The association between childbirth and later Parkinson’s onset is a non-biological artefact of societal change.

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# Author note

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Abstract

**Background:** Uncontrolled studies have reported an association between later Parkinson’s onset in women and a history of giving birth, with age of onset delayed by nearly three years per child.

**Objectives:** We tested this association in two independent datasets but also examined men with Parkinson’s, as a control to test for non-biological explanations.

**Methods:** We analysed the Parkinson’s Progressive Markers Initiative incident sample (PPMI: 145 women, 276 men) and a prevalent sample surveyed by the New Zealand Brain Research Institute (NZBRI: 210 women, 394 men).

**Results:** The association was present in both women and men in the PPMI study, and absent in both in the NZBRI study. This is consistent with generational differences, common to men and women, which confound with age of onset in incident-dominant samples.

**Conclusion:** Despite being replicable in certain circumstances, the association between childbirth and later Parkinson’s onset is an artefact of generational cohort differences.

*Keywords:* Parkinson disease, sex differences, pregnancy, epidemiology, cohort effects, sex hormones

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# Introduction

The incidence and prevalence of Parkinson’s are lower in women than in men.1,2 The cause is unknown but could include increased exposure to risk factors for men, such as head injury or occupational use of toxins.3 Conversely, there could be protective factors that apply differentially to women, possibly including exposure to endogenous or exogenous sex hormones such as œstrogens and progestogens.

There are plausible mechanisms for how female sex hormones could play a neuroprotective role in neurodegenerative conditions like Parkinson’s.4 Many observational human studies have therefore examined whether there is evidence that increased hormone exposure actually protects against Parkinson’s. This requires operationalising an individual woman’s lifetime hormonal exposure level, using numerous measures such as fertile life span (the duration between age at menarche and age at menopause), the use of oral contraceptives, or a history of hormone replacement therapy. Despite some positive studies, however, a recent meta-analysis showed little epidemiological support for relationships between any such measures and a lower subsequent risk of Parkinson’s.5

Lv *et al.’s* meta-analysis included an examination of five case-control and three cohort studies of the effect of parity (number of childbirths) upon the risk of women subsequently developing Parkinson’s. This also showed no effect, with the relative risk of Parkinson’s between the highest and lowest number of births being 0.99 (95% CI: 0.79–1.25). Two studies have, however, reported another childbirth-related association that was not specifically addressed in that meta-analysis: among women who have been diagnosed with Parkinson’s, they found that each additional childbirth was associated with a later age of onset6,7. That is, although having children might not reduce the risk of developing Parkinson’s *per se,* it might at least slow the pathological process, leading to a later onset. The associations were of a surprisingly large magnitude: Haaxma *et al.*6 found symptom onset in women to be later by 2.7 years per childbirth (95% CI: 0.8 – 4.6) while Frentzel *et al.*7 independently found a delay of 2.6 years per birth (95% CI: 0.05 – 5.1). (In neither report was the effect as linear as these single coefficients might imply, however.)

A challenge in such observational studies can be applying valid control groups. For example, women without Parkinson’s lack the disease outcome measures (such as age of onset), while men with Parkinson’s lack the predictor hormonal measures. For the specific factor of number of children, however, men with Parkinson’s can and should be used as a control, to test for non-biological explanations. Therefore we sought to examine the association between number of children and Parkinson’s age of onset, in both men and women. If the relationship is absent in men, it would provide better evidence for a protective hormonal effect specific to women. If a similar relationship holds in men, it would strongly indicate a non-biological cause, common to both sexes.

# Methods

## Data sets

*PPMI:* We downloaded the openly-available Parkinson’s Progressive Marker Initiative dataset (PPMI, see Acknowledgments). PPMI is an incident study, with a mean 7 months (SD 7 months) between diagnosis and study entry (see Table 1). Data on participants’ number of children was available for 421 of the study’s 423 idiopathic Parkinson’s cases.

