Y  | Y  | Y  | U  | Y  | U  | U  | U   | Y   |
| (15)    | Y  | Y  | Y  | Y  | Y  | Y  | Y  | U  | U  | U   | U   |
| (6)     | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y  | U  | U   | Y   |
| (16)    | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y   | Y   |
| (17)    | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y  | U   | Y   |
| (9)     | Y  | Y  | Y  | Y  | U  | Y  | Y  | Y  | Y  | Y   | Y   |

N, no; NA, not applicable; U, unclear; Y, yes. Q1: Were the two groups similar and recruited from the same population? Q2: Were the exposures measured similarly to assign individuals to both the exposed and unexposed groups? Q3: Was the exposure measured using a valid and reliable method? Q4: Were confounding factors identified? Q5: Were strategies to overcome the confounding factors stated? Q6: Were the groups/participants free of the outcome at the start of the study (or at the time of exposure)? Q7: Were the outcomes measured using a valid and reliable method? Q8: Was the follow-up period reported and sufficiently long for outcomes to occur? Q9: Was follow-up complete, and if not, were the reasons for loss to follow-up described and explored? Q10: Were strategies to address incomplete follow-up used? Q11: Was an appropriate statistical analysis method used?

Table 3 - Results of the critical assessment of the included studies (Cross-sectional) using the Joanna Briggs Institute approach.

| Studies | Q1 | Q2 | Q3 | Q4 | Q5 | Q6 | Q7 | Q8 |
|---------|----|----|----|----|----|----|----|----|
| (18)    | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y  |
| (19)    | U  | U  | Y  | Y  | U  | U  | U  | U  |
| (20)    | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y  |
| (21)    | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y  |

N, no; U, unclear; Y, yes. Q1: Were the inclusion criteria clearly defined? Q2: Were the study participants and context described in detail? Q3: Was exposure measured using a valid and reliable method? Q4: Were objective and standard criteria used to measure the exposure in both patients and control individuals? Q5: Were confounding factors identified? Q6: Were strategies to overcome the confounding factors used? Q7: Were the results measured using a valid and reliable method? Q8: Was an appropriate statistical analysis method used?

Table 4 - Results of the critical assessment of included studies (case-control) using the Joanna Briggs Institute approach.

| Studies | Q1 | Q2 | Q3 | Q4 | Q5 | Q6 | Q7 | Q8 | Q9 | Q10 |
|---------|----|----|----|----|----|----|----|----|----|-----|
| (22)    | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y  | Y   |

N, no; NA, not applicable; U, unclear; Y, yes. Q1: Were the groups comparable, except for the presence of disease in patients and absence of disease in control individuals? Q2: Were patients and control individuals matched appropriately? Q3: Were the same criteria used for the identification of patients and control individuals? Q4: Was exposure measured using a valid, standard, and reliable method? Q5: Was the same method used to measure the exposure in both patients and control individuals? Q6: Were confounding factors identified? Q7: Were the same method used to measure the exposure in both patients and control individuals? Q8: Were outcomes assessed using a standard, valid, and reliable method in patients and control individuals? Q9: Were strategies to overcome the confounding factors stated? Q8: Were the exposure measured using a valid and reliable method? Q10: Was the exposure measured using a standard, valid, and reliable method in patients and control individuals? Q9: Was the exposure measured using a valid and reliable method? Q10: Was the exposure period sufficiently long to be meaningful? Q10: Was an appropriate statistical analysis method used?

found that a high-quality diet or a healthy eating pattern was associated with a lower risk of developing PD.

In the search for the present review, no studies were identified associating diet, MS, and PD. However, the relationship between diet and PD can be explained by the findings of a study performed in mice (24); the study found that long-term high-fat diet suppresses receptors activated by Peroxisome proliferators, increases inflammation and gliosis, and decreases dopaminergic neurons and dendritic spines in the midbrain black substance. In addition, PD mice showed neurodegenerative deficits, high anxiety, and movement disorders. Thus, it appears that the changes induced by a long-term high-fat diet interfered with both the development of morphofunctional changes in the central nervous system to facilitate the onset of PD and the existing disease.

