

# Executive function in children with neurodevelopmental conditions: a systematic review and meta-analysis

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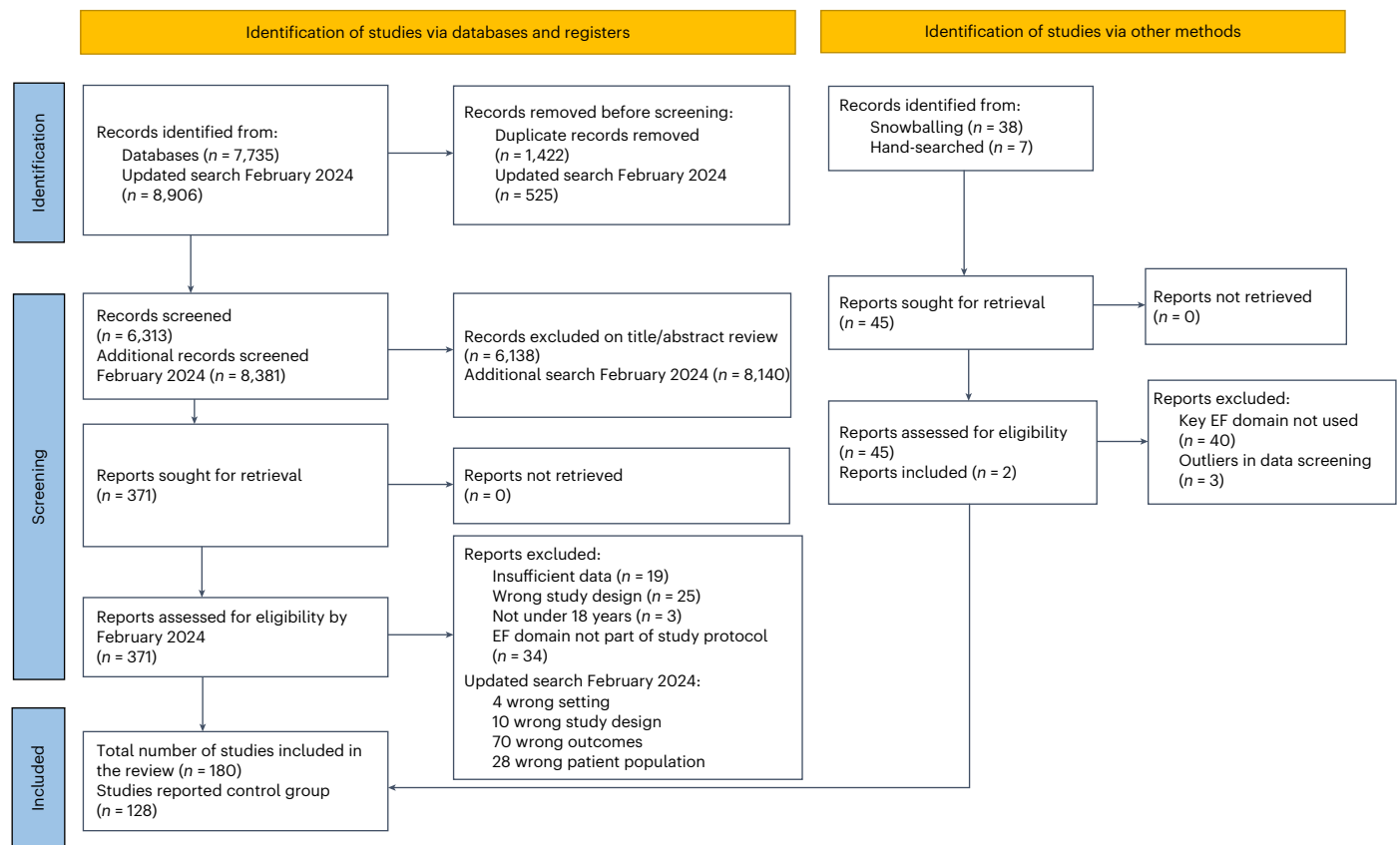
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Executive function (EF) delays are well documented in paediatric neurodevelopmental conditions (NDCs). There is no consensus about whether EF delay represents a transdiagnostic feature of NDCs. This systematic review and meta-analysis synthesized 180 studies reporting two or more NDC comparisons on EF, examined differences between NDCs, and the moderating effects of gender, age, publication year, DSM editions and assessment types. Studies using established EF measures across seven domains (attention, fluency, set-shifting, set-switching, response inhibition, planning and working memory) in participants under 18 were included. Summary effects were compared: (1) for all reported NDCs relative to control, (2) for each individual NDC relative to control and (3) between NDC groups. Results confirmed that EF delay was a transdiagnostic feature of neurodevelopmental delay, with a moderate effect size of delay across all NDCs ( $g = 0.56$ , 95% confidence interval (CI) 0.49–0.63) compared with control. This effect increased with comorbidities ( $g = 0.72$ , 95% CI 0.59–0.86), DSM-5 criteria and informant measures. Comparisons between NDCs revealed few differences: children with tic disorders (TD) showed smaller EF delays, children with attention-deficit/hyperactivity disorder (ADHD) showed larger delays in attention, response inhibition, planning and working memory compared with TD and specific learning disorders, while children with autism spectrum disorders showed greater delays on set-switching compared with ADHD. Findings support transdiagnostic models of neurodevelopment to further a developmentally sensitive science that can reveal how EF delays contribute to brain circuitry, symptom profiles and functioning, and ultimately support early interventions and outcomes for all children with NDCs.

Executive function (EF) is an umbrella term used to conceptualize a range of cognitive processes including planning, working memory, attention, inhibition, self-monitoring, self-regulation and initiation<sup>1,2</sup>. It encompasses a network of functional cognitive abilities that allow for real-world engagement, which are often refined in developmental periods<sup>2,3</sup>. Impairments in EF can include difficulties in sustaining attention, impulsivity and an inability to transition and flexibly manage multiple tasks<sup>1,4–8</sup>. Their presence may also signal divergence in brain

development and can play a key role in both causal and maintaining factors in neurodevelopmental conditions (NDCs)<sup>9,10</sup>. Together, these processes are involved in a range of emotional, behavioural and social functions. Theoretical models over past decades propose that EF is comprised of core components that emerge early in development and are built upon over time, resulting in complex, higher-order cognitive abilities that are critical for daily functioning<sup>11,12</sup>. These models have been informed and supported by a substantial amount of empirical

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**Fig. 1 | The PRISMA diagram of this systematic review.** The retrieval processes are illustrated, covering the stages of identification, screening and inclusion of studies. Note that the use of *n* indicates the number of studies included at each stage.

