Impact of parental rheumatoid arthritis on risk of autism spectrum disorders in offspring: A systematic review and meta-analysis

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Background: To investigate the association of risk of offspring autism spectrum disorder (ASD) with both maternal and paternal rheumatoid arthritis (RA).

Methods: The Embase, Medline, Cochrane Library databases were searched for studies that investigated the association of parental RA with risk of offspring ASD. The primary outcome was the associations of maternal/paternal RA with the risk of offspring ASD. Subgroup analyses were conducted based on the timing of maternal RA diagnosis (i.e., before/after childbirth) and geographical location (i.e., Western vs. Asian countries) of studies.

Results: Ten studies published between 2005 and 2022 involving 6,177,650 participants were analyzed. Pooled results revealed a significant association between maternal RA and the risk of ASD (OR = 1.246, p < 0.001, 10 studies), while there was no association of paternal RA with the risk of offspring ASD (OR = 1.104, p = 0.253, four studies). Subgroup analysis demonstrated no correlation between diagnosis of maternal RA before childbirth and the risk of offspring ASD (OR = 1.449, p = 0.192, four studies), while there was a significant association of maternal RA regardless of the timing of diagnosis with the risk of offspring ASD (OR = 1.227, p = 0.001, six studies). Subgroup analysis on geographical location showed a significant association of maternal RA with the risk of offspring ASD regardless of the study location (all p < 0.05).
Conclusion: Our findings supported an association between maternal RA and an elevated risk of ASD in offspring. However, given the limited numbers of studies investigating the risk of offspring ASD in mothers diagnosed with RA before childbirth, further studies are warranted to elucidate this issue.

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KEYWORDS
autism spectrum disorder, rheumatoid arthritis, meta-analysis, offspring, autoimmune diabetes

Introduction

Autism spectrum disorder (ASD), a type of neurodevelopmental disorder, is characterized mainly by deficits in social communication and the presence of repetitive and restrictive behaviors (1). The global prevalence of ASD is estimated to be about 1 in 100 children (2). Children with ASD also frequently present with other comorbid conditions (3) and associated behavioral problems (4) that could pose a significant burden to their family (5) and also society (6). Despite the rise in public awareness of ASD and the increase in prevalence of ASD diagnosis (2), there are still no FDA-approved medications for treating the core symptoms of ASD (7) and early behavioral intervention remains the only currently acceptable strategy for achieving favorable outcome for children diagnosed with ASD (8).

Although available evidence endorses the importance of early treatment for children suffering from ASD (9), early signs of ASD can be difficult to observe especially for those with high-functioning [i.e., normal intelligence quotient (IQ)] autism (10). Therefore, early identification of those at risk of ASD is critical for the prompt implementation of appropriate interventions (11). Despite the complex and possibly multifactorial etiologies of ASD, most previous research suggested that genes, environment, and gene-environment interactions are the key factors in the pathogenesis of ASD (12). Therefore, the association between several parental illnesses and risks of ASD in their offspring has been an important area in ASD research not only to improve the understanding of the genetic influence on the development of ASD but also to help identify children who may be at an increased risk of the disease (13, 14).

A variety of autoimmune or autoinflammatory diseases in parents, such as inflammatory bowel diseases and inflammatory arthritis, were found to be associated with higher risks of ASD in offspring (13). In particular, rheumatoid arthritis (RA), which is one of the most extensively studied parental autoimmune diseases for its possible link to offspring ASD, has been found to be associated with a higher risk of offspring ASD in many observational studies (15–17). Indeed, a previous meta-analysis also demonstrated an elevated risk of offspring ASD in mothers diagnosed with RA, suggesting a possible link between parental autoimmune diseases and offspring ASD (18). However, that meta-analysis was only able to include five observational studies and did not investigate the correlation between offspring ASD and paternal RA (18). Moreover, some studies only selected mothers who were diagnosed with RA prior to childbirth (19, 20), while others included mothers with their diagnosis of RA being made in any time frame regardless of its relation to childbirth (21, 22). Although maternal RA with an onset prior to childbirth may predispose offspring to the risk of brain maldevelopment through exposing the fetus to an unfavorable pregnancy environment (23–25), the impact of the timing of maternal RA onset with reference to childbirth was not addressed in that meta-analysis (18).