*NZBRI:* At the New Zealand Brain Research Institute (NZBRI) we conducted two online surveys on risk factors, in which 604 people with idiopathic Parkinson’s (210 women, 394 men) provided valid data on number of children and age of onset (see Table 1). The first survey (192 valid cases) recruited from our ongoing longitudinal study of a convenience prevalence sample from the local Canterbury region. The second survey (412 valid cases) was nation-wide, recruiting from the membership (excluding Canterbury) of the Parkinson’s New Zealand charitable trust. In both surveys, cases ranged from the recently-diagnosed to those with a long history of disease (those with dementia had assistance to complete the survey).

## Analyses

Data were fit with simple Bayesian linear models, using the *brms* package8 in the R statistical environment9. Parameters were given default weakly informative priors. Four chains of 4000 iterations each were run.

Disease onset was defined as the age at which a formal diagnosis of Parkinson’s was made. When testing hypotheses that there was an association between the number of children and age (of onset or birth), in each case we compared a model containing the number of children as a predictor to an intercept-only model (i.e. a model of no association, which simply estimated the mean age). Bayes factors indicated how much more likely it was that incorporating the number of children as a predictor provided a better fit to the data than an intercept-only model.

## Reproducibility

The code and the anonymised dataset extracts used to carry out the analyses and generate this manuscript are publicly available for inspection and comment at <https://github.com/nzbri/pd-parity>.

Table 1: Demographic characteristics of the PPMI and NZBRI samples.

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| --- | --- | --- | --- | --- | --- |
| Study | Sex | n | Mean age | Mean age at diagnosis | Mean years disease duration |
| PPMI | Men | 276 | 62.2 (9.7) | 61.6 (9.7) | 0.5 (0.5) |
| PPMI | Women | 145 | 60.7 (9.6) | 60.1 (9.6) | 0.6 (0.6) |
| NZBRI | Men | 394 | 71.7 (7.5) | 63.8 (9) | 7.9 (5.8) |
| NZBRI | Women | 210 | 70.3 (7.3) | 62.9 (9) | 7.5 (5.8) |

# Results

## PPMI sample

We first attempted to replicate the association between number of children and later disease onset in women in the PPMI sample. The linearly-modelled delay in diagnosis associated with children was 1.1 years per child (uncertainty interval [-0.01, 2.1]). That estimate is the slope of the line in the upper-right panel of Figure 1B, with a Bayes factor of 9.9 indicating substantial evidence for an association. Similar results were, however, found for the men in this study, with a slope of 1.3 years per child [0.3, 2.3] (see the upper-left panel of Figure 1B). A Bayes factor of 29.9 provided even stronger evidence for the association. That is, as the relationship between number of children and later disease onset was not unique to women, this strongly argues against the underlying cause being additional exposure to œstrogens.

We also tested whether women actually had a later age of disease onset. As indicated in Table 1, the mean age at diagnosis was actually slightly *earlier* for women, by -1.6 [-3.5, 0.4] years, with the posterior distribution implying a probability of only 0.06 of a later onset in the population.

## NZBRI sample

There was no evidence of an association between number of children and age of onset in the NZBRI sample. The slope for women was only 0.3 years per child [-0.6, 1.3], with a Bayes factor of 1.5 indicating no evidence for a positive association. For men the results were similar, with a slope of 0.3 years per child [-0.4, 1.0] and a Bayes factor of 1.3.

The mean age at diagnosis was marginally earlier for women in this sample, by -0.9 years [-2.5, 0.6], with a posterior probability of 0.12 of onset actually being later in the population.

## Testing a non-biological alternative explanation

Due to societal changes in many countries, older people are more likely to have had larger families than would people born more recently. Figure 1A shows there was indeed such a relationship, for the men and women in both studies (for ease of comparison with the age-of-onset effect, we plot number of children on the *x*-axis, although this does not reflect the direction of causality). We then modelled how many years earlier we would expect a patient to have been born, given how many children they had. *PPMI study.* Women: -1.1 years per child, uncertainty interval [-2.15, 0.0], Bayes factor 11.3. Men: -1.3 years per child [-2.32, -0.3], Bayes factor 31.4. *NZBRI study.* Women: -0.6 years per child, [-1.35, 0.2], Bayes factor 2.7. Men: -0.9 years per child [-1.49, -0.4], Bayes factor 98.2. (The coefficients are negative, as we are modelling year of birth, rather than age.)