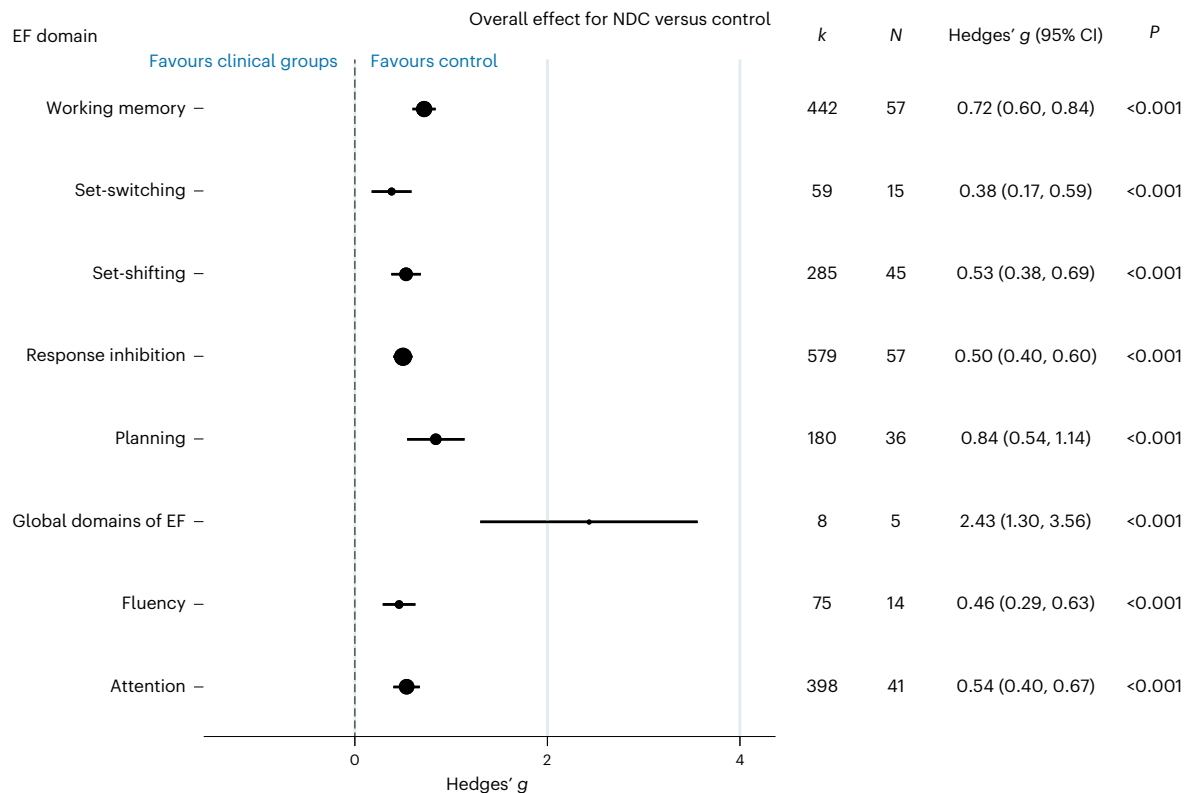
data<sup>13,14</sup> and set the groundwork for this review. This review aims to investigate both higher-order and more foundational domains within the paediatric population. EF studies presents considerable debate on how EF domains overlap or present as separate faculties. Further, the literature attempts to capture these faculties through the use of various EF outcomes. Considering the body of literature in this area, this review will focus on attention, set-shifting, set-switching, fluency, planning, working memory and response inhibition. This also enables consideration of broader EF domains, such as attention<sup>2</sup>, where impairments have been consistently reported in NDCs such as attention-deficit/hyperactivity disorder (ADHD)<sup>15</sup>. Impairments in EF are believed to impact quality of life in children<sup>16</sup> and can contribute to lifelong challenges<sup>1,3</sup>. These impairments are thought to be moderated by variables such as sex, type of EF measure and age<sup>17</sup>. For example, some studies have suggested that EF difficulties may increase during adolescence<sup>18</sup>, while others do not<sup>19–21</sup>. Overall, delays in EF could be considered to represent a broad transdiagnostic cognitive phenotype of NDCs. There has, however, been limited research examining this across NDCs.

EF impairments have traditionally been studied within specific NDCs<sup>22,23</sup>. Paediatric EF studies have generally focused on emerging core EF domains of inhibition, working memory and cognitive flexibility<sup>11,24</sup>. These domains are believed to serve foundational cognitive faculties that allow for the development of higher-order EF functions, such as planning and problem-solving<sup>11</sup>. Delays in each of these domains have been reported in children with NDCs in the first years of life<sup>1,3</sup>. So far, age-based effects have also been examined within disorders. For example, deficits in working memory are well documented in children with ADHD and specific learning disorders (SLD) across different developmental stages<sup>25–27</sup>.

Cross-disorder studies<sup>23,28</sup> further show that there may be some differences in sub-domain EF outcomes based on the specific disorder phenotype. In support, some studies comparing children diagnosed with autism spectrum disorder (ASD) or ADHD suggest that children with ASD show marked delays in cognitive flexibility and planning, while children with ADHD have delays in inhibition and working memory<sup>29–32</sup>. Other studies<sup>33</sup> have reported larger delays in response inhibition, cognitive flexibility/switching and working memory for children with ASD compared with those with ADHD<sup>33</sup>. Such comparisons are important to inform the understanding of causal and maintaining features that might differentiate clinical profiles of conditions. Further, they could be used to screen for diagnoses, to confirm co-occurring NDCs and to assess for clinical severity.