Therefore, to fill this gap in knowledge, the current updated meta-analysis aimed at investigating the association of the risk of offspring ASD with both maternal RA and paternal RA as well as assessing the difference in risk of offspring ASD between mothers who were diagnosis with RA prior to childbirth and those with the diagnosis of RA being made in any time frame.

Methods

This meta-analysis was reported based on recommendations from the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA). The protocol was previously registered in PROSPERO (CRD42022358470).

Search strategy

Three independent databases including Embase, Medline, and Cochrane Library were searched for the identification of studies that investigated the association of parental RA with the risk of offspring ASD. We also searched Google Scholar and the reference lists of the included studies to retrieve other relevant articles. There were no language and ethnicity restrictions applied for determining study eligibility. The database research was conducted from inception until
TABLE 1 Summary of characteristics of included studies in the current meta-analysis (n = 10).

| References       | Design    | RA                                | ASD cases/Total participants | Estimated risks (95% CI) | Parents age (years) | Children age (years) | Female (%) in ASD in offspring | NOS scores | Country       |
|------------------|-----------|-----------------------------------|------------------------------|--------------------------|---------------------|----------------------|-------------------------------|------------|---------------|
| Atladóttir et al. (16) | Cohort study | Mother (b/a childbirth) | 3,325/689,196 | 1.70 (1.07, 2.54) | N/A | N/A | 16.15 | 7 | Denmark |
| Chiu et al. (15) | Cohort study | Father | 297/263,791 | 1.42 (0.7, 2.87) | 1.49 (1.01, 2.2) | 61.66 | 33.95 | 19.26 | 8 | Taiwan |
| Croen et al. (30) | Case control | Father (2 years before and 2 years after childbirth) | 407/2,502 | 0.858 (0.103, 7.142) | 30.01 | N/A | 18 | 8 | U.S. |
| Croen et al. (19) | Case control | Mother (before childbirth) | 663/1,587 | 2.09 (0.74, 5.89) | N/A | N/A | 17.6 | 8 | U.S. |
| Lee et al. (20) | Cohort study | Father (b/a childbirth) | 2/4,506 | N/A | 1.38 (0.35, 5.53) | 61.66 | 33.95 | 19.26 | 8 | Taiwan |
| Li et al. (31) | Cohort study | Father (b/a childbirth) | 13,885/1,386,260 | 1.03 (0.82, 1.31) | 1.13 (0.94, 1.37) | N/A | 9.7 | 52.2 | 8 | Taiwan |
| Mouridsen et al. (22) | Case control | Father (b/a childbirth) | 111/441 | 0.627 (0.03, 13.15) | 0.49 (0.06, 4.12) | 65.58 | 5.4 | 26.1 | 7 | Denmark |
| Rom et al. (17) | Cohort study | Father (b/a childbirth) | 8985/1,917,723 | 1.33 (0.97–1.82) | 1.31 (1.06, 1.63) | 1.39 (1.11, 1.75) | N/A | 9.7 | 52.2 | 8 | Denmark |
| Spann et al. (21) | Case control | Father (b/a childbirth) | 4,600/18,400 | 0.9 (0.6, 1.4) | 1.1 (0.8, 1.5) | N/A | 8 | 13.92 | 7 | Finland |
| Tsai et al. (32) | Cohort study | Mother (b/a childbirth) | 10,631/1,893,244 | 1.42 (0.60, 3.40) | 31.97 | 6.24 | 47.8 | 8 | Taiwan |

ASD, autism spectrum disorder; b/a, before and after (indicating diagnosis of RA regardless of time of childbirth); CI, confidence interval; N/A, not available; NOS, the Newcastle-Ottawa Scale; RA, rheumatoid arthritis.

September 7, 2022. The major terms used for literature search included "Rheumatoid Arthritis" AND "Autism" for literature search. Medical Subject Headings (MeSH), and Embase Subject Headings (Emtree) were also used to facilitate searching. The detailed search strategies for these databases are shown in Supplementary Table 1.

Studies selection and data extraction

The titles and abstracts of the retrieved articles were independently screened by two authors for eligibility based on the following criteria: (1) observational studies such as cohort or population-based investigations that reported an association between parental RA and risk of offspring ASD; (2) availability of information about odds ratio (OR), relative risk (RR), hazard ratio (HR), or sufficient details (e.g., number of cases) for risk assessment. We excluded the following forms of publications: letters, conference abstracts, meta-analyses, case reports, reviews, and animal experimental studies. For two or more institution-based cohort studies that reported findings on the same or overlapping population, the one with the largest sample size was selected. A third reviewer was consulted for any disagreement.