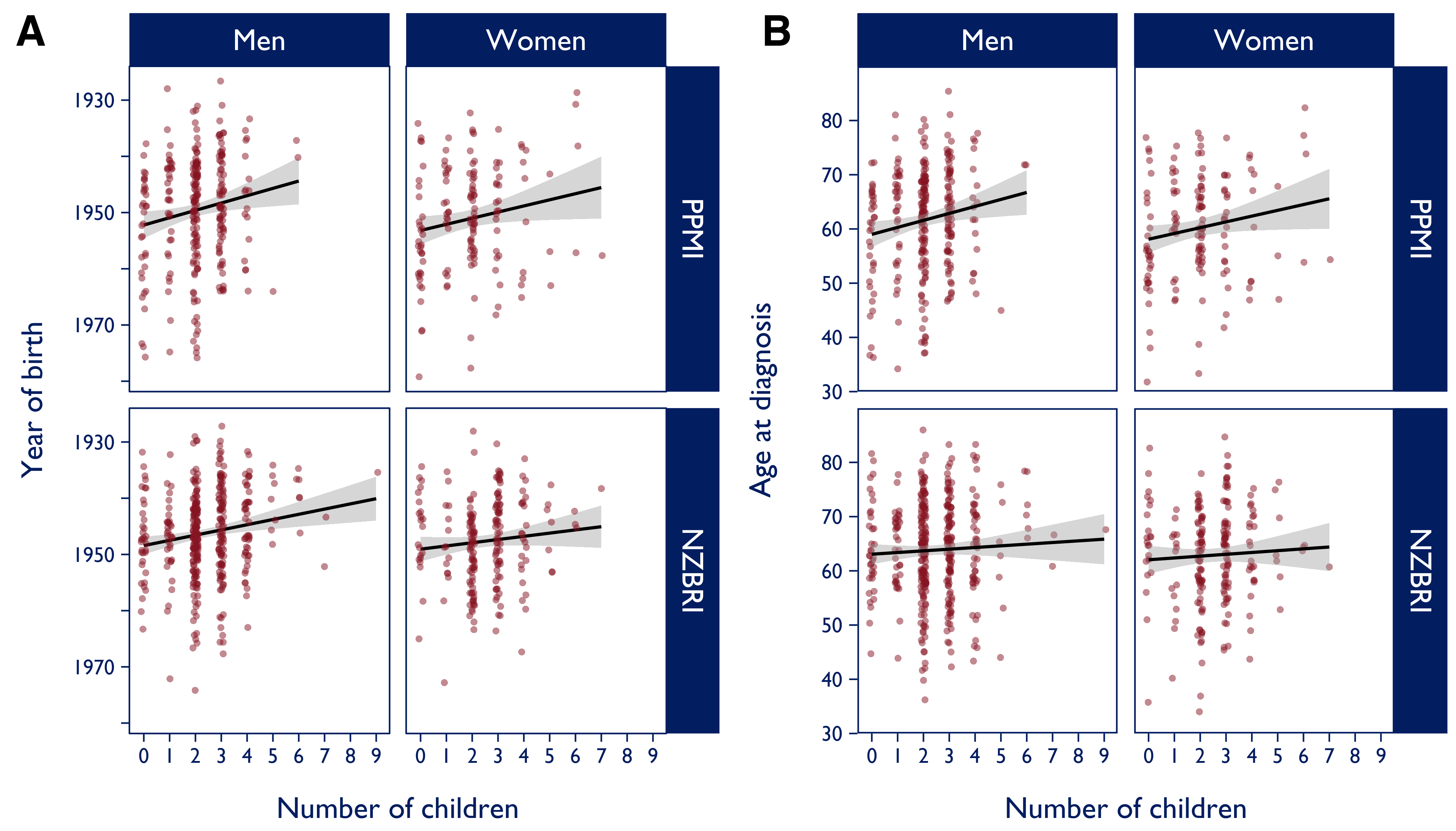


Figure 1: (A). In both the PPMI and NZBRI studies, in both women and men, there was a background cohort effect in which older patients tended to have had more children than patients born more recently. The magnitude of this effect (i.e. the slope of each line) was each additional child being associated with patients having been born (approximately) one year earlier. (B). The PPMI is an incident study and hence patient age and age-of-disease-onset are tightly correlated. Accordingly, the PPMI age-of-onset association almost perfectly reflects the underlying background societal association of older people having more children (at a similar magnitude of approximately one year per child). The NZBRI survey, however, was of a prevalent sample. The relationship between current age and age of onset therefore becomes de-coupled. Accordingly, the age-of-onset association was absent for both men and women in this sample.

# Discussion

The association between having more children and later disease onset in the men of the PPMI sample is sufficient to discount a hormonal cause for the relationship (and reinforces the need to examine relevant control comparisons whenever possible). We tested a non-biological explanation, in common to both men and women, in which older patients are simply more likely than younger ones to have had larger families. In an incident study like PPMI, patients’ ages, and their ages of disease onset, are almost perfectly correlated. Hence the diagnosis-age effect in the PPMI study (top row of Figure 1B) almost perfectly resembles the simple background generational cohort effect (top row of Figure 1A). The reported associations are therefore a classic example of the epidemiological flaw of mistaking an age effect for a cohort one.10

The association between number of children and diagnosis age was not found in either sex in the NZBRI study (bottom row of Figure 1B), although the background age effect (bottom row of Figure 1A) was similar to that seen in the PPMI study. This makes sense, because in a prevalence sample, the relationship between patient age and their age of onset becomes de-coupled, as the sample includes many people with longstanding disease. Take, for example, a pair of people, one with an age of onset at 50 and another of onset at 75. In an incident study, participants are recruited soon