## EF measures in the paediatric population

Assessments of EF include both performance and informant-based measures. Performance EF tasks typically involve practical tasks of EF capacities (for example, errors on a computer-based response inhibition task; placing objects in an instructed order) and are purported to objectively tap into discrete EF domains. There is, however, a high degree of overlap between EF domains and their underlying neurobiology, raising questions about their functional independence<sup>8</sup>. As such, many measures of EF likely tap multiple domains. For example, tasks that measure set-shifting, such as the Wisconsin Card Sorting Test, may be impacted by a working memory component<sup>34</sup>. Other measures, such as informant-based measures, are proposed to possess greater ecological validity given their reliance on the reporting of observed everyday behaviours<sup>1</sup>. Informant-based measures typically describe how a child's EF profile directly relates to their daily functioning, encompassing a greater number of observed



**Fig. 2 | Forest plot summaries overall effects (Hedges'  $g$ ) for EF in NDCs versus controls.** Here  $N$  refers to the number of studies across EF comparisons made within the meta-analysis, and  $k$  refers to the number of outcome measures reported. Mean effects and their 95% CI are presented by the central black circle

and the horizontal line. The size of the black circle reflects the number of studies included in each comparison. The dashed line at  $g = 0$  indicates no effect, while the  $P$  values show whether the effect was statistically significant. The  $P$  values reported from the random-effects models are two-sided.

functional deficits<sup>17,22</sup>. Such measures might have more ecological validity, but come at a potential cost of being less objective<sup>25–27</sup>. The debate between the ecological validity and objective reliability of these measures has been long-standing. Some authors have argued that performance-based measures may provide more objective scaling of EF performance, while others have pointed to the utility of informant and self-report measures to predict functioning outcomes; nonetheless, each provides unique clinical utility for a range of paediatric contexts<sup>35–37</sup>.

## Study aims

There is a lack of research examining EF as a transdiagnostic phenotype in paediatric NDCs. This meta-analysis aims to synthesize the existing literature of EF in NDCs. It reviews seven EF constructs identified in paediatric literature (attention, fluency, set-shifting, set-switching, response inhibition, planning and working memory). Given that EF is a component of cognition across neurodevelopment, it is critical to evaluate its discriminating and/or shared profile across NDCs. This meta-analysis aimed to review studies that tested EF measures across two or more NDCs in children with or without a control comparison. Further, this study aimed to consider how the type of assessment measure (informant or performance-based), DSM criteria and demographic factors (age and gender) influenced the results of the primary analysis. We predicted that:

1. NDCs would be associated with significant delays across EF domains, when compared with controls.
2. Effect size of delay would be moderated by the type of assessment and reported sex, such that informant-based measures and a higher percentage of males captured in studies would be associated with larger effect sizes.

3. Comparisons of EF profiles between NDCs would show differences in the severity of delay overall and for EF sub-domains.

## Results

### Primary outcome: neurodevelopmental groups and controls

An overall meta-analysis comparing eligible studies that compared two or more NDCs with controls was completed (see Fig. 1 for the literature search process and Supplementary Table 3 for study characteristics). To facilitate the analysis and accurate interpretation of our findings, we systematically reported Hedges'  $g$  effect size estimates ( $g$ ), with 95% confidence interval (CI) when appropriate, and reported variances (as detailed in Methods). The neurodevelopmental groups showed significant impairment in their overall EF ( $N = 114$  studies,  $k = 2031$  outcomes,  $g = 0.56$ , 95% CI = 0.49 to 0.63,  $P < 0.001$ ,  $\tau^2 = 0.14$ , prediction interval  $-0.17$  to 1.29). These effect sizes overall were statistically significant for both performance ( $N = 110$  studies,  $k = 1911$  outcomes,  $g = 0.51$ ,  $P < 0.001$ ) and informant-based measures ( $N = 14$  studies,  $k = 120$  outcomes,  $g = 1.49$ ,  $P < 0.001$ ) and across all domains of EF (Fig. 2).

Funnel plot asymmetry was detected with Egger's test, indicating a possible small-study effect ( $\beta = -0.31$ ,  $P < 0.001$ ). A trim-and-fill analysis did not find any missing studies (Supplementary Fig. 1). Inspection of the funnel plot revealed that asymmetry was driven by the informant measurement outcomes (with  $g > 2$ ). Five studies were removed as they contributed to plot asymmetry (Supplementary Table 4). The overall effect stayed the same ( $g = 0.56$ , 95% CI = 0.49 to 0.63,  $P < 0.001$ ,  $\tau^2 = 0.14$ , prediction interval  $-0.17$  to 1.29).

A meta-analysis was conducted comparing paediatric NDCs and their control group (Fig. 3). Significant delays in overall EF were observed for all neurodevelopmental groups, except FASD, and across all sub-domains (Fig. 2). Effect sizes were moderate to large, except for TD where a small effect size was observed. For FASD, the sample

collected was imprecise with a small number of studies ( $N = 4$  studies) to contribute to this observation. For those conditions with the greatest number of studies, effect sizes were moderate in size (ADHD, ASD and SLD), suggesting similar delays. Analysis was then conducted between those studies that specifically recruited comorbid NDC groups of children (for example, children diagnosed with both ASD and SLD) against those studies that only required a single NDC diagnosis for inclusion. Studies focused on recruiting comorbid NDCs showed a larger effect size of EF delay ( $N = 58$  studies,  $k = 427$  outcomes,  $g = 0.72$ , 95% CI = 0.59 to 0.86,  $P < 0.001$ ) (as shown in Supplementary Table 5) against studies that only required diagnosis of a single NDC ( $N = 113$  studies,  $k = 1601$  outcomes,  $g = 0.53$ , 95% CI = 0.46 to 0.60,  $P < 0.001$ ).

### Moderator analysis

There was a small, though significant, moderator effect of reported sex, based on the percentage of males ( $N = 92$  studies,  $k = 1746$  outcomes,  $\beta = -0.003$ , 95% CI =  $-0.005$  to  $-0.001$ ,  $P = 0.016$ ,  $Q_{(1)} = 5.84$ ,  $R^2 = 1.33\%$ ). The higher the percentage of males, the smaller was the effect size of NDC compared with control, but this accounted for a negligible degree of variance. The moderator effect of age was not significant ( $N = 111$  studies,  $k = 1995$  outcomes,  $g = -0.027$ , 95% CI =  $-0.076$  to  $0.021$ ,  $P = 0.267$ ,  $Q_{(1)} = 1.23$ ,  $R^2 = 0.82\%$ ).

