Caffeine Exposure and the Risk of Parkinson's Disease: A Systematic Review and Meta-Analysis of Observational Studiess

João Costa^{a,b,*}, Nuno Lunet^{c,d}, Catarina Santos^{c,d}, João Santos^a and António Vaz-Carneiro^a

Abstract. Several studies conducted worldwide report an inverse association between caffeine/coffee consumption and the risk of developing Parkinson's disease (PD). However, heterogeneity and conflicting results between studies preclude a correct estimation of the strength of this association. We conducted a systematic review and meta-analysis of published epidemiological studies to better estimate the effect of caffeine exposure on the incidence of PD. Data sources searched included Medline, LILACS, Scopus, Web of Science and reference lists, up to September 2009. Cohort, case-control and cross-sectional studies were included. Three independent reviewers selected the studies and extracted the data on to standardized forms. Twenty-six studies were included: 7 cohort, 2 nested case-control, 16 case-control, and 1 cross-sectional study. Quantitative data synthesis of the most precise estimates from each study was accomplished through random effects meta-analysis. Heterogeneity was quantified using the l^2 statistic. The summary RR for the association between caffeine intake and PD was 0.75 [95% Confidence Interval (95%CI): 0.68–0.82], with low to moderate heterogeneity ($l^2 = 28.8\%$). Publication bias for case-control/cross-sectional studies may exist (Egger's test, p = 0.053). When considering only the cohort studies, the RR was 0.80 (95%CI: 0.71–90; $l^2 = 8.1\%$). The negative association was weaker when only women were considered (RR = 0.86, 95%CI: 0.73–1.02; $l^2 = 12.9\%$). A linear relation was observed between levels of exposure to caffeine and the RR estimates: RR of 0.76 (95%CI: 0.72–0.80; $l^2 = 35.1\%$) per 300 mg increase in caffeine intake. This study confirm an inverse association between caffeine intake and the risk of PD, which can hardly by explained by bias or uncontrolled confounding.

Keywords: Caffeine, meta-analysis, Parkinson's disease, relative risk, risk assessment

INTRODUCTION

Parkinson's disease (PD) is the second most common neurodegenerative disease with an estimated world-wide prevalence of 0.5 to 4% among the elderly [1]. The underlying neuropathological lesion is the degeneration of the pigmented neurons of the substantia nigra, locus caeruleus, and other brain stem dopaminer-

gic cell groups, with the subsequent loss of dopaminergic neurons terminals in the striatum. The continuous depletion of dopamine is responsible for most of the debilitating motor disturbances of the disease. The cardinal signs include bradykinesia, rigidity, rest tremor, gait disturbances, and postural instability [2].

There is not a single cause of PD, and multiple etiological factors with complex interactions are thought to be responsible for the development and progression of the disease [3,4]. The results of genetic and epidemiological studies suggest that genetic factors are particularly important in early-onset cases of PD [5,6] while the environmental component is probably more

^aCenter for Evidence-Based Medicine, Faculty of Medicine, University of Lisbon, Portugal

^bCentro de Investigación Biomédica en Red de Enfermedades Neurodegenerativas (CIBERNED), Spain

^cDepartment of Hygiene and Epidemiology, Porto University Medical School, Porto, Portugal

^dInstitute of Public Health of the University of Porto (ISPUP), Porto, Portugal

^{*}Correspondence to: João Parracho da Costa, MD, PhD, Centro de Estudos de Medicina, aseada na Evidência (CEMBE), Piso 3, Faculdade de Medicina de Lisboa, Av Prof Egas Moniz., 1649-028 Lisboa, Portugal. Tel./Fax: +351 217940424; E-mail: joaoncosta@sapo.pt.

relevant in the development of PD at older ages (above 50 years) [1].

There is a long list of environmental and lifestyle factors that have been associated with PD, either as risk or protective factors for the development of the disease. Infections, place of birth at early life, drinking wellwater, occupational exposure to welding, heavy metals or pesticides, and lack of vigorous exercise have all been referred to in the literature as putative risk factors [7–12]. On the other hand, smoking and consumption of coffee, tea, or nonsteroidal anti-inflammatory drugs are thought as possible protective factors [13]. Among all these factors, the most well studied in the literature are cigarette smoking and lifetime coffee consumption. In fact, the associations between smoking and coffee and lower risk of PD were first mentioned in the literature many years ago [14,15]. Since then, several large epidemiological studies conducted in the US, Europe, and Asia reported a dose-dependent inverse association between exposure to these factors and the risk of developing PD. These inverse associations were corroborated in family-based case-control studies, thus emphasizing smoking and caffeine as important covariates in any genetic or epidemiological studies of PD [16].

The strength of the evidence for the described inverse associations is weaker for coffee/caffeine than for smoking, because there are fewer studies and the magnitude of the effect is lower. A meta-analysis by Hernán and colleagues, published 8 years ago, found a polled relative risk of PD of about 60% and 30% lower among smokers and coffee drinkers in comparison to nonsmokers and non-coffee drinkers, respectively [17]. These results were based on a large number of studies (44 case-control and 4 cohort studies) for the smoking association, but on only 13 studies (8 case-control and 5 cohort studies) for coffee drinking. In addition, there is heterogeneity between studies results, and some of the studies published since then failed to show a significant negative association [18-21] or suggested significant differences between men and women, especially postmenopausal women on estrogen replacement therapy [22,23]. There are also conflicting findings in the few available data about the putative association of caffeine and the rate of progression of PD or the age of motor symptoms onset. Recent studies failed to identify any consistent relation either with the rate of progression [24,25] or the age of motor symptoms onset [26,

In view of the results of these more recent studies, we conducted a systematic review and a meta-analysis of the literature to quantify the association between caffeine intake and PD.

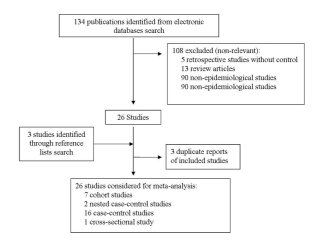


Fig. 1. Systematic review flow-chart.

MATERIALS AND METHODS

Search strategy and selection criteria

Potential eligible studies were identified through an electronic search of the databases Medline, LILACS (Latin America and Caribbean), Scopus and Web of Science (Fig. 1). The latest search of these databases was performed on September 2009. The search strategy for Medline combined the terms (coffee OR caffeine) with (Parkinson or Parkinson disease) together with a search filter developed for the retrieval of epidemiological studies (Cohort Studies OR Case Control Studies OR Prospective Studies OR Follow-Up Studies OR Cross-Sectional Studies OR Retrospective studies OR Epidemiological OR Incidence OR Risk Factors OR Risk Assessment OR Risk Reduction OR Relative Risk OR Behavior Regression Analysis OR Multivariate Analysis OR Proportional Hazards Models). All terms were searched as MeSH (Medical Subjects Headings) and free-text words. Moreover, the reference lists of relevant studies were cross-checked for potential additional studies not identified by the electronic search. We screened titles, keywords, and abstracts and obtained full copies of potentially suitable reports. There were no language restrictions and reports published as a full paper or abstract were considered as long as relevant data could be extracted.