Regarding obesity, another human study found that the availability of dopamine D2 receptors (verified by positron emission tomography) was lower in obese individuals (n=10) than in control individuals (n=10). Individuals with lower D2 values had a higher BMI. Dopamine modulates motivation and reward circuits; hence, the deficiency of this neurotransmitter in obese individuals may consolidate pathophysiological nutrition to compensate for the alteration in the circuit (25).

A cohort study assessed the relationship between the levels of gamma-glutamyltransferase (γGT) and the risk of PD and the possible interaction between γGT and obesity or MS (15). It is already known that obesity, diabetes mellitus, and components of MS are related to the serum activity of γGT (26,27); moreover, γGT may be associated with the risk of PD development through neuroinflammation and oxidative stress, which are possible etiological factors of PD (28).

A study included in this review (15) evaluated secondary data of 6,098,405 individuals of both sexes (age: >40 years) who were followed up for an average of 6.4 years and found sex difference in the association between the serum level of γGT and the risk of PD. In men with higher serum γGT activity, the risk of developing PD was low; on the contrary, in women with higher serum γGT activity, the risk of developing PD was high regardless of age, income, BMI, smoking, alcohol consumption, and exercise level. Thus, obese women are more likely to develop PD. According to the authors, the low incidence of PD among obese men may be because of protective effect of uric acid on the development of PD (29).

In this sense, another study found a higher mean uric acid level in men with PD and MS (20). However, this study aimed to investigate the progression of PD in 1022 individuals of both sexes (individuals with MS: n=396 [mean age: 63.9 years] and individuals without MS: n=626 [mean age: 59.9 years]) using a cross-sectional design. To assess the progression of PD, we used the UPDRS, the official reference scale for assessing PD (30) and the SDMT, a cognitive test to assess the processing speed and attention of participants (31). The group with MS had higher UPDRS scores, indicating faster progression of the disease, than the group without MS; however, no difference was found in the SDMT scores between the groups. The authors suggest that treatment of MS may be a new approach to delay the progression of PD.

However, a cross-sectional study evaluating the association of MS with the occurrence of falls in 194 elderly individuals with PD (mean age: 73 years) (21) using the UPDRS found no relationship between any component of MS and the severity of PD or occurrence of falls. We believe that the small sample size affected the findings and cognitive performance because no difference was observed between individuals who reported falls and the control individuals. Similarly, a recent study found that executive dysfunction is an independent risk factor for falls in PD (32).

A cohort study investigated the effect of MS on cognition in 787 individuals with PD (16). All participants were more
Author Contributions
Souza APS, Barros WMA and Silva JML contributed to research conception, data collection, interpretation of results and critical review of the manuscript. Silva MRM, Silva ABJ, Fernandes MSS and Santos MERA contributed to data analysis and interpretation, and manuscript drafting. Silva ML, Carmo TS, Silva RKP and Silva KG contributed to manuscript drafting and critical review. Souza SI and Souza VON reviewed the manuscript.