The following details were extracted from all eligible studies: publication-specific information (e.g., first author’s name, journal, and year of publication) and study characteristics including population size, country, follow-up duration, number of cases, and details for risk calculation [e.g., ORs with corresponding 95% confidence interval (CI)]. If a study reported both unadjusted and adjusted data, we collected the latter for analysis.
Primary and secondary outcomes

The primary outcomes included the association of maternal RA with the risk of offspring ASD and that of paternal RA with offspring ASD risk regardless of parental age and the timing of RA diagnosis. Two subgroup analyses were performed. The first subgroup investigation was conducted to evaluate the potential impact of pre-childbirth maternal RA (e.g., teratogenicity, pregnancy-related conditions) on the risk of offspring ASD after separating the maternal participants into those diagnosed with RA before childbirth and those whose diagnosis of RA was made without referring to the timing of childbirth, followed by comparing their differences in the risk of offspring ASD relative to that in their respective controls (i.e., mothers without diagnosis of RA). The second subgroup analysis focused on the effect of geographical location (i.e., Western vs. Asian country) on the risk of offspring ASD. The diagnosis of RA and ASD was in accordance with that of each study.

Quality assessment

Two reviewers independently judged the risk of bias in the retrieved studies based on the Newcastle-Ottawa Scale (NOS), which categorizes the possible sources of bias in a cohort study into eight items pertinent to selection, comparison, and outcome. Disagreements were solved by a third reviewer. A nine-point scale was used to grade the included studies (i.e., low, moderate, and high quality for a score of 0–3, 4–6, and 7–9, respectively).

Statistical analysis

With the comprehensive Meta-Analysis (CMA) V3 software (Biostat, Englewood, NJ, USA), pooled effect size was estimated using reported patient number or raw data of OR. HR and RR were directly considered to be OR as previously reported (18), taking into account the rarity of ASD. Because this
FIGURE 2
Forest plot showing association of maternal rheumatoid arthritis (RA) with risk of offspring autism spectrum disorder (ASD) regardless of the timing of maternal RA diagnosis. CI, confidence interval.

| Study name          | Odds ratio | Lower limit | Upper limit | p-Value | Odds ratio and 95%CI | Relative weight |
|---------------------|------------|-------------|-------------|---------|----------------------|------------------|
| Atladottir 2009     | 1.700      | 1.070       | 2.700       | 0.025   |                      | 6.28             |
| Chiu 2022           | 1.490      | 1.010       | 2.199       | 0.045   |                      | 8.87             |
| Croen 2019          | 2.090      | 0.741       | 5.886       | 0.164   |                      | 1.25             |
| Lee 2021            | 1.380      | 0.347       | 5.485       | 0.647   |                      | 0.71             |
| Li 2022             | 1.130      | 0.936       | 1.364       | 0.203   |                      | 37.90            |
| Rom 2018            | 1.310      | 1.056       | 1.624       | 0.014   |                      | 29.05            |
| Sann 2019           | 1.100      | 0.803       | 1.506       | 0.552   |                      | 13.61            |
| Croen 2005          | 0.858      | 0.103       | 7.142       | 0.897   |                      | 0.30             |
| Mouridsen 2007      | 0.491      | 0.058       | 4.123       | 0.512   |                      | 0.30             |
| Tsai 2018           | 1.326      | 0.550       | 3.196       | 0.530   |                      | 1.74             |
|                     | 1.246      | 1.110       | 1.400       | 0.000   |                      |                  |

Results

Identification of eligible studies

Our database research retrieved 1,666 records, of which 217 were excluded for being duplicates. Following title and abstract screening, 1,429 records were further removed. Of the remaining 20 articles eligible for full-text reading, 10 were excluded because of being review articles (n = 6), conference abstract (n = 1), and failure to meet the inclusion criteria (n = 3). Finally, 10 studies were eligible for inclusion in this systematic review (15–17, 19–22, 30–32). The flowchart of study selection is demonstrated in Figure 1.