The studies with cohort, case-control or cross-sectional designs that evaluated the relation between exposure to coffee/caffeine and the risk of PD (all diagnostic criteria were considered) or PD mortality were eligible for the systematic review. We excluded studies addressing the effects of short-term exposure to coffee

or caffeine and those that evaluated associations other than the risk of PD, such as the rate of progression. No studies were excluded *a priori* for weakness of design or data quality.

Data extraction

Three authors independently assessed the identified studies (JC, JS, and CS). Study details were obtained independently, written on predefined standardized forms, and cross checked for accuracy. Disagreements were resolved by consensus after repeated examination of the articles.

The information abstracted included the study characteristics (publication year, country of origin, study period, study design and length of follow-up), participant characteristics (number, age and gender), selection of cases and controls in case-control studies, assessment of coffee/caffeine intake and outcome (criteria for definition of PD or PD mortality), adjustment for potential confounders, and estimates of the association between different measures of coffee/caffeine exposure and PD.

When different risk estimates were available in the same publication, we opted for those that reflected the greatest degree of control for potential confounder, to the largest number of categories of exposure among caffeine consumers, or to the most comprehensive assessment of caffeine intake, applying these criteria consecutively. If results were provided separately for different caffeine-containing beverages or food items we opted for those referring to coffee consumption. Stratumspecific Relative Risk (RR) estimates [according to gender, use of Hormonal Replacement Therapy (HRT), or genetic polymorphisms related to caffeine metabolization] were extracted whenever available. Ross and colleagues [28] provided adjusted RRs but the highest category of exposure was used as reference and crude RR estimates were computed using the lowest exposure as reference. The crude estimates, however, were not meaningfully different from the adjusted ones. Ascherio and coworkers [29] provided RR estimates for both coffee and caffeine intake. The latter was provided graphically and the former was extracted, but there were no meaningful differences in the RR estimates per exposure level for coffee and caffeine.

Three studies [15,30,31] had matched case-control designs and did not provide Odds Ratio (OR) estimates for the association between caffeine intake and PD, or the data necessary for the calculation of valid estimates. Since the OR for drinkers vs. non-drinkers computed

using information from these studies is available in the meta-analysis by Hérnan et al. [17], which reported to have contacted the authors for additional information, we used the estimates they computed. Haack and collaborators [32] provided the OR for drinkers vs. non-drinkers in their report but it is slightly different from the provided by Hérnan et al. [17] and we used the latter in our meta-analyses.

When there was more than one publication for the same study, we used the one providing more detailed information on the relation between coffee/caffeine intake and PD, using the same criteria applied when more than one estimate was available from the same study, or referring to the longer follow-up (for cohort designs).

The samples evaluated by Ascherio and colleagues in 2001 [29] and in 2003 [33] overlap partially and we used the results referring to males presented in the study published in 2001, as these are not available in the 2003 study, and the results referring to females published in 2003.

Data synthesis

Each study is summarized in Table 1 and Fig. 2. The forest plot corresponding to Fig. 2 represents the RR estimates provided in each study for the association between caffeine intake and PD. Several estimates from the same study may be provided, referring to different exposure levels or to stratum-specific analyses.

Quantitative data synthesis was accomplished through random effects meta-analysis (DerSimonian and Laird method) (Fig. 3). Relative risks (cumulative incidence ratios or incidence density ratios) and ORs were treated the same and are referred to as RR. A cumulative random effects meta-analysis (Fig. 4) was conducted to allow a better understanding of the time trends in the understanding of the relation between caffeine intake and PD.

Summary estimates for exposure to caffeine were computed considering the individual RR estimates corresponding to coffee, coffee, and tea or caffeine intake (from caffeinated beverages or caffeinated beverages and chocolate), as available from each article, under the assumption that coffee is the main contributor for caffeine intake.

Since more than one RR estimate was available from several studies, only the most precise measures of association were used from each report (except for stratum-specific estimates, which were considered separately as if obtained from different studies). This criterion was followed for selection of a single estimate per study

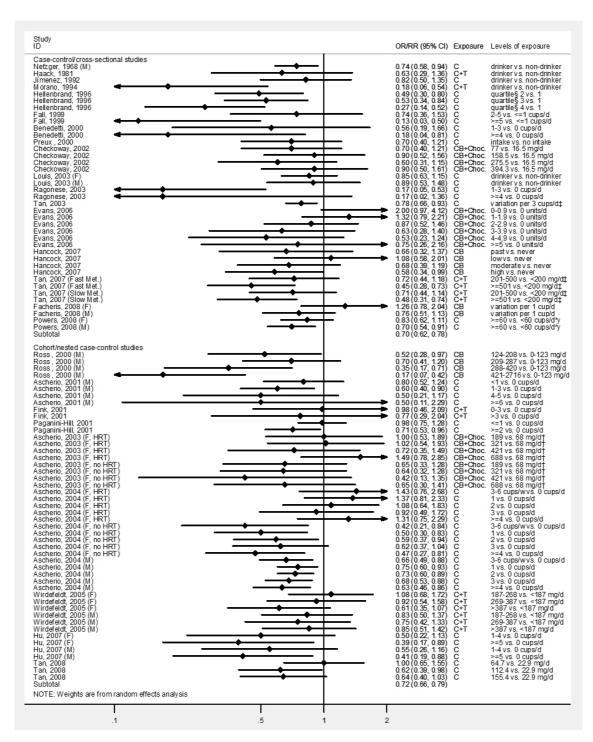


Fig. 2. Relative Risk estimates for the association between caffeine and Parkinson's disease, according to sources of caffeine intake and levels of exposure. Legend: ID – Identification; OR/RR – Odds Ratio/Relative Risk; M – Male; F – Female; C – coffee; C – tea; C – coffee and tea; C – caffeinated beverages; C – caffeinated beverages and chocolate; C – Fast metabolizers; C – Slow metabolizers; C – Hormonal Replacement Therapy; C – day; C – week; C – the exposures correspond to the median of each fifth of the distribution; C – consumption in C – C – C – levels of exposure not further specified

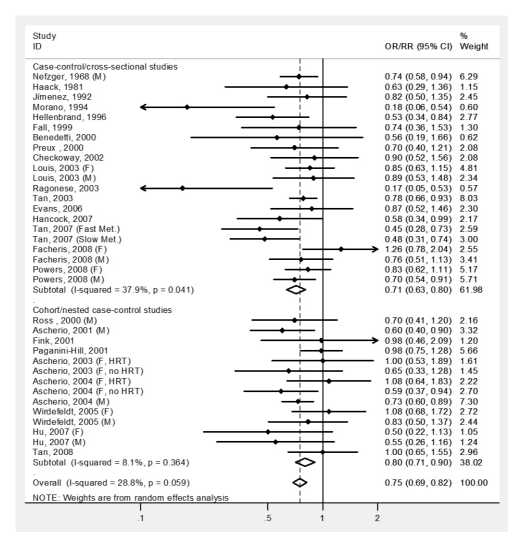


Fig. 3. Meta-analysis for the association between caffeine and Parkinson's disease, including the most precise RR estimates from each individual study. Legend: ID-Identification; $OR/RR-Odds\ Ratio/Relative\ Risk$; M-Male; F-Female; F ast I metabolizers; I slow I metabolizers; I met

when RRs were provided for different categories of exposure. If the precision of RR estimates was the same for more than one category we conservatively chose the one corresponding to the RR closest to 1.