The effect of DSM edition used in assessments was significant ( $N = 84$  studies,  $k = 1,452$  outcomes,  $P < 0.001$ ,  $Q_{(2)} = 75.25$ ,  $R^2 = 22.92$ ). DSM-III ( $g = 0.49$ , 95% CI = 0.31 to 0.66,  $P < 0.001$ ) had a similar effect size to DSM-IV ( $g = 0.47$ , 95% CI = 0.39 to 0.55,  $P < 0.001$ ). However, effect sizes increased significantly with the use of the DSM-5 edition ( $g = 0.92$ , 95% CI = 0.81 to 1.03,  $P < 0.001$ ). This effect held when examining only performance-based measures ( $N = 80$  studies,  $k = 1363$  outcomes,  $P < 0.001$ ,  $Q_{(2)} = 19.45$ ,  $R^2 = 25.38\%$ ), with the use of DSM-III ( $g = 0.43$ , 95% CI = 0.27 to 0.59,  $P < 0.001$ ), DSM-IV ( $g = 0.46$ , 95% CI = 0.39 to 0.53,  $P < 0.001$ ) and DSM-5 ( $g = 0.73$ , 95% CI = 0.61 to 0.85,  $P < 0.001$ ), when each single disorder (TD, ASD, ADHD, SLD and developmental coordination disorder (DCD) was separately removed from the analysis, and when comorbid NDCs were removed from analysis. Informant-based measures did not show significant differences ( $N = 11$  studies,  $k = 89$  outcomes,  $P = 0.592$ ,  $Q_{(1)} = 0.29$ ,  $R^2 = 0\%$ ). Publication year was not significantly different ( $N = 111$  studies,  $k = 1964$  outcomes,  $g = 0.001$ , 95% CI =  $-0.006$  to  $0.007$ ,  $P = 0.868$ ,  $Q_{(1)} = 0.028$ ,  $R^2 = 0.17\%$ ).

Moderator analyses also revealed that there were significant moderator effects based on the type of EF assessment measure ( $N = 113$  studies,  $k = 2028$  outcomes,  $Q_{(1)} = 717.47$ ,  $P < 0.001$ ,  $R^2 = 19.08\%$ ). Effect sizes for performance measures ( $g = 0.49$ , 95% CI = 0.43 to 0.56,  $P < 0.001$ ) were significantly smaller compared with informant measures ( $g = 1.47$ , 95% CI = 1.38 to 1.56,  $P < 0.001$ ).

### EF profiles in specific NDCs

The next set of analyses examined whether the observed delay found on EF measures differed according to the neurodevelopmental diagnosis (see Supplementary Table 6 for details). To address this question, we conducted a series of analyses between each NDC using studies that provided enough power ( $N \geq 3$  studies). This was informed by the presence of underlying heterogeneity as well as a sufficient number and balance of studies within subgroups to allow for cross-condition analyses to take place within each cell (Fig. 4 and Supplementary Table 8). In the majority of cases, there were not sufficient data to explore both performance and informant outcomes separately (results from these analyses where available are provided in Supplementary Fig. 2).

**ADHD and ASD.** Studies comparing ADHD and ASD showed that there were no significant differences on overall EF measures between conditions ( $N = 39$  studies,  $k = 284$  outcomes,  $g = 0.076$ , 95% CI =  $-0.026$  to  $0.18$ ,  $P = 0.146$ ,  $t^2 = 0.088$ , prediction interval  $-0.52$  to  $0.67$ ). The sub-domain comparisons, however, revealed that children with ADHD performed better on measures of set-shifting compared with children

with ASD ( $N = 16$  studies,  $k = 42$  outcomes,  $g = -0.19$ , 95% CI =  $-0.35$  to  $-0.019$ ,  $P = 0.029$ ).

**ADHD and TD.** There was a significant effect for overall EF domain differences ( $N = 12$  studies,  $k = 60$  outcomes,  $g = 0.52$ , 95% CI = 0.22 to 0.82,  $P < 0.001$ ,  $t^2 = 0.26$ , prediction interval  $-0.53$  to  $1.56$ ), suggesting that the ADHD group showed poorer performance than the TD group. Sub-domain analyses for individual EF domains showed differences in attention ( $N = 7$  studies,  $k = 18$  outcomes,  $g = 0.46$ , 95% CI = 0.12 to 0.80,  $P = 0.008$ ), response inhibition ( $N = 6$  studies,  $k = 13$  outcomes,  $g = 0.46$ , 95% CI = 0.023 to 0.9,  $P = 0.039$ ) and working memory ( $N = 3$  studies,  $k = 6$  outcomes,  $g = 1.49$ , 95% CI = 0.013 to 2.98,  $P = 0.048$ ). Children with TD performed better than children with ADHD on these domains.

**ADHD and SLD.** Studies comparing ADHD and SLD showed no significant results for a combined effect on EF measures ( $N = 33$  studies,  $k = 213$  outcomes,  $g = 0.099$ , 95% CI =  $-0.066$  to  $0.26$ ,  $P = 0.238$ ,  $t^2 = 0.20$ , prediction interval  $-0.80$  to  $1.0$ ). The sub-domain analyses revealed that the SLD group performed better on tests of planning ( $N = 12$  studies,  $k = 20$  outcomes,  $g = 0.58$ , 95% CI = 0.29 to 0.88,  $P < 0.001$ ) and set-shifting ( $N = 10$  studies,  $k = 15$  outcomes,  $g = 0.43$ , 95% CI = 0.09 to 0.76,  $P = 0.013$ ).

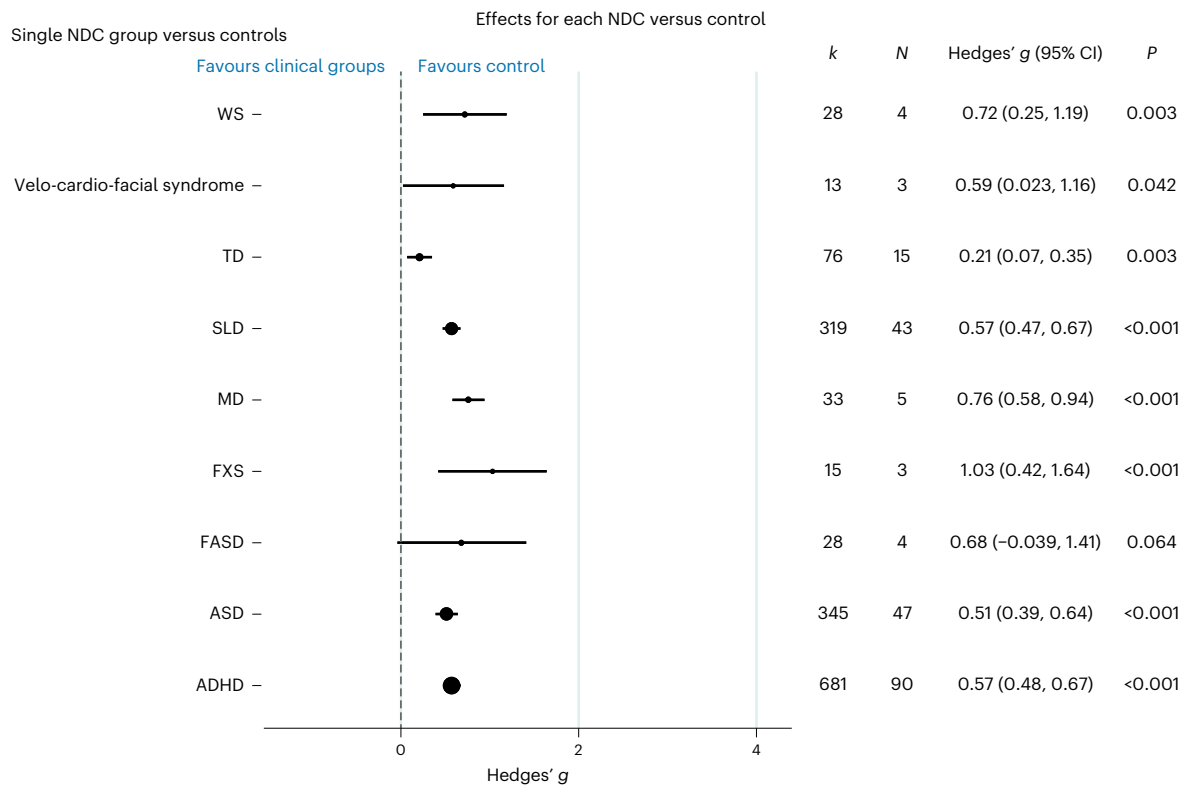
**ADHD and DCD.** Findings from studies comparing ADHD and DCD suggested no overall differences between groups ( $N = 3$  studies,  $k = 18$  outcomes,  $g = -0.37$ , 95% CI =  $-0.94$  to  $0.20$ ,  $P = 0.203$ ,  $t^2 = 0.20$ , prediction interval  $-1.42$  to  $0.68$ ). Sub-domain analyses were not conducted owing to an insufficient number of studies ( $N < 3$  studies).