References
1. Saklayen MG. The Global Epidemic of the Metabolic Syndrome. Curr Hypertens Rep. 2018;20(2):12. https://doi.org/10.1007/s11906-018-0812-z
2. Morrell JS, Lofgren IE, Burke JD, Reilly RA. Metabolic syndrome, obesity, and related risk factors among college men and women. J Am Coll Health. 2012;60(1):83-9. https://doi.org/10.1080/07448481.2011.582208
3. Yahia N, Brown CA, Snyder E, Cumper S, Langolf A, Trayer C, et al. Prevalence of Metabolic Syndrome and Its Individual Components Among Midwestern University Students. J Community Health. 2017;42(4):674-67. https://doi.org/10.1007/s10970-016-0360-5
4. Mercers MCD, Coelho JMF, Lua I, Silva DSE, Gomes AMT, Santana AIC, et al. Burnout syndrome and metabolic syndrome: a cross-sectional population-based study. Arch Environ Occup Health. 2021;76(5):266-74. https://doi.org/10.1080/19338244.2020.1819186
5. Yadav LC, Rani V, Deep C, Singh RK, Palle K. Oxidative Stress in Metabolic Disorders: Pathogenesis, Prevention, and Therapeutics. Oxid Med Cell Longev. 2016;2016:9137629. https://doi.org/10.1155/2016/9137629
6. Nam GE, Kim SM, Han K, Kim NH, Chung HS, Kim JW, et al. Metabolic syndrome and risk of Parkinson disease: A nationwide cohort study. PLoS Med. 2018;15(6):e1002640. https://doi.org/10.1371/journal.pmed.1002640
7. Roh JH, Lee S, Yoon JH, et al. Metabolic Syndrome and Parkinson’s Disease Incidence: A Nationwide Study Using Propensity Score Matching. Metab Syndr Relat Disord. 2021;19(1):1-7. https://doi.org/10.1089/met.2020.0060
8. De Pablo-Fernández E, Breen DP, Bouloux PM, Barker RA, Foltynie T, Warrier TT. Neuroendocrine abnormalities in Parkinson’s disease. J Neurol Neurosurg Psychiatry. 2017;88(2):176-85. https://doi.org/10.1136/jnnp-2016-314601
9. Sääksjärvi K, Knelt P, Männistö S, Lyytinen J, Heliovära M. Prospective study on the components of metabolic syndrome and the incidence of Parkinson’s disease. Parkinsonism Relat Disord. 2015;21(10):1148-55. https://doi.org/10.1016/j.parkreldis.2015.07.017
10. Seppi K, Ray Chaudhuri K, Coelho M, Fox SH, Katzenschlager R, Perez Lloret S, et al. Update on treatments for nonmotor symptoms of Parkinson’s disease—an evidence-based medicine review. Mov Disord. 2019;34(2):180-98. https://doi.org/10.1002/mds.27602
11. Lee A, Gilbert RM. Epidemiology of Parkinson Disease. Neurol Clin. 2016;34(4):955-65. https://doi.org/10.1016/j.ncl.2016.06.012
12. He R, Yan X, Guo J, Xu Q, Tang B, Sun Q. Recent Advances in Biomarkers for Parkinson’s Disease. Front Aging Neurosci. 2018;10:305. https://doi.org/10.3389/fnagi.2018.00305
13. Malfiehnert F, Seppi K, Poeae W. The Concept of Prodromal Parkinson Disease. J Parkinsons Dis. 2015;5(4):681-97. https://doi.org/10.3233/JPD-150685
14. Postuma RB, Berg D. Advances in markers of prodromal Parkinson disease. Nat Rev Neuro. 2016;12(11):622-34. https://doi.org/10.1038/nrneurol.2016.152
15. You O, Kim R, Jung YJ, Han K, Shin CM, Lee Y. Serum gamma-glutamyltransferase activity and Parkinson’s disease risk in men and women. Sci Rep. 2020;10(1):1258. https://doi.org/10.1038/s41598-020-8306-x
16. Peng Z, Dong S, Tao Y, Hao Y, Zheng Z, Huang W, et al. Metabolic syndrome contributes to cognitive impairment in patients with Parkinson’s disease. Parkinsonism Relat Disord. 2018;55:68-74. https://doi.org/10.1016/j.parkreldis.2018.05.013
17. Denham M, Langlois M, Dupré N, Simard M. The influence of vascular risk factors on cognitive function in early Parkinson’s disease. Int J Geriatr Psychiatry. 2018;33(2):288-97. https://doi.org/10.1002/gps.4735