The dose response relation between caffeine intake and PD was assessed through visual inspection of a scatter plot representing the RR estimates from each study (in a log scale) according to the exposure to caffeine (Fig. 5), and quantified by weighted least squares regression (WLS). All the RR estimates (for each level of exposure and for each stratum-specific analysis) obtained from studies providing RR estimates for at least two categories of exposure compared with the referent were plotted and included in the regression model.

This information was obtained from 15 studies [19,20, 22,23,26,28,29,33–40], corresponding to 69 RR estimates. The exposures corresponding to each RR estimate were those provided by the authors (e.g. median of each distribution quantile) or assumed to correspond to the midpoint of each index category range subtracted by the midpoint of the reference category range. For this purpose, we assumed that the open-ended upper category had the amplitude of the preceding stratum. The caffeine intake corresponding to each category of exposure or the information to compute it was provided by most studies. For three studies conducted in the USA [22,26,36] we assumed that a cup of coffee corresponds to 137 mg of caffeine (based on the estimates

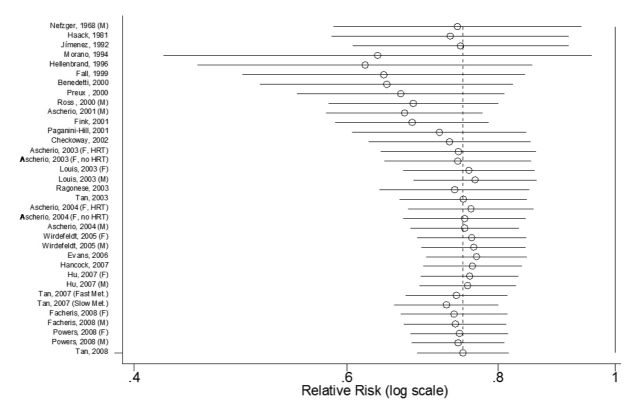


Fig. 4. Cumulative meta-analysis for the association between caffeine and Parkinson's disease, including the most precise Relative Risk estimates from each individual study. Legend: M – Male; F – Female; Fast Met. – Fast metabolizers; Slow Met. – Slow metabolizers; HRT – Hormonal Replacement Therapy.

used in other studies conducted by Ascherio et al.) and for one Italian study [37] the caffeine contents of a cup was assumed to be 75 mg (under the assumption that espresso coffee was more frequently consumed in this setting).

In all analyses heterogeneity was quantified using the I² statistic [41]. Publication and publication-related biases were examined through visual inspection of the funnel plot (Fig. 6). The Begg adjusted rank correlation [42] and the Egger's regression asymmetry test [43] were used for further assessment of these biases through hypothesis testing. All analyses were conducted with STATA®, version 9.2.

RESULTS

Systematic review

The search yielded a total of 134 reports (Fig. 1). A total of 26 epidemiological studies met criteria for inclusion in the systematic review, including 7 cohort [22,

23,28,29,33,35,40,44], 2 nested case-control [20,36], 16 case-control [15,16,18,19,21,26,30–32,34,37–39, 45–47] and one cross-sectional study [48]. The main characteristics of the studies and the respective results on the relation between caffeine intake and cognitive impairment are summarized in Table 1 and Fig. 2.

The publication year ranged from 1968 to 2008. The studies were conducted mainly in the USA (13 out of 26 [15,16,19,21,22,26,29,32,33,36,47,48], one of which in an Asian population [28]); in Europe (two in Spain [30,31], two in Sweden [20,?], one in Finland [23], one in France [18], one in Germany [45], one in Italy [37], one in the United Kingdom [38]); and in China [39,40,46].

Among the case-control studies, information on caffeine intake obtained from proxies or exclusion of cognitively impaired subjects was referred to in 4 [21,32, 34,46] and 3 reports [19,38,45], respectively. The study by Louis and colleagues [48] used both these strategies to minimize information bias. An accurate definition of the study base is not always possible with the information provided by the authors, but at least 5 were hospital based [15,30,31,37,38]. In cohort designs, the

Table 1

Main characteristics of the studies included in the systematic review

1	7		Evaluation of exposure	
First author	Lype of study	Outcome assessment	Timing of exposure	
Country	Sample characteristics		•	Control of confounding
Publication year	Follow-in (Cohort studies)	Definition of Parkinson Disease/Parkinsonism	Validation of the method	
			Items evaluated about caffeine exposure	
COHORT STUDIES	IES			
Ross	Cohort Study	Review of hospital records, local neurologists	24-hour recall methods and food frequency	
USA	Japanese-American men	After 1991, 3-step process:	questionianes	
2000 130 441	(Honolulu-Asia Aging Study)	1. Self declared diagnosis of PD (structured	1 week before	
7000 [70, 44]	Age: 43-08 y M/F: 8,006 (all M)	2. Evaluation by trained technician; recognition	Validated	
		of PD clinical signs (tremor, gait disturbances,		Age and Smoking
	Follow-up: Direction, 27 v. (moditar): Completeness: MS	bradykinesia) 2. Pofowol to gitudy normalogist - oritonia for DD	Coffee, tea (green and black), other caffeinated	
	Durauon. 27 y (median), Compreteness. No	5. Neteria to study neurologist – criteria for f.D. diagnosis (consensus from 2 neurologists):	beverages, and caneme non onder sources.	
		Progressive disorder		
		Any two of: marked response to levodopa, asymmetry at onset, or initial onset tremor		
	Cohort Study	Absence of office possible cause		
	fance atomorphisms	: :	SFFO (mail)	
Ascherio	Male health professionals	Questionnaire (mail)		
431	(Health Professionals' Follow-Up Study)	Confirmation of diagnosis with the treating	1 year before	Age, smoking, alcohol, BMI,
OSA	Age: 40-73 47,351: all M	neurologist or by review of the medical records At least 2. Tremor rigidity, bradykinesia	Validated	physical activity
2001 [29]	Follow-in:	Response to levodopa	Coffee. tea. cola and chocolate	
	Duration: 9.2 y (mean); Completeness: > 97%			
	Cohort Study		Other other model and control of	
Fink	Participants in the Original Framingham Study who attended the 12th.		Suucinied questionnaire	
	17th or 22nd biennial examination	Physical examinations	NS	:
USA 2001 1351	Age: 69 y (mean) M/F: 2,382 / 3,746	Tremor, rigidity, bradykinesia Absence of other possible cause	NS	Age, gender, smoking
	Follow-up: Duration: 10 v (for each index examination): Completeness: NS		Coffee, tea	