**ASD and SLD.** A review of studies comparing children with ASD and SLD revealed no differences between groups ( $N = 4$  studies,  $k = 9$  outcomes,  $g = 0.046$ , 95% CI =  $-0.7$  to  $0.79$ ,  $P = 0.904$ ,  $t^2 = 0.53$ , prediction interval  $-1.57$  to  $1.66$ ). No significant results were found when conducting sub-domain analyses. We note that this sub-domain comparison did not include tests of set-shifting.

**Down syndrome (DS) and Williams syndrome (WS).** Studies reporting EF measures between DS and WS suggested no overall significant differences between groups ( $N = 4$  studies,  $k = 32$  outcomes,  $g = -0.035$ , 95% CI =  $-0.40$  to  $0.32$ ,  $P = 0.846$ ,  $t^2 = 0.12$ , prediction interval  $-0.80$  to  $0.73$ ). There were not sufficient studies to explore sub-domain differences.

## Discussion

Our results suggest that EF delay is a transdiagnostic feature of paediatric NDCs, with an overall moderate effect size of delay compared with control. Comparisons between specific NDCs showed few differences on sub-domains. Children diagnosed with TD showed small EF delays overall, while children with ADHD showed larger delays on attention, response inhibition and planning on some comparisons to children with TD and SLDs. Children with ASD showed poorer set-shifting when compared with children with ADHD. We then conducted moderator analyses to understand factors that might contribute to larger effect sizes. Moderator analysis showed that EF delay increased with the percentage of females in each study, but the meaningful contribution of reported sex was marginal. There was no influence of age. While there was no moderating effect of publication year, EF delay was substantially increased with the use of the DSM-5 criteria compared with previous DSM editions. The effect of DSM-5 appeared stable regardless of specific NDCs or measures. Informant-based measures of EF, relative to performance-based measures, showed much larger EF delays. Results suggest EF delay is a transdiagnostic phenotype of NDCs and highlight an urgent need for research that captures data across NDCs. Such data are needed to inform a broader transdiagnostic science of



**Fig. 3 | Forest plot summaries overall effects (Hedges' *g*) for each single NDC versus control.** Here *N* refers to the number of studies reported, and *k* refers to the number of outcome measures reported. Mean effects and their 95% CI are presented by the central black circle and the horizontal line. The size of the black circle reflects the number of studies included in each comparison. The dashed line at *g* = 0 indicates no effect, while the *P* values show whether the effect was statistically significant. The *P* values reported from the random-effects

models are two-sided. FASD, fetal alcohol spectrum disorder; velo-cardio-facial syndrome (including 22q11DS), WS, fragile X syndrome (FXS), motor disorders (MD). Other group comparisons, including DS, neurofibromatosis-1, intellectual disability, Turner syndrome, communication disorders (speech, language and specific language impairments), Prader-Willi syndrome, developmental disability and prenatal alcohol exposure, were not conducted (*N* ≤ 2).

neurodevelopment<sup>38</sup> and to inform how delays impact circuits, symptoms and functioning across child development.