Study characteristics and risk of bias

Ten studies published between 2005 and 2022 involving 6177650 participants were analyzed (15–17, 19–22, 29, 30, 32). In terms of research designs, six (15–17, 20, 29, 32) and four (19, 21, 22, 30) were cohort and case-control studies, respectively. The method for identification of ASD and RA is shown in Supplementary Table 2. The sample sizes of the studies ranged from 441 to 1,893,244. Of the 10 studies, six documented a relationship between the risk of offspring ASD and mothers diagnosed with RA regardless of timing of maternal RA diagnosis (15–17, 21, 22, 29), four offered data on the risk of offspring ASD associated with the diagnosis of maternal RA prior to childbirth (15, 19, 20, 32), and four provided information about the risk of offspring ASD linked to paternal RA (15, 17, 21, 29). Five studies provided details regarding parental age at the time of offspring ASD diagnosis, which was between 29.05 and 66.6 years. In respect of ethnic backgrounds, six studies were conducted in Western countries (16, 17, 19, 21, 22, 30) and four in Asian countries (15, 20, 29, 32). The mean NOS score of the included studies was 7.7 (SD: 0.48). Detailed information for each included study is provided in Table 1.

Outcomes

Primary outcome: Association between maternal/paternal rheumatoid arthritis and risk of offspring autism spectrum disorder

Pooled results revealed a significant association between maternal RA and the risk of ASD (OR = 1.246, 95% CI: 1.11 to 1.4, p < 0.001; I² = 0%, ten studies) (Figure 2; 15–17, 19–22, 29, 30, 32). Sensitivity analysis demonstrated no significant influence on the results by omitting certain studies. Both funnel plot and Egger’s test indicated a low risk of publication bias (Egger’s test: p = 0.595) (Figure 3).

Our analysis demonstrated no impact of paternal RA on the risk of offspring ASD (OR = 1.104, 95% CI: 0.932 to 1.309,
$p = 0.253, I^2 = 2.127\%$) (Figure 4; 15, 17, 21, 29). The pooled result was not significantly influenced by removing certain studies on sensitivity analysis.

**Subgroup analysis: The impact of timing of maternal rheumatoid arthritis diagnosis on risk of offspring autism spectrum disorder**

The impact of timing of maternal RA diagnosis on the risk of offspring ASD is shown in Figures 5, 6. Analysis of the four studies focusing on mothers with pre-childbirth diagnosis of RA (15, 19, 20, 32) demonstrated no significant association between maternal RA and offspring ASD (OR = 1.449, 95% CI: 0.83 to 2.53, $p = 0.192, I^2 = 0\%$, four studies) with consistent findings on sensitivity analysis (Figure 5).

On the other hand, analysis of maternal RA regardless of the timing of diagnosis revealed a significant positive association of maternal RA with the risk of offspring ASD (OR = 1.227, 95% CI: 1.087 to 1.385, $p = 0.001, I^2 = 36.91\%$, six studies) (15–17, 21, 22, 29). The pooled result was not significantly influenced by excluding certain trials on sensitivity analysis (Figure 6).

**Subgroup analysis: Impact of geographical location on the association between maternal rheumatoid arthritis and risk of offspring autism spectrum disorder**

Findings of subgroup analysis based on geographical location (i.e., Western vs. Asian countries) are shown in Figures 7, 8. Pooled results from studies conducted in Western countries (i.e., European/United States) revealed a significant correlation between maternal RA and offspring RA (OR = 1.295, 95% CI: 1.101 to 1.524, $p = 0.002, I^2 = 0\%$, six studies) (16, 17, 19, 21, 22, 30; Figure 7). Sensitivity analysis demonstrated a loss of significance in this association when one study was removed (17) (OR = 1.283, 95% CI: 0.99 to 1.663, $p = 0.06$).

Focusing on Asian countries, the pooled results also disclosed a significant link between maternal RA and the risk of offspring ASD (OR = 1.198, 95% CI: 1.015 to 1.413, $p = 0.032, I^2 = 0\%$, four studies) (15, 20, 29, 32; Figure 8). Nevertheless, sensitivity analysis showed inconsistent results when one study (15) was excluded (OR = 1.142, 95% CI: 0.951 to 1.37, $p = 0.155$).