Table 1, continued

th Study) The Parson Parson of the model of the recent		Cohort Study			
refue (Natures) Health Study) And the registron than so in 11 studies Control May 2015 of 12 studies of the registron than so in 11 studies Control May 2015 of 12 studies of the registron than so in 15		Colloct State)	Organization of the Company of the C	SFFQ (mail)	
Age; 26.55 Age	Ascherio	Female registered nurses in 11 states	(man)	-	Age, smoking, alcohol, age at
Follow-up: Cohert Study (Cincer Percention Study II) Follow-up:	115.4	(Nurses' Health Study)	Confirmation of diagnosis with the treating	l year before	menopause, type of
Follow-up: Fol	VSO	9.434 person-vears): all	neurologist or by review of the medical records	Validated	contraceptives, and hormone
Follow-up: Cohort Study Age: 239 y (medium: 75 for M; 56 for P) Age: 230 y (medium: 57 for M; 56 for P) Age: 230 y (medium: 57 for M; 56 for P) Age: 230 y (medium: 57 for M; 56 for P) Age: 230 y (medium: 57 for M; 56 for P) Age: 230 y (medium: 57 for M; 56 for P) Age: 230 y (medium: 57 for M; 56 for P) Age: 230 y (medium: 57 for M; 56 for P) Age: 230 y (medium: 57 for M; 56 for P) Age: 230 y (medium: 57 for M; 56 for P) Age: 230 y (medium: 57 for M; 56 for P) Age: 230 y (medium: 57 for M; 56 for P) Age: 232 y (medium: 57	2003[33]		At least 2: Tremor, rigidity, bradykinesia		use or duration of use
Cohort Study	,	Follow-up: Duration: 18 v (mean); Completeness: > 98%	Response to levodopa	Coffee, tea, cola and chocolate	
Colort Study Colo		Cohort Study		-:	
(22) Cancer Percention Study 1) Authors 12 (22) Completeness: 100% (Cohert Study Orthochards Study Completeness: 100% (Cohert Study Orthochards Study Orthochards Study Orthochards Study Orthochards Study Orthochards Study Canalidate Institution is Register (Cohert Study Orthochards Study Orthochards Study Orthochards Study California) (Cohert Study Orthochards Study California) (Cohert Study Orthochards S	Accharia	American Cancer Society volunteers	DD as underlying or contributing cause of death	Structured questionnaire	
Age: 230 y (modeline: 57 for Mt. 56 for F) National Death Index	ASCILICITIO	Cancer Prevention Study II)	The as underlying of continuing cause of ucau		
Maironal formation: 1984-1988; Completeness: 100% National Social Insurance Institution's Register Cohort Study	USA	Age: ≥30 y (median: 57 for M; 56 for F)	National Death Index	2	Age, smoking, alcohol
Follow-up: Follow-up: Pollow-up: Pol		M/F: 301,164 / 238,058		NS	
Cohort Study 4 cross-sectional population surveys in 1982, 1987, 1992 and 1997 Consultant (usually specialist in neurology) National Social Insurance Institution's Register Age: 25-44	2004 [22]	989-1998: Completeness:	Idiopathic Parkinson's disease (ICD 9th revision)	Coffee, tea and sodas	
4 cross-sectional population surveys in 1982, 1987, 1992 and 1997 Age: 32-74 y Age: 32-74 y Age: 32-74 y Age: 32-74 y Article 1902 and 1997 Consultant (usually specialist in neurology) Bollow-up: Cohort Study Cohort Study Cohort Study Cohort Study Cohort Study Cohort Study california Nested Case-Control Study within a prospective cohort study of cases were study of cases were for study of cases were marked for age, gender, vital status and death date Age: 4-74 y (mean: 57 y) Age: 4-74 y (mea		Cohort Study	National Social Insurance Institution's Register	Self-administered questionnaire at home.	
Age: 25-74 y Motien lation; 12.9 y; Completeness: 74-88% Consultant (usually specialist) Cohort Study Cohort	Hu		data	1	Age, BMI, systolic blood pressure, total cholesterol.
Follow-up: Chinese, belonging to the two major dialect groups, and residue in government-built housing estates (Singapore Chinese Perlam Chinese, belonging to the two major dialect groups, and residue in government-built housing estates (Singapore Chinese Perlam Chinese, belonging to the two major dialect groups, and residue in government-built housing estates (Singapore Chinese Perlam Stardy) Age: 4744 y (mean: 57 y)	Finland		Consultant (usually specialist in neurology)	w Z	education, leisure-time
Follow-up: Cohort Study Cohort		210,01 / 0/2/11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Medical history, clinical examination (tremor,	NS	physical activity, smoking,
Cohort Study Ethnic Chinese, belonging to the two major dialect groups, and residing in government-built housing estates (Singapore Chinese Health Study) Health Study. Health Study Age: 45-71/956 / 35,262 Duration: 1993-2005; Completeness: 90% TED CASE-CONTROL STUDIES Cases and controls were selected from all residents that answered the health survey questionnaire sent by mail. Controls from the large cases vere evaluated by a movement death date and sord as socious services and controls were selected from all residents that answered the cases were evaluated by a movement death date and sord as a death date and so discharge database and two hospital discharge database and controls were reached for age, gender, vital status and death date cases were evaluated by a movement arched for age, gender, vital status and death date cases were evaluated by a movement arched for age, gender, vital status and death date and two hospital discharge databases of cohort anothers for PD. Filton-up: Cases identified through review of hospital discharge databases of cohort members for PD. Resident of the CIS Nation the day and two hospital discharge databases of cohort members for PD. Resident of the cases were evaluated by a movement and two discharge databases of cohort members for PD. Resident of the cases were evaluated by a movement and two discharged discharged discharged discharged discharged discharged discharged databases of cohort members for pa	2007 [23]	Follow-up: Duration: 12.9 y; Completeness: 74-88%	bradykinesia, stiffness, etc) and other relevant diagnostic methods	Coffee and tea	arconol and rea consumption, and history of diabetes
Ethnic Chinese, belonging to the two major dialect groups, and residing in government-built housing estates (Singapore Chinese Health Study). Hore Age: 45-74 (im-person interviewer) Health Study. Health Study. MF: 27,956 / 35,262 Duration: 1993-2005; Completeness: 90% Nested Case-Control Study within a prospective cohort study of health survey questionmaire end bright survey questionmaire end bright survey questionmaire and death date cases / Ontroles: 395 (MF: NS) / 2,320 Age: 15-74 (im-person interviewer) interviewer) NS Saw of the cases were evaluated by a movement displace registrics NS Saw of the cases were evaluated by a movement displacement and severed the health survey questionmaire end of the CIS survey questionmaire sor by mail. Controls from the large cases / Controls: 395 (MF: NS) / 2,320 Age: 15-74 (im-person interviewer) interviewer) NS Saw of the cases were evaluated by a movement displace registrics NS NS Health survey questionmaire of cases identified through review of hospital review of death certificates and report of cases / Controls: 395 (MF: NS) / 2,320 Age: 75 y		Cohort Study	Follow-up interviews, nationwide hospital		
Health Study. Age: 45-74 y (mean: 57 y) Age: 75 y Age: 45-74 y (mean: 57 y) Age: 45-74 y (mean: 57 y) Age: 75 y Age: 75 y	Tan		discharge database and two hospital-specific Parkinson's disease registries	SFFQ (in-person interview at home made by a trained interviewer)	;
Health survey questionnaire send from all residents that an awared the health survey questionnaire send from the large color study were selected from all residents that answered the health survey questionnaire send from the large color study were supported from the large color study were succeeded from the large color study were matched form all residents from the large color study were succeeded to the succeeded from the large color study were succeeded to the succeeded from the large color study were succeeded to the succeeded from the large color study study that succeeded to the succeeded from the large succeeded from the large succeeded from the large succeeded from the large succeeded fr	i			SZ	Age, year of interview,
M.F. 27,956 / 35,262 Diagnosis criteria were those from the Advisory Pollow-up: Duration: 1993-2005; Completeness: 90% Council of the US National Institute of Council of the US National Institute of Neurological Disorders and Stroke Coffee, black and green tea and sodas	Singapore	Age: 45-74 y (mean: 57 y)	88 % of the cases were evaluated by a movement disorder enecialists/ namelogist	!	gender, dialect, smoking and
Follow-up: Duration: 1993-2005; Completeness: 90% TED CASE-CONTROL STUDIES Nested Case-Control Study within a prospective cohort study of 13,979 residents of Leisure World Laguna Hills (Leisure World Laguna Hills Leisure World Laguna Hills (Leisure Wor	2008 [40]	M/F: 27,956 / 35,262	Diagnosis criteria were those from the Advisory	Validated	
Nested Case-Control Study within a prospective cohort study of 13.979 residents of Leisure World Laguna Hills (Leisure World Laguna Hills (Lei		Follow-up: Duration: 1993-2005; Completeness: 90%	Council of the US National Institute of Neurological Disorders and Stroke	Coffee, black and green tea and sodas	
Nosted Case-Control Study within a prospective cohort study of 13.979 residents of Leisure World Laguna Hills (Leisure World 13.679 residents of Leisure World Laguna Hills (Leisure World Cases identified through review of hospital NS Isolated Cases identified through review of hospital NS Control age, gender, vital status and death date Cases /Controls: 395 (M/F: NS) / 2,320 Age: 75 y Coffee and tea	NESTED CASE	C-CONTROL STUDIES			
Cases and controls were selected from all residents that answered the flexible graph of death certificate and report of controls: 395 (M/F: NS) / 2,320 Cases // Controls: 395 (M/F: NS) / 2,320 Cases // Controls: 395 (M/F: NS) / 2,320 Cases // Controls: 395 (M/F: NS) / 2,320	Posserini Hill	Nested Case-Control Study within a prospective cohort study of 13-97 residents of Leisure World Laguna Hills (Leisure World Laguna Hills (Leis		Health survey questionnaire	
Cases and controls were selected from all residents that answered the learling urganises of the peath survey questionnaire sent by mail. Controls from the large cohort study were matched for age, gender, vital status and death date Cases /Controls: 395 (M/F: NS) / 2,320 Age: 75 y Cases /Controls: 956 (M/F: NS) / 2,320	r agamm-mm	Study, California).	Cases identified through review of hospital	NS	Age, gender, smoking,
color study were matched for any 2, 320 Controls: 395 (M/F; NS) / 2,320 Age: 75 y	USA	Cases and controls were selected from all residents that answered the	discharge diagnoses of conort members for PD, review of death certificates and report of	o P	alcohol, blood pressure
	2001 [36]	cohort study were matched for age, gender, vital status and death date Cases (Controls: 395 AMF: NS). / 230		Coffee and tea	
		Age: 75 y			