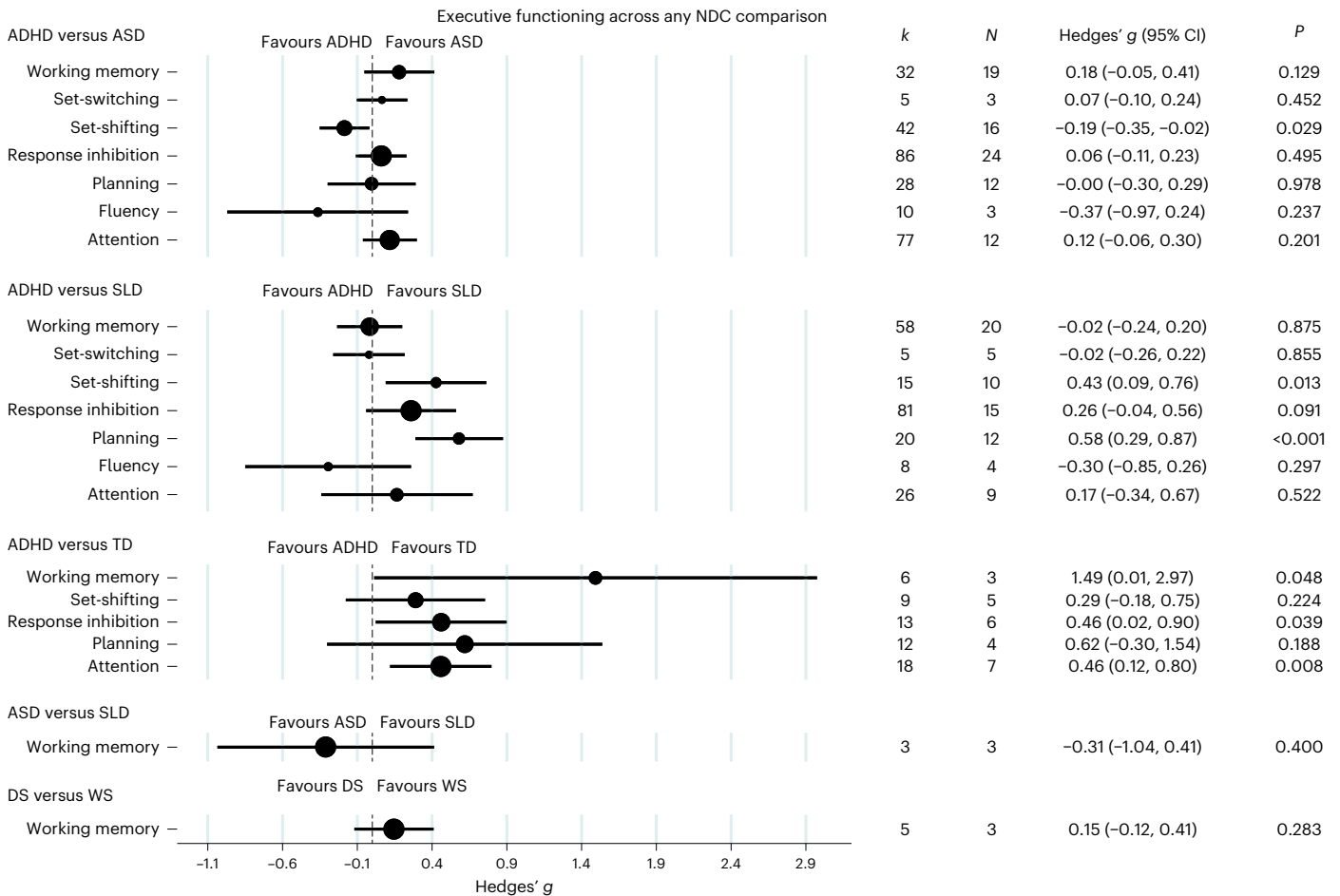
The findings of this study suggest a broad EF delay across all NDCs, with little evidence that specific NDCs were associated with greater overall EF delays. The executive system is believed to be one of the most recently evolved brain systems that operates as a control centre, managing other cognitive abilities (for example, attention, learning, social development and memory)<sup>1,3</sup>. It facilitates flexibility and adaptability<sup>39</sup> to novel and complex situations. While its circuitry is facilitated by the prefrontal cortex, it is highly connected and reliant on other brain circuitry<sup>6</sup>. It follows then that any divergence in brain development in EF systems is likely to impact this interconnected system.

Research has also shown how genetic, neurochemical and environmental<sup>40,41</sup> factors linked to NDCs, as well as critical periods of neurodevelopment, may all influence EF maturation. Importantly, the effect of EF delay in comorbid recruited studies was larger than for studies that only required single NDCs for recruitment. Comorbidity has long been associated with poorer outcomes within reviews of specific NDCs<sup>22</sup>. Such findings suggest that EF delay may provide a sensitive marker to differentiate NDC complexity and progression. A transdiagnostic NDC research framework<sup>38</sup> is now needed to further evaluate modifiers of EF across child development that can better incorporate the genetic and environmental contributors to these effects across development and functioning<sup>42,43</sup>.

Only one difference emerged between specific NDC conditions on the overall EF outcome. Results showed that children with TD had a smaller EF delay compared with children with ADHD, which was consistent with a small effect size of delay for children with TD when

compared with control populations. Past studies have suggested that children with TD may experience greater impacts from circuits that drive motor function in combination with EF delays<sup>44</sup>, and this circuitry may additionally impact functioning. Further research is required to understand such interactions between motor and EF performance and the circuits that underpin outcomes in child development.

While EF delay was largely transdiagnostic, some sub-domain comparison showed differences between NDCs. Children with ADHD showed greater delay on attention, response inhibition and working memory, compared with children with TDs, and planning and set-shifting compared with children with SLDs. Attention disruption is central to the diagnostic criteria of ADHD<sup>45</sup> and the results reinforce this on both informant and performance-based measures. Results offer some support to the specificity of attention-based models of ADHD<sup>15,46,47</sup> that highlight brain regions (such as the dorsal anterior cingulate cortex) involving selective and sustained attention. According to the supervisory attentional system<sup>46</sup>, delayed attention can cause challenges in disengaging from habitual behaviour and exerting novel responses, which, in turn, implicates response inhibition and set-shifting<sup>48</sup>. While such models allude to the unique role attention holds within the EF network and its multi-layered nature<sup>2,49</sup>, others take a multifactorial approach and highlight the trajectory EF domains take from simple to complex EF networks<sup>6,11</sup>. These models emphasize the interconnected relationship attention has with other EF domain areas as EF networks become more mature<sup>50</sup>. In regards to other conditions, children with ASD were more delayed on tests of set-shifting (also known as concept formation) compared with their ADHD counterparts<sup>22,51</sup>. This delay has been linked to the phenotype of



**Fig. 4 | Effect sizes for seven areas of EF across NDC comparisons.** Mean effects and their 95% CI are presented by the central black circle and the horizontal line. The size of the black circle reflects the number of studies included in each

comparison. The dashed line at  $g = 0$  indicates no effect, while the  $P$  values show whether the effect was statistically significant. The  $P$  values reported from the random-effects models are two-sided.

rigidity and repetitive behaviours, as well as difficulties in processing stimuli and social information in complex environments<sup>52,53</sup>. Taken together, these findings inform our understanding that while NDCs share an overall transdiagnostic EF profile<sup>54,55</sup>, nuanced atypicalities could lead to distinct cognitive profiles<sup>22,33,43,51,56</sup>.

Interestingly, DSM edition was a significant moderator, suggesting that the effect size of EF delay for NDCs has almost doubled in size since the introduction and use of the DSM-5. These findings could not be accounted for by any one specific diagnosis, since the removal of a single diagnosis, or removal of comorbid NDCs, from this analysis did not alter the significance. The effect was also found when limiting analysis to only performance or informant reports, so it is unlikely to be due to EF measurement changes. It is also unlikely to be accounted for a real increase in EF delay over the last 40 years, since publication year was not a significant moderator. There has, however, been a gradual increase in the percentage of studies focusing on ASD and a reduction in the percentage of studies focusing on TDs (Supplementary Table 7). Furthermore, there have been important changes in criteria that permit both ASD and ADHD as comorbid conditions. We cannot rule out that a changing focus of NDC research may contribute to this effect. Further prospective research is required to determine whether this effect of DMS-5 assessment criteria is a direct consequence of the changing criteria or changing research priorities since the release of DSM-5.