**Discussion**

Although a previous meta-analysis demonstrated an elevated risk of offspring ASD in mothers suffering from autoimmune diseases, that study only included five studies for RA and investigated only the maternal side (18). Our meta-analysis was updated with 10 studies and was the first to investigate the risk of offspring ASD on the paternal side. To assess the possible influence of pregnancy, we further conducted subgroup analysis on the risk of offspring ASD in mothers diagnosed with RA before childbirth. Albeit statistically non-significant, our subgroup analysis showed a 1.44-fold increase in the risk of offspring ASD in mothers diagnosed with RA before childbirth compared to those without the diagnosis before childbirth, suggesting a potential impact of RA-related maternal
changes before childbirth on the risk of ASD in offspring. Whether the absence of statistical significance was attributed to the availability of only four studies for analysis remains unclear. In contrast, comparison of the risk of offspring ASD between a mixed population of mothers diagnosed with RA before and after childbirth and those without RA demonstrated a relatively small 1.25-fold but significant increase in the risk of offspring ASD. On the other hand, the results of our study
failed to demonstrate an elevated risk of offspring ASD in fathers diagnosed with RA. Overall, our results not only showed an elevated risk of ASD in offspring associated with maternal RA rather than paternal RA but also suggested an impact of RA-related changes in maternal physical condition (i.e., systemic inflammation or medications) before childbirth on the risk of offspring ASD that warrants further investigations.

Consistent with the results of a previous meta-analysis that demonstrated a 39% increase in risk of ASD in offspring of mothers diagnosed with ASD compared with those without RA (18), our study showed a 25% elevated risk. Compared with that study (18), the inclusion of five more studies (15, 20, 21, 29, 30) in the present meta-analysis provided more robust evidence to support such an elevated risk. Nevertheless, most of the new studies in the current meta-analysis were conducted in the same geographic locations as the previous meta-analysis (18), including Taiwan, U.S., and Denmark with only one new country (Finland) being included in our updated study. Therefore, the similarities in geographic locations and ethnicities of the study populations may partly explain the consistent findings between the previous study (OR: 1.39) (18) and the present meta-analysis (OR: 1.25). Focusing on ethnicity, our subgroup analysis demonstrated comparable increases in risk between Western and Asian countries, suggesting that an elevated risk of ASD in offspring of mothers with RA may be a universal phenomenon. Further studies are required to elucidate the validity of this finding.

There are several possible mechanisms that may explain this elevated risk of offspring ASD in mothers with RA. First, genetic inheritance, which is considered to be one of the most important causes of ASD, is supported by several studies that found a shared genetic liability between autoimmune diseases and ASD, such as human leukocyte antigen (HLA) alleles (33–35). Interestingly, this elevated risk in offspring ASD was less obvious or non-significant on paternal side as reported in previous studies (15, 17, 21, 29) and also in our meta-analysis. Therefore, if shared common genetic paths are the main reason for the elevated risk, such a genetic influence seemed only to inherit through the maternal side. However, a previous study investigating a shared risk allele between ASD and RA (i.e.,
HLA-DR4) during pregnancy ruled out genomic imprinting and mitochondrial allele as the causes of an increased ASD risk in offspring from the mothers (33). Therefore, this elevated risk of offspring ASD on the maternal side may be better explained by changes in maternal condition before childbirth, in which teratogenic effects from shared risk alleles (33) and possible adverse effects from autoimmune mediators (36, 37) or medications (38, 39) during pregnancy may have a role to play in fetal development.

Taking into consideration the importance of environmental factors, especially pregnancy, in the etiology of ASD (36, 37), we further conducted subgroup analysis of studies that focused on mothers diagnosed with RA before childbirth to investigate the effects of pregnancy and other factors before childbirth on the risk of offspring ASD. Although our results showed a 1.44-fold increase in the risk of ASD in offspring among mothers suffering from RA before childbirth compared to those without RA, such an increased risk failed to reach statistical significance \( (p = 0.116) \) possibly due to the limited numbers of studies (i.e., only four). In contrast, despite the apparently smaller increase in risk from a mixed population comprising women diagnosed with RA before and after childbirth (OR: 1.25), the increase was probably attributable to a larger number of studies (i.e., ten) with a narrower confidence interval.