Table 1, continued

	Nested Case-Control Study			
			Questionnaire sent to the twins in 1967 or in	
Wirdefeldt	Cases were 476 (M/F: 230/246) twins (mean age of 75 y) identified through the Swedish Innatient Discharge Register and the Cause of	Swedish Inpatient Discharge Register and the	1973	Matched for age, sex, genetic and familial environmental
-	Death Register.	Cause of Death Register	NS	factors (twins)
Sweden	Two control groups: (1) randomly selected twins unrelated to the	:		
2005 [20]	cases matched for birth year, gender and questionnaire source of the exposure data (external control subjects: $n=3.380$); and (2) co-twins	Idiopathic Parkinson's disease (ICD criteria)	NS	Smoking, alcohol and educational level
	of the cases (co-twin control subjects; n=415 same-sex twin pairs). Mean Age: 75.3 years		Coffee and tea	
CASE-CONTROL STUDIES	OL STUDIES			
	Case-Control Study		In-narcon interview	
Nefzger	Cases were successive PD inpatients recruited from every neurologist		in person merupa	
115.4	(up to a maximum of 10 patients) in Veteran's hospitals throughout the Neurological examination	Neurological examination	NS	Matched for age
	Controls were the first patient registered after the PD case without	Clinical diagnosis (criteria NS)	NS	of the second
1968 [15]	psychiatric or extrapyramidal disease (matched for age) Cases/Controls: 198 (all M) / 198 Ann. NS (>550% had >65 v)		Coffee	
	Case-Control Study		In-person interview (proxy if physically unable to	
Haack	Construction accomples of flavors associated accomples of DD accomples for a		answer questions)	
	cases were recruited from metrical records of FD cases seen by a neurologist in central Kentucky	Neurological examination	SZ	Matched for age, gender and
USA	Controls were identified trough door-to-door in neighbourhood	At least 2. brodylinesis resting tramor rigidity	2	race
1981 [32]	(matched for age, gender and race).	A reast 2. Otacy antesia, resume a curot, rigiany.	NS	
	Age: 65 y (range, 25-89)		Coffee and tea	
	Case-Control Study		Personal interview assessing coffee drinking	
Jiménez	Cases were unselected PD patients recruited from an outpatient		habits	
	movement disorder clinic (Madrid)	Neurological examination	S weare hefore	Matched for age and gender
Spain	Controls were patients presenting in the emergency room at the same hospital complaining of minor non-neurologic ailments (matched for	Clinical diagnosis (criteria NS)	3 2000	Smoking and alcohol
1992 [31]	age and gender).	· ·	N.)
	Cases/Controls: 128 (M/F: 68/60) / 256 Age: 65 v		Coffee	
	Case-Control Study		Orrodizanciis	
Morano	Cases were unselected outpatients making the first visit to one of two		Çuestionnane	
	neurology clinics (Cáceres)	Neurological examination	NS	
Spain	Controls were patients presenting in the emergency room at the same		5	Matched for age and gender
1994 [30]	hospital complaining of minor nonneurologic ailments (matched for	Clinical diagnosis (criteria NS)	N.	
lec' Loca	Lases/Controls: 74 (M/F: 33/41) / 148		Coffee and tea	
	Agr. 00 y			