Assessment type had a significant contribution to the overall effect. Findings of larger EF delays when using performance versus informant-based measures have also been supported by previous literature<sup>22,37</sup>. The administration of performance-based measures

evaluates EF in cumulative scores that tap into accuracy of individual cognitive domains, whereas informant-based measures are often based on reporting observed challenges on rating scales, which are correlated with functionality or real-world behavioural performance of the individual. A large number of studies utilize performance-based measures, which allows for EF domains to be siloed and more precisely and objectively assessed<sup>37</sup>. However, some research has shown that informant EF measures may be better at differentiating those with a clinical diagnosis as it captures the degree and type of impairment in everyday life<sup>22</sup>. Taken together, these findings suggest that research is needed to delineate the use of informant and performance-based measures across development and their utility to understand when informant or performance-based measures are best utilized, with some research suggesting the increased value of using both<sup>37</sup>.

### Limitations

Although many studies control for IQ, we did not include IQ as a primary covariate in our analysis given the overlap between IQ and EF domains. We also note the low number of studies used to identify EF patterns in lesser investigated NDCs. This meta-analysis is limited by the existing literature and the absence of broad investigations of neurodevelopment that incorporates many between-disorder comparisons. This study highlights the urgent need for future work to address these gaps in transdiagnostic developmental research (Supplementary Table 8) where greater focus may be placed on other non-diagnostic contributors to EF delay. The results here show that when a cross-disorder evaluation of EF domains is applied, it leads to a more integrated and nuanced understanding of the EF impairment across sub-domains and overall

in child development. Considering that this review was limited by the number of studies captured within a particular NDC, we note lack of generalizability to lesser investigated NDCs and lack of generalizability to lesser utilized experimental methods (that is, longitudinal studies). Finally, we acknowledge that many of the included studies did not control for comorbidity and the changing criteria in DSM iterations is a limitation for diagnostic specific comparisons given the changes to discrete diagnostic classification over time.

## Conclusion

The conclusions drawn here demonstrate that EF delay is a transdiagnostic feature of paediatric NDCs that increases in severity with greater diagnostic complexity. Future research should target transdiagnostic and distinct EF profiles across the developmental trajectory with a focus on determinants of variance in the EF phenotype, such as its neurobiological underpinnings (for example, functional brain connectivity and neurotransmission) and environmental factors that moderate outcomes across age. Delays in EF may also provide useful and stable transdiagnostic markers for the identification of neurodevelopmental delays that can be used relatively early in life. Subtle differences observed here between NDCs may also further offer potential to develop precision medicine approaches for specifying the circuits underpinning the development of different NDCs, leading to targeted supports for learning, cognition and daily living.

## Methods

This review adheres to the Preferred Reporting Items for Systematic Reviews and Meta-analyses (PRISMA) guidelines<sup>57,58</sup>. The review was prospectively registered with PROSPERO (CRD42020210785).

### Study selection

The review included peer-reviewed studies in any language published from 1980 to 19 February 2024 (see Fig. 1 for the PRISMA diagram). Cross-sectional and longitudinal cohort studies were considered if they included children under 18 years of age and reported on at least two NDC comparison groups (see Supplementary Table 1 for search strategy). NDCs were identified if they were listed within the DSM-5 and assessed with reliable diagnostic measures.

### Search strategy and study variables

The literature search was conducted through MEDLINE, Embase and PsycINFO databases using detailed search criteria of EF measures (that is, 'BRIEF', 'Wisconsin Card Sorting Test', 'Go-No-Go task' and 'Stroop test') and a full range of paediatric NDCs (for example, 'ASD', 'SLD', 'ADHD' and 'TD'). Terms used to describe NDCs in previous DSM editions were also included in the search strategy, as were NDCs identified in The SAGE Handbook of Developmental Disorders<sup>59</sup>. See Supplementary Tables 1 and 2 for the search strategy and more detailed descriptions of included EF measures and NDCs. Each NDC comparison then needed sufficient data (at least three studies) to be included for further analysis. The search strategy captured information such as whether the diagnosis was made through a clinical interview, DSM criteria, standardized measures or other modes of classification. In addition, studies must have included established outcome measures looking at either informant measures or performance measures of EF (Supplementary Table 1).

The authors (A.K.S., C.S., N.P. and M.M.) screened for initial eligibility based on title and abstract, and then for the full-text screenings using Covidence (Extraction 1)<sup>60</sup>. An independent reviewer addressed any disagreements (E.A.D.). The inter-rater reliability, kappa value, was 0.87, indicating high agreement between reviewers<sup>61</sup>.

In data extraction, EF measure outcomes (that is, commissions or omissions errors in a task like Go-No-Go) reported in each comparison group were extracted as mean values and standard deviation scores at a single time point (or baseline results were extracted in the case of

longitudinal studies). For studies reporting multiple measures derived from psychometric tests, experimental tasks and/or self/informant measures, each outcome measure was considered separately. Where there were missing data, efforts were made to contact authors regarding missing data by email at least once; however, no author was able to address these requests. In addition, all study authors were contacted for unpublished data to mitigate 'the file drawer effect'. One author was able to address this request and their data are included in the results (Supplementary Table 3). The data extraction process was completed using a customized Excel spreadsheet (version 2404).

### Quality assessment

Quality review of studies was completed by assessors (A.K.S., C.S., M.M. and N.P.) using the Checklist for Cross-Sectional and Cohort Studies within the JBI Critical Appraisal Tools. Studies were based on the JBI critical appraisal tool with 80% of ratings double coded. Agreement between raters (kappa value) was 0.97, indicating high agreement between reviewers<sup>61</sup>.