Because the peak incidence of RA for adults is in their sixties (40) which is long after the fertile period, the genetic influence of maternal RA may be largely diluted if we only included mothers diagnosed with RA prior to childbirth, as a large portion of mothers who carried RA or ASD-related genes were still asymptomatic and mis-classified into the non-RA group for comparison. Indeed, two of our included studies recruiting mother diagnosed with RA regardless of the timing of diagnosis (i.e., before vs. after childbirth) have demonstrated a mean maternal age of around sixties (15, 22), while the mean age of mothers was around thirty in studies that only included those diagnosed with RA before childbirth (20, 30, 32). Therefore, we would expect an underestimation of the risk of offspring ASD in studies that only included mothers with RA before childbirth to assess maternal genetic influence on the risk of ASD in offspring. Nevertheless, a higher risk of offspring RA was observed in our subgroup of mothers suffering from RA prior to childbirth compared to that of a mixed population of mothers with RA diagnosis both before and after childbirth, implicating that pre-childbirth conditions (e.g., pregnancy) may have an important contribution to the risk of ASD in their offspring. Such a proposal was further supported by a study suggesting that risk alleles such as HLA-DR4 may have a teratogenic effect (33). Consistently, the results of other investigations have highlighted that certain conditions during pregnancy such as inflammation or infection (36, 37) may contribute to an increased risk of offspring ASD. For instance, previous animal studies have demonstrated that the elevated levels of cytokines in fetal circulation from in utero exposure in mothers with autoimmune diseases were associated with abnormalities in fetal neurodevelopment (36, 37). A clinical correlation between immune mediators and fetal neurodevelopment was further reflected by the finding that the increased level of serum immunoglobulin G (IgG) in women diagnosed with autoimmune diseases (41) could access the fetal compartment during gestation (42) with a higher concentration being found in the circulation of children with autism than that in those with normal development (43). Finally, medications used to relieve pain or control inflammation during pregnancy have also been shown to be associated with an elevated risk of offspring ASD (38, 39), despite the report of a protective effect of NSAID during pregnancy against offspring ASD in one study (44). Therefore, our findings may still implicate the potential impacts of factors before childbirth (e.g., teratogenicity, increased susceptibility during pregnancy) on the risk of offspring ASD.

There are several limitations in the present study. First, although we doubled the number of included studies compared to the previous meta-analysis (18), the additional studies were still conducted in three main geographic locations including Taiwan, Europe, and U.S.; therefore our results may not be extrapolated to populations with other geographic or ethnic backgrounds. Indeed, genetics may be one of most important factors in the etiologies of ASD; a previous study found that the association between certain risk alleles in autoimmune disease and ASD was only observed in Western countries but not in the Chinese population (45), implying a variation in influences of risk alleles among people of different ethnicities. Second, the availability of only four studies for investigating the risk of ASD in offspring of mothers diagnosed with RA before childbirth and the non-significant increase in risk of offspring ASD in this subgroup warrant further studies to verify our findings. Second, because the non-significant increase in the risk of offspring ASD of mothers diagnosed with RA before childbirth was derived from only four studies, further studies are warranted to verify our findings. Third, despite the known associations of other confounders such as an increased body weight of mother (46), medical conditions such as diabetes (47), and certain habits such as smoking (48) with elevated risks of both ASD (49) and offspring RA (48), unavailability of relevant information in most of the included studies precluded the conduction of meaningful subgroup analyses. Fourth, because most of the included studies based their diagnoses of ASD and RA on medical registry or medical records instead of independent validation, a lack of adequate case definition or ascertainment of exposure were the main sources of bias. Fifth, despite our adoption of OR for risk assessment, three studies only provided data on HR (17, 20, 29). Therefore, bias may arise from our using HR to replace OR in our analysis. Finally, since our finding of an absence of
association between the risk of offspring ASD and paternal RA were derived from only four studies, this preliminary finding remains to be verified.

**Conclusion**

Our finding supported an association between maternal RA and an elevated risk of ASD in offspring. While shared genetic liability may partly explain this correlation, conditions before childbirth may also have an important role to play. However, given the lack of solid case definition and the limited numbers of studies investigating the risk of offspring ASD in mothers diagnosed with RA before childbirth, further studies are warranted to elucidate the maternal effects of RA before childbirth (e.g., pregnancy) on the risk of offspring ASD.

**Data availability statement**

The original contributions presented in the study are included in the article/Supplementary material, further inquiries can be directed to the corresponding author.

**Author contributions**

C-KS and Y-SC: conceptualization and literature search. I-WC and H-JC: methodology. WC and R-FT: trial selection. H-YF: data analysis. H-YF and C-WL: data extraction. C-KS, Y-SC, and K-CH: writing—original draft preparation. K-CH and C-KS: writing—review and editing. All authors had read and agreed to the published version of the manuscript.

**Conflict of interest**

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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**Supplementary material**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmed.2022.1052806/full#supplementary-material

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