Table 1, continued

	Case-Control Study		SFFQ (in-person interview made by a trained	
Hellonbuond	Good was all DD notionty accomited from nine Common namedowic		interviewer)	
пепепогапа	cases were all FD pauents recruited from filling definition (and 465 y) older notions were not recruited in order to	Naurological examination	Before diamosis of DD (cases) or 1 was hefore	Matched for age and gender
German	minimize memory deficits)	iveniological examination	frontrols)	
	Controls were randomly selected from the same neighbourhood or	UK Brain Bank criteria		Smoking, education and total
1996 [45]	region		Validated	energy intake.
	Cases/Controls: 342 (M/F: 224/118) / 342 Age: 56 y		Coffee and tea	
	Case-Control Study	Neurological examination	SFFQ (mail - proxy information when patients	
Fall	Cases were selected from prescription records and medical reports.		seemed directed disease property)	
Sweden	Controls were randomly selected from population registry in the same	At least 1: hypokinesia, tremor, rigidity. No earlier freatment with neuroleptic drugs.	15 years before	Age, gender, smoking, alcohol, occupation/exposure.
	area (Central Health Care District in Ostergotland County).	Response to levodopa	SIN	food factors
1999 [34]	Cases /Controls: 113 (M/F: NS) / 263	Progressive course Absence of atvnical features	22	
	Age: 40-75 y (mean age: 63 y for cases and 57 for controls)	15	Coffee and tea	
	Case-Control Study	Neurological revision of medical records (ICD codes) and full neurological examination in 27%	Neurological revision of medical records	
Benedetti	Cases were recruited from records linkage system of the Rochester	of the cases.	NS	Matched for one and nender
USA	Epidemiology Project. Controls were randomly recruited from the community (matched for	At least 2: bradykinesia, resting tremor, rigidity,	Validated by telephone interview to a subsample	Marched for ago and general
2000 [26]	gender and age). Cases/Controle: 106 (M/F: 121/75) / 106	postural instability Absence of other possible cause or atypical	of participants (direct and proxy interview)	Smoking, Alcohol
	Age: 71 y (range, 41-97)	features Response to levodopa	Coffee	
	Case-Control Study			
Prelix	Cases were PD inpatients and outpatients of the Limoges University		Standard questionnaire (physician personal interview)	Matched for age and gender
	Hospital. Controls were innotients and outpotients from other bosmital	Neurological examination	Lifatima	
France	departments (matched for age and gender).	1. d d d d	FIEGUIE	Age, smoking, PD familial
2000 [18]	Cases and controls had to live in the region of Limousin for at least 20	UN Brain Bank criteria	NS	history, urban area, toxic
	years. Cases /Controls: 140 (M/F: 74/66) / 280		Coffee and tea	products
	Age: 71 y			
	Case-Control Study	N	Structured in-person questionnaire by a nurse	
Checkoway †	Cases selected from diagnosis logs at neurology and general clinics	neurological examination of neurological panel review of charts	practitioner at subjects home	
	and from pharmacy database (Washington); MMSE to establish		During most of adult life	Age, gender, ethnicity and
USA	cognitive performance.	At least 2: bradykinesia, resting tremor, rigidity,		education.
2002 [19, 61]	Controls were from health cooperative enrolees (matched for gender, age, geographic location and year of enrolment)	postural instability Absence of other possible cause or atypical	NS	
	Cases/Controls: 210 (M/F: 131/79) / 347 Age: 71 v (range, 37-88)	features	Coffee, tea, cocoa, cola drinks and chocolate	

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Ragonese	Case-Control Study	Neurological examination	Structured questionnaire	
Italy	Cases were consecutive outpatients at neurological clinics. Controls randomly selected from population records of the	At least 2: bradykinesia, resting tremor, rigidity, postural instability	Years of coffee consumption (0; 1-40; >40)	Education, smoking and
2003 [37]	municipality (matched for gender, age and place of residence) Cases/Controls: 150 (M/F: 68/82) / 150	Unilateral onset or asymmetry Response to levodopa	NS	alcohol
	Age: 60 y (range, 31-81)	Progressive course	Coffee	
Tan	Case-Control Study		Structured questionnaire (interview)	Matched for age, gender and race
Singapore	Cases randomly selected from movement disorders database.	Neurological examination	NS	
2003 [46]	= '	UK Brain Bank criteria	Validated (information from caregivers and family members)	Tea, alcohol, smoking, head injury, stroke, hypertension,
	Cases/Confrois: 200 (Wr: 115/85) / 200 Age: 65 y (range, 43-88)		Coffee and tea	presence of heart conditions, toxin exposure and farm dwelling
	Case-Control Study		SFFQ (mail)	
Evans	Cases were consecutive outpatients of Caucasian descent fulfilling Queen square brain bank criteria; MMSE to establish cognitive	Neurological examination	1 month before	Matched for age and gender
UK	performance. Controls were friends of participants, outpatients without PD and	Oueen Square Brain Bank criteria	Validated	:
2006 [38]	randomly recruited from a volunteer panel (matched for age and gender) Cases/Controls: 106 (M/F: 65/41) / 106		Coffee, tea, chocolate milk, caffeinated soft drinks and chocolate	Sensation seeking score
	Age: 65 y (range, 38-81) Family-based Case-Control Study		Structured questionnaire (telenhone)	
Homosoli	ranniy-based Case-Connot study		Structured questionnaire (telephone)	
напсоск	Cases recruited through physician- and self-referrals to an academic medical center clinic (Miami).	Neurological examination	At reference age, 10 and 20 years before the reference age	Age, gender, smoking and
2007 TSA [16]	Controls were siblings, spouses, parents of subjects, other branches of At least 2: bradykinesia, resting tremor, rigidity, family. Absence of atypical features.	At least 2: bradykinesia, resting tremor, rigidity. Absence of atypical features.	SN	NSAIDs
[01] 150	Cases/Controls: 356 (M/F: 235/121) / 317 Age: 66 y (mean, PD cases)		Coffee, tea and soft drinks	
Tan	Case-Control Study		Standard questionnaire with a semiquantitative food frequency section	
Singapore	Cases were consecutive patients diagnosed with PD by a neurologist. Controls were volunteers from similar geographical regions (matched	Neurological examination	NS	Age, gender and smoking
2007 [39]	for age, gender and race). Cases/Controls: 418 (M/F: 243/175) / 468	UK Brain Bank criteria	Validated	
	Age: 70 y (mean, PD cases)		Coffee and tea	