after diagnosis, and therefore this pair come from different generations: one would have been born approximately 25 years before the other. In a prevalent sample, however, that tight relationship breaks down. Both patients could even be the same age: one could have been recently diagnosed at age 75, while the other, also aged 75, would have a 25 year history of disease. We therefore predict that artefactual associations between number of children and the age of Parkinson’s onset are most likely to be observed in samples that have a preponderance of incident cases. In a pure incident sample, the magnitude of the association should closely match the background societal association between age and family size (in each case, roughly one year per child in the PPMI sample).

Lastly, we found that women’s age of onset was marginally *earlier* compared to men. This is not inconsistent with the two previous reports: in the Frentzel *et al.* sample7, the women’s age of onset (57.1) was similar to the men’s (57.3), while Haaxma *et al.*6 found it to be non-significantly later (53.4 vs 51.3, p = 0.06). At a national population level, further analysis of our previously reported drug-tracing epidemiological study2,11 of 10 500 [10 300,10 700] incident cases of Parkinson’s found that the standardised age of onset for women was again, if anything, slightly earlier for women (69.8 [69.6, 70.0]) compared to men (70.5 [70.5,70.6]). That is, even if the association between childbirth and later onset was not artefactual, there might not be a sex difference that it could be required to explain.

# Acknowledgments

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# TODO

* add uncertainty intervals on plots based on the models, as per the noisy prediction analysis (don’t need to worry about any random effects, only fixed effects are involved).
* “As per the PPMI Data Use and Biospecimen Use Agreements, investigators are reminded that they must submit all manuscripts and abstracts to the PPMI Data & Publications Committee (DPC) for administrative review prior to submission to a journal or conference. Review of submitted manuscripts and abstracts is guaranteed within one week of submission. To submit to the DPC, please upload the publication for review via the PPMI website.”
* Submit figure to figshare.com and cite in caption.

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