18. Thaler A, Shanhar-Tsfatfy S, Shaked Y, Gurevich T, Omer N, Bar-Shira A, et al. Metabolic syndrome does not influence the phenotype of LRRK2 and GBA related Parkinson’s disease. Sci Rep. 2020;10(1):9329. https://doi.org/10.1038/s41598-020-66319-9
19. Okenkrag G, van der Hart M, Roesser J, Sommergrad P. Peripheral Triphenyl - Kynurenine: Metabolism Associated with Metabolic Syndrome is Different in Parkinson’s and Alzheimer’s Diseases. Endocrinol Diabetes Metab J. 2017;14(4):http://researchopenworld.com/wp-content/uploads/2017/11/EDM-2017-113 Gregory-F-Okenkrag-A.pdf
20. Leehey M, Luo S, Sharma A, Wilks AA, Bambridge JL, Wong PS, et al. Association of metabolic syndrome and change in Unified Parkinson’s Disease Rating Scale scores. Neurology. 2017;89(17):1789-94. https://doi.org/10.1212/WNL.0000000000004572
21. Laidisio A, Lo Monaco MR, Vetrano DL, Bentivoglio AR, Bernabei R, et al. Association of metabolic syndrome with falls in patients with Parkinson’s disease. Clin Nutr. 2017;36(2):559-63. https://doi.org/10.1016/j.clnu.2016.02.004
22. van den Heuvel LL, du Plessis S, Stalder T, Acker D, Kirschbaum C, Carr J, et al. Hair glucocorticoid levels in Parkinson’s disease. Psychoneuroendocrinology. 2020;117:101404. https://doi.org/10.1016/j.psyneuen.2020.104704
23. Liu YH, Jensen GL, Na M, Mitchell DC, Wood GC, Still CD, et al. Diet Quality and Risk of Parkinson’s Disease: A Prospective Study and Meta-Analysis. J Parkinsons Dis. 2021;11(1):337-47. https://doi.org/10.3233/JPD-202260
24. Kao YC, Wei YW, Tsai KJ, Wang LC. High Fat Diet Suppresses Peroxisome Proliferator-Activated Receptors and Reduces Dopaminergic Neurons in the Substantia Nigra. Int J Mol Sci. 2019;20(23):570. https://doi.org/10.3390/ijms20(23)2016;34(4):955-65. https://doi.org/10.1016/j.bjpt.2020.11.005
25. Westerholm PJ, Voutilainen E, Erkkila A, Lanne T, Hu пи R, et al. Association of metabolic syndrome and risk of type 2 diabetes mellitus in men and women from the general population. J Intern Med. 2005;258(6):527-35. https://doi.org/10.1111/j.1365-2966.2005.01572.x
26. Lee DS, Evans JC, Robins SJ, Wilson PW, Albano I, Fox CS, et al. Gamma glutamyl transferase and metabolic syndrome, cardiovascular disease, and mortality risk: the Framingham Heart Study. Arterioscler Thromb Vasc Biol. 2007;27(12):127-33. https://doi.org/10.1161/01.ATV.0000251993.20724.0o
27. Poewe W, Seppi K, Tanner CM, Halliday GM, Brundin P, Volkmann J, et al. Parkinson disease and REV Dis Primers. 2017;3:15013. https://doi.org/10.1038/nrdp.2017.13
28. Davis JW, Grandinetti A, Waslien C, Ross GW, White LR, Korenz MH. Observations on serum uric acid levels and the risk of idiopathic Parkinson’s disease. Am J Epidemiol. 1996;144(7):480-4. https://doi.org/10.1093/oxfordjournals.aej.a008954
29. Goetz CG, Tilley BC, Shaftman SR, Stebbins GT, Fahn S, Martinez-Martin P, et al. Movement Disorder Society-sponsored revision of the Unified Parkinson’s Disease Rating Scale (MDS-UPDRS): scale presentation and clinimetric testing results. Mov Disord. 2008;23(15):2129-70. https://doi.org/10.1002/mds.22340
30. Roozendaal B. A Symbol digit modality test. West Psychol Serv Los Angeles. 1973;10:10.3389/fnagi.2018.00305
31. Pelicioni PHS, Menant JC, Henderson EJ, Latt MD, Brodie MA, Lord SR. Mild and marked executive dysfunction and falls in people with Parkinson’s disease. Braz J Phys Ther. 2021;25(4):437-43. https://doi.org/10.1016/j.fisioter.2020.11.005