Table 1, continued

	Family-based Case-Control Study		Structured auestionnaire (telephone - self or	
			proxy for deceased or incapacitated subjects)	
Facheris	Cases were 604 (M/F: 336/238) patients (mean age of 65 y, range 32	Namological axamination		
2008	to 22) reterror sequentiany to the department of recurously of the Mayo Clinic (Rochester).	iventiligation evaluation	From birth to the age at onset	Age, gender, smoking and
	iblings (n=446); and (2) unrelated controls	Detailed protocol (not specified))	education
USA [21]			Validated	
	ethnicity, selected randomly from Medicare and Medicaid services (if			
	older than 65 y) or using random digit dialling (if younger than 65 y).		Coffee, tea and caffeinated sodas	
	Case-Control Study			
			Standardized, self-administered questionnaire.	
Powers	Cases recruited sequentially through movement disorder clinics of			
	NeuroGenetics	Neurological examination	Lifetime	Ana nandar athnicitu
USA	Research Consortium (New York, Oregon, Washington and Georgia).			emoling NCAIDs and state
	Controls were spouses and blood relatives of patients, and community UK Brain Bank criteria	UK Brain Bank criteria	NS	shoung, warner and state
2008 [47]	volunteers.			
	Cases/Controls: 1,186 (M/F: 790/396) / 928 Aoe: 69 v (range 25-97)		Coffee	
	1250.00 J (141156) 20 01)			
CROSS-SECTION	CROSS-SECTIONAL STUDIES			
	Cross-sectional evaluation of coffee, smoke and parkinsonism		Food frequency questionnaire	
Louis	•	Neurologic and neuropsychological examinations	1 moon backons	A my mandam affaminites
IISA	Participants in the Washington Heights-Inwood Columbia Aging		ı year betore	Age, gender, emmenty, smoking years of education
	Project cohort (random selection from healthy Medicare beneficiaries) At least 2: bradykinesia, resting tremor, rigidity, Mean age: 77 v	At least 2: bradykinesia, resting tremor, rigidity, nostural instability	Validated	and dementia
2003 [48]	M/F: 655 / 1,471	Formal money.	Coffee	

BMI - Body Mass Index; ICD - International Classification of Diseases; NS - Not specified; NSAIDs - nonsteroidal antiinflammatory drugs; PD - Parkinson Disease; SFFQ - Semiquantitative Food-Frequency Questionnaire; UPDRS - Unified Parkinson's Disease Rating Scale

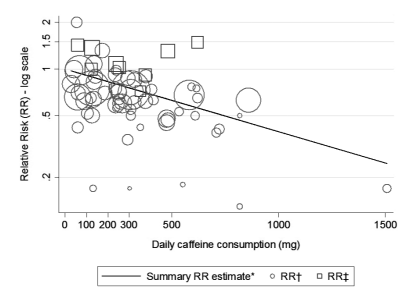


Fig. 5. Dose-response relation for the association between caffeine intake and the risk of Parkinson's disease. Legend: * Summary RR estimated by weighted least squares regression; † RRs for the comparison of each category of exposure with the reference category, obtained from each individual study; ‡ RRs for the comparison of each category of exposure with the reference category, obtained from the studies providing stratum-specific estimates for women under Hormonal Replacement Therapy [29,33].

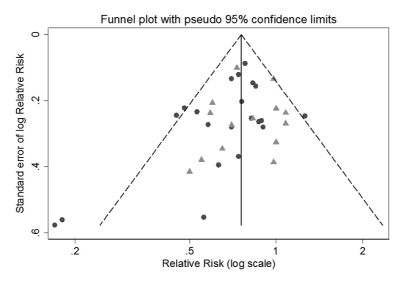


Fig. 6. Meta-analysis funnel plot, including the most precise Relative Risk estimates from each individual study. Legend: Circles – case-control/cross-sectional studies; Triangles – cohort/nested case-control studies.

estimated mean age of the participants at the time of baseline evaluation ranged from 42 to 77 years [33].

Different sources of caffeine were accounted for in the reports reviewed, and the results used for metaanalysis refer to coffee consumption in most studies (n=15), to coffee and tea consumption in 5 studies, and 6 studies extended exposure assessment to all caffeinated beverages or caffeinated beverages and products containing chocolate. Seventeen out of 25 studies provided RR estimates for different categories of exposure, with an estimated daily exposure to caffeine ranging from 27.4 mg to 1507 mg, and the reference categories including different proportions of non-caffeine consumers and consumers of different amounts of caffeine. From one study we used the RR estimate for the variation in the consumption of one cup of coffee per day. The remaining studies only compared drinkers with non-drinkers.

The clinical diagnosis of PD, based on a set of predefined clinical criteria, was the outcome in most studies. Information obtained from medical records and national medication or inpatient databases were occasionally considered as a complementary source in 6 studies [19, 20,28,29,36,40], as well as from death certificates in 4 studies [20,28,33,36]. In two studies [16,40] some patients had PD defined by self report and not confirmed by a clinical diagnosis, death certificates or medical records. One study [22] assessed PD mortality as the sole outcome.

Regarding potential confounding factors, smoking was considered taken into account in 7 studies [15,16, 19,20,31,32,38]. Exposure to heavy metals and use of pesticides or herbicides was accounted for by 2 authors [18,46]. Age and gender were controlled for in all studies except that by Ragonese et al. [37], either by stratified analysis, matching or multiple regression.

Meta-analyses

The summary RR for the association between caffeine intake and PD was 0.75 (95% CI: 0.68–0.82), with low to moderate heterogeneity ($I^2=28.8\%$). The summary RR estimates were homogeneous ($I^2=8.1\%$) and slightly higher among the cohort/nested case-control studies, and the I^2 was 37.9% among the case-control studies (Fig. 3). The negative association was weaker when only women were considered for analysis (summary RR = 0.86, 95% CI: 0.73–1.02, 9 estimates from 7 studies, $I^2=12.9\%$) than when only men were considered for analysis (summary RR = 0.72, 95% CI: 0.65–0.81, 9 estimates from 9 studies, $I^2=0.0\%$) or both genders were considered (summary RR = 0.68, 95% CI: 0.57–0.81, 17 estimates from 16 studies, $I^2=50.3\%$)

The search date of the previous most recent systematic review on the risk of PD and caffeine exposure was 2001 [17]. The results of the cumulative meta-analysis (Fig. 4) show that since year 2001 the number of studies on this topic nearly doubled, corresponding to 14 new published studies (5 cohort/nested case-control and 9 case-control/cross-sectional studies). The results of these new studies allowed us to calculate a total of 23 RR estimates that were included in the present meta-analysis and confirmed the observation of a consistent and robust association between caffeine intake and PD. The summary RR was 0.72 (95%CI: 0.61–0.84) at the end of 2001 and is currently 0.75 (95%CI: 0.68-0.82), with no meaningful variation in heterogeneity (I²: 26.6% in 2001 vs. 32.6% in 2008).

A linear relation was observed between levels of exposure to caffeine and the RR estimates (Fig. 5), corresponding to a summary RR of 0.76 (95% CI: 0.72–0.80) per 300 mg increase in caffeine intake, with moderate heterogeneity ($I^2=35.1\%$). Excluding the estimates corresponding to women under HRT from the studies by Ascherio et al. [22,33], the heterogeneity decreased ($I^2=27.6\%$).

Publication bias

The visual inspection of the funnel plot (Fig. 6) suggests that case-control/cross-sectional low precision studies yielding a positive association between caffeine intake and PD may be underrepresented in our meta-analysis, which is confirmed by the Egger's regression asymmetry test (p=0.053) and the Begg adjusted rank correlation test (p=0.037). On the other hand, for co-hort/nested case-control studies, the funnel plot is symmetric and there is no evidence of statistically significant publication bias (Egger's regression asymmetry test: p=0.821; Begg adjusted rank correlation test: p=0.412).

DISCUSSION

The present meta-analysis shows a 25% reduction in risk of PD among caffeine consumers. The results also indicate a linear dose-response relation, with higher intakes of caffeine being associated with a lower risk of PD.