### Data items

Group-level summary data (for example, sample size, means and standard deviations of each group) from each was extracted for all measures reporting outcomes for EF. When studies reported data from multiple measures or subgroups, all eligible data were extracted. When studies reported outcomes across several manuscripts, data were combined into a single study to avoid double-counting of studies. All meta-analyses were conducted with the metafor (version 3.8-1), meta (version 6.5-0), dplyr (version 1.1.2), readxl (version 1.4.3) packages in RStudio (2023.06.1 Build 524) and R (version 4.3.3) using multivariate models in order to account for non-independence among effect sizes within studies. The unit of analysis utilized within this model was the standardized mean difference (calculated as Hedges' *g*) on each measure between each comparison group. When making NDC versus control group comparisons, a positive effect size indicated that the control group performed better than the NDC group. The data analysis was planned a priori and was completed in four stages. The initial analysis combined all EF outcomes to assess the overall EF effect size across NDCs when compared with controls. Publication bias was assessed using Egger's test and trim-and-fill methods and illustrated in funnel plots. The second analysis examined subgroup comparison of the individual EF domains. The third analysis examined the estimated effect size of difference for each NDC compared with control conditions. The next step involved examining between study variability and moderator impact for overall EF and individual EF domains. This included 'type of measure', 'gender', 'age', 'DSM edition' and 'year of publication', which were assessed as covariates in meta-regression analyses. The final step involved the analyses of individual EF domains for each of the NDC comparison groups (that is, ASD versus ADHD and so on). By convention, effect sizes (Hedges' *g*) with 95% CI are described as small ( $\leq 0.30$ ), moderate ( $> 0.30$  and  $< 0.60$ ) and large ( $\geq 0.60$ ). Heterogeneity (that is, variance between studies) was quantified using  $I^2$ . The variance explained by the moderators in the meta-regression model was quantified as  $R^2$  and formally tested using Cochran's *Q*.

### Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

### Data availability

The data used to undertake this systematic review and meta-analysis are freely available ([https://github.com/CarterSunUSYD/Transdiagnostic\\_EF\\_meta.git](https://github.com/CarterSunUSYD/Transdiagnostic_EF_meta.git)). Source data are provided with this paper.

### Code availability

The R codes used for the meta-analysis are freely available ([https://github.com/CarterSunUSYD/Transdiagnostic\\_EF\\_meta.git](https://github.com/CarterSunUSYD/Transdiagnostic_EF_meta.git)).

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## Author contributions

A.K.S., A.J.G. and E.A.D. collaborated on the initial protocol design and development of this study and subsequent paper. A.K.S. and C.S. were involved in article searches, screening, coding of article risk-of-bias variables and data extraction using a co-designed template. M.M. and N.P. supported the updated search, screening, data extraction and critical appraisal with C.S., under the close guidance of A.J.G. and E.A.D. Data analysis in R was primarily completed by C.S., with contributions from A.K.S. A.K.S., C.S. and A.J.G. wrote the initial draft paper and K.A.B., E.A.D., A.L., N.P. and M.M. all provided extensive comments on drafts. The production of statistical figures and visualization was assisted by the Sydney Informatics Hub at the University of Sydney. All co-authors reviewed and provided substantial input to finalize the paper for submission.

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## Competing interests

The authors declare no competing interests.

## Additional information

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
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**Kelsie A. Boulton**<sup>1,2</sup> & **Adam J. Guastella** <sup>1,2</sup> 

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### Software and code

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**Data collection** Manuscript screening and data extraction were completed through Covidence (Extraction 1) (<https://www.covidence.org>) and a customised excel spreadsheet (Version 2404).

**Data analysis** Meta-analysis was conducted through open source software, RStudio (2023.06.1 Build 524) and R (version 4.3.3), and customised codes are available on GitHub ([https://github.com/CarterSunUSYD/Transdiagnostic\\_EF\\_meta.git](https://github.com/CarterSunUSYD/Transdiagnostic_EF_meta.git)). The package used to conduct the meta-analysis was metafor (Version 3.8-1), meta (Version 6.5-0), dplyr (Version 1.1.2), readxl (Version 1.4.3).

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The data used to undertake this systematic review and meta-analysis are freely available on GitHub ([https://github.com/CarterSunUSYD/Transdiagnostic\\_EF\\_meta.git](https://github.com/CarterSunUSYD/Transdiagnostic_EF_meta.git)). Databases used in this study including MEDLINE, Embase and PsycINFO.

## Research involving human participants, their data, or biological material

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Reporting on sex and gender	The meta-analysis extracted gender information from the original text of each eligible studies.
Reporting on race, ethnicity, or other socially relevant groupings	This review did not explicitly extract race and ethnicity information for the quantitative meta-analysis.
Population characteristics	This review included all studies that measured executive functioning using established tests in participants under 18 years of age and describing populations comparing two or more paediatric neurodevelopmental conditions.
Recruitment	Participant demographics and diagnostic conditions were extracted directly from each included study. Manuscripts reporting on the same cohort were nested into one study group to reduce potential bias from repeating samples in the meta-analysis.
Ethics oversight	Not applicable for meta-analysis.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

## Field-specific reporting

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## Behavioural & social sciences study design

All studies must disclose on these points even when the disclosure is negative.

Study description	This quantitative meta-analysis includes both cross-sectional and longitudinal cohort studies.
Research sample	This review includes children under 18 years of age and reported on at least two neurodevelopmental conditions listed within the DSM-5 and assessed with reliable diagnostic measures. We searched all studies available on databases. These studies were chosen to inform our understanding of published data comparing different neurodevelopmental conditions.
Sampling strategy	The review included peer-reviewed studies in any language published from 1980 to the 19th of February 2024. Studies were included if they met our inclusion criteria (under 18 years of age, at least two neurodevelopmental conditions recruited in each published articles). Meta-analysis was conducted when at least 3 studies were reported in each comparison group. This was informed by the presence of underlying heterogeneity as well as sufficient number and balance of studies within subgroups, enabling cross-condition analyses within each cell.
Data collection	Researchers were not blinded to experimental condition during data extraction. In data extraction, EF measure outcomes (i.e., commissions or omissions errors in a task like Go-No-Go) reported in each comparison group were extracted as mean values and standard deviation scores at a single time point (or baseline results were extracted in the case of longitudinal studies). For studies reporting multiple measures derived from psychometric tests, experimental tasks and/or self/informant measures, each outcome measure was considered separately. Where there was missing data, efforts were made to contact authors regarding missing data by email at least once, however, no author was able to address these requests. In addition, all study authors were contacted for unpublished data to mitigate 'the file drawer effect'. One author was able to address this request and their data is included in the results.
Timing	The search of eligible studies was updated in February 2024.

Data exclusions	Five studies were removed as they contributed to data asymmetry ( $g > 2$ ). Comparisons between neurodevelopmental conditions were not conducted if there were less than three eligible studies. Details are outlined in the methods section and was preregistered on PROSPERO.
Non-participation	No participants were recruited in this systematic review and meta-analysis.
Randomization	Not applicable as this is a systematic review and meta-analysis study.

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We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

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### Methods

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## Plants

Seed stocks	Not applicable.
Novel plant genotypes	Not applicable.
Authentication	Not applicable.