From a biological point of view, caffeine (1,3,7trimethylxanthine) and its major metabolite, paraxanthine (1,7-dimethylxanthine), are antagonists of the adenosine A2A receptors. The expression of these receptors in the brain is particularly prominent in the striatum, which is the target of the dopaminergic neurons that degenerate in PD. Similar to other more specific A2A antagonists, caffeine attenuates neurotoxicity in experimental animal models of PD [49,50]. A recent study by Nakaso and collaborators provided further evidence for a possible neuroprotective effect of caffeine, showing that caffeine activates specific neuroprotection signaling pathways and prevents apoptotic cell death in a PD model using human dopaminergic neuroblastoma cells [51]. Therefore, there is a plausible rational biological mechanism based on the pharmacological actions of caffeine for the inverse association between coffee drinking and PD found in several epidemiological studies.

The primary candidate component that is believed to be responsible for the neuroprotective effect of coffee is caffeine. In fact, a negative association has also been reported for other non-coffee sources of caffeine, such as tea [46], but not for decaffeinated coffee [28]. However, coffee is a complex chemical mixture reported to contain more than a thousand different chemicals, including carbohydrates, lipids, nitrogenous compounds, vitamins, minerals, alkaloids, and phenolic compounds [52]. Thus, the possibility exists that other components of coffee or tea may also play a role. However, our systematic review was designed to test the effect of caffeine on the risk of PD under assumption that coffee is the main contributor to caffeine intake, and the specific effect of other caffeine containing beverages, such as tea, was not evaluated.

The negative association between caffeine intake and PD was consistent throughout different methodological approaches. Unlike those observed for cohort designs, the results from case-control/cross-sectional studies were somewhat heterogeneous, but consistently pointed to a protective effect, despite the observation that the strength of the association differed substantially across studies. Publication bias seems to have occurred for case-control/cross-sectional, but not for cohort/nested case-control studies, which may contribute to explain the stronger negative association observed among the former. Moreover, the homogeneity across results of cohort designs probably reflects a lower potential for bias with this methodological approach. Control selection is more likely to be biased in hospital-based case-control studies. Patients with PD may have an associated cognitive impairment [53], especially among older individuals, and this makes information bias more likely when exposure assessment is retrospective and exposure information is not collected from proxies. Also, patients with motor disability, such as in PD, may be less likely to drink coffee and this can only be accounted for with prospective designs or assessing exposure before the occurrence of the disease.

It has been suggested that PD patients may have a premorbid personality which may be responsible for particular addictive personality characteristics [54–58]. In PD, the progressive degeneration of the striatum with low endogenous dopamine and serotonin levels may lead to a low sensation seeking behavior (cause-effect bias) [59]. PD patients may therefore be less prone to smoke and drink coffee and alcohol, all lifestyle confounders with a potential neuroprotective or symptomatic effect in PD. Evans and coworkers [38] ad-

dressed this issue and raised the possibility of an existing neurobiological link between low sensation-seeking trait, which may underlie the parkinsonian personality, and the hypothetical protective effect of coffee in Parkinson's disease.

Cohort designs are less prone to information bias, but also have potential limitations related to the enrollment of non-inception cohorts and resulting from incomplete follow-up. Another potential source of bias is the definition of PD cases because of the lack of information regarding PD diagnoses in medical records and death certificates. Bias may also arise from exposure classification and quantification of coffee/caffeine consumption due to the different methods used in the studies, the low accuracy (recall bias) and reproducibility of the quantitative questionnaires, and the high variability of caffeine concentrations in coffee beverages.

It has been recognized that smokers have a lower risk of PD [17] and confounding by smoking habits is therefore an inherent problem when addressing the association between caffeine and PD. The majority of results available, however, were adjusted for smoking and other potential sources of bias, which makes confounding unlikely to be responsible for our conclusions.

The methodological options in our meta-analysis also need to be discussed. From studies presenting RRs for different categories of exposure we selected the most precise estimates to compute the summary RR for caffeine consumers vs. non-consumers, which allowed us to include all the available studies in the analysis. The precision of the individual RR estimates is not dependent on the direction of the association, and with this criterion the selection of the exposures corresponding to the largest number of participants is the most likely. However, if the categories of exposure in each individual study are defined to include a similar number of participants per group, this criterion leads to the selection of the estimates reflecting the weaker associations. This contributed to a slight underestimation of the summary RR, as well as an overestimation of homogeneity, especially for the cohort studies among which the definition of exposure categories with a similar number of participants was more frequent. The precision of the summary estimates, however, is underestimated by considering only part of the overall sample from each study in the meta-analysis.

For trend estimation we conducted a weighted linear regression adjusted through the origin, which implies the assumption of independence between all categories of exposure, an assumption that within each study is not met because all risk estimates depend on a common referent group, ultimately leading to an underestimation of the slope variance. This contributes to spurious precision of our estimates but allows the computation of point estimates less prone to bias as it allows inclusion of most studies providing information for different categories of exposure. The use of a method that allows the correction for the lack of independence across RR estimates for different exposures would lead to the exclusion of some of the studies, as it requires more information than is provided by many studies [60].

A meta-analysis conducted by Hernán et al., published in 2002, concluded that smoking habits and coffee intake were independently associated with a lower risk of Parkinson's disease [17]. Despite the different options for meta-analysis, our review included nearly twice more individual studies and reaches robust conclusions that confirm the negative association between caffeine and PD. Also, the present meta-analysis adds to the previous one by confirming a linear dose-response relation, as previously suggested by Hernán and colleagues. Recent studies aiming to evaluate interactions with hormonal replacement therapy in women or hepatic caffeine metabolization were included [22,33,39], contributing to a broader view of the problem.

Ascherio and collaborators suggested gender differences in the relation of caffeine intake and the risk of PD: in men, a strong inverse association was found, whereas in women a U-shaped relationship was observed, with the lowest risk occurring at moderate intakes [29]. These authors further investigated this difference in two different cohorts and found an interaction between the use of postmenopausal hormones and caffeine intake in the risk of PD, with an increased risk among women on hormonal replacement therapy with a high caffeine intake [22,33]. The use of postmenopausal estrogens seems to modify the effects of caffeine on the risk of PD, although the reasons for this interaction are not yet clear.

The individual variability in the metabolism of coffee compounds related to genetic polymorphisms was also recently addressed [39]. The main endogenous system responsible for caffeine metabolism in humans is the cytochrome P450 1A2 (CYP 1A2). The study conducted by Tan and coworkers stratified the results for CYP 1A2 genetic polymorphism and demonstrated a similar dose-dependent PD protective effect of caffeine in individuals with fast and slow metabolizing status [39].

In conclusion, our data confirm an inverse association between caffeine intake and the risk of PD, with a dose-response relation, and more consistency in cohort studies and among men, which cannot be fully explained by bias or uncontrolled confounding. The understanding of the mechanisms for the protective effect of caffeine exposure warrants further investigation in PD.

DISCLOSURE STATEMENTS

Authors' disclosures available online (http://www.j-alz.com/disclosures/view.php?id=266).

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