Eczema and related atopic diseases are associated with increased symptom severity in children with autism spectrum disorder

C. Jameson1, K. A. Boulton1,2, N. Silove2,3 and A. J. Guastella1,2✉

© The Author(s) 2022

Growing evidence indicates that autism spectrum disorder (ASD) has diverse genetic, neurological, and environmental factors that contribute to its neurodevelopmental course. Interestingly, childhood ASD is often accompanied by skin disorders, such as eczema, and other related atopic manifestations. This link may be due to the shared embryonic origin of epidermal and neural tissue. Accordingly, we consider the potential influence of a skin-brain co-vulnerability and ensuing atopic cascade on ASD symptomatology by investigating whether atopic disorders (asthma, allergies, eczema and hay fever) are associated with increased symptom severity in children with ASD. Overall, 45 atopic and 93 non-atopic children with ASD were assessed using the ADOS-2 on scores of total, social and non-social symptoms. Differences in ASD symptom severity were further evaluated as a function of atopic disease type. Atopic children displayed greater symptom severity overall and in the social domain, relative to non-atopic participants. Atopic children were 2.4 times more likely to experience overall impairments classified within the ADOS-2 highest-level severity bracket and 2.7 times more likely to show social difficulties in this range. Moreover, those reporting comorbid eczema displayed increased symptom severity relative to both their non-atopic peers and those reporting asthma and allergies. Taken together, findings indicate that atopic disorders, and particularly comorbid eczema, are associated with increases in ASD symptom severity. Findings provide grounds for future investigations into this link between childhood skin diseases and ASD symptom severity to advance our understanding of neurodevelopment and to develop targeted assessment and intervention opportunities.

Translational Psychiatry (2022) 12:415; https://doi.org/10.1038/s41398-022-02185-5

INTRODUCTION

Autism spectrum disorders (ASD) represent a group of neurodevelopmental conditions characterised by reduced social responsivity and restricted and stereotyped interests and behaviours [1]. The aetiology of ASD is underpinned by a complex interplay between neurological, genetic, and environmental factors that culminate in heterogeneous clinical presentations [2]. Consequently, ASD is increasingly recognised as representing a broad range of neurodevelopmental disorders and subtypes that are associated with diverse physical and mental health comorbidities [3]. The role of atopic diseases in neurodevelopment has received growing attention as a potential modifier of neurodevelopment [4–6]. Although atopic disorders are relatively common, affecting between 5 and 15% of all children, accumulating evidence indicates prevalence of such diseases is increased amongst those with ASD [6, 7]. For instance, in a recent meta-analysis including over one million pooled participants, Tasi et al. [8] determined that children with ASD were 1.49 times more likely to experience atopic dermatitis relative to their typically developing peers (95% CI: 1.2–1.8). Similarly, Magalhães et al. [9] reported that 86.6% of participants with Asperger’s syndrome (N = 15) had at least one atopic disease, compared to <7% in a neurotypical cohort. Further, children with ASD appear 4.5 times more likely to report food allergy (95% CI: 3.0–7.0) [7], and experience significantly increased rates of asthma and allergic rhinitis relative to those without ASD (OR 1.69, 95% CI: 1.11–2.59 and OR 1.66, 95% CI: 1.49–1.85 respectively) [10]. Notably, these findings have been reported across large and multi-racial samples, suggesting this link extends beyond ethnic or cultural differences. Overall, whilst a positive association between atopic manifestations and ASD has been repeatedly observed, this association with neurodevelopmental outcomes remains poorly understood.

Research examining whether atopic disorders are associated with the severity of neurodevelopmental delay in ASD is scarce. In the only study to date, Mostafa, Hamza and El-Shahawi (2008) evaluated reports of atopic disease (bronchial asthma, atopic dermatitis, and allergic rhinitis) amongst children diagnosed with autism (N = 50) relative to an age- and sex-matched neurotypical cohort (N = 50) [11]. The researchers observed that children with ASD who scored in the severe range on the Childhood Autism Rating Scale (CARS) (N = 20) showed significantly increased rates of atopic disease (80%), compared to both children without an autism diagnosis (10%) and those with mild to moderate autism presentations (33%). The authors, however, did not investigate
symptom domains associated with ASD, nor did they distinguish between cutaneous and non-cutaneous atopic disorders, despite growing recognition of a skin-brain axis stemming from a shared embryonic origin [12, 13].

The skin-brain axis is of emerging interest in the field of neurodevelopment and has prompted discussion as to whether atopic eczema’s apparent link with ASD may be unique from that of non-cutaneous atopic conditions. For instance, from 3 to 4 post-conception weeks, the brain, skin, and skin appendages develop in synchronisation in utero, all originating from one of three primary germ layers, the ectoderm [12]. Accordingly, whilst the neuroectoderm differentiates to form early derivatives of the brain, such as the neural tube, the ventral ectoderm develops into a monolayer epidermis [13]. It, therefore, seems plausible that atopic eczema may be uniquely associated with neurodevelopmental outcomes relative to non-cutaneous atopic disorders.

Interestingly, cutaneous syndromes also often mark the onset of the “atopic march”, with epidermal dysfunction facilitating subsequent allergic sensitisation and atopic respiratory disorders [14]. Accordingly, in eczematous patients, sub-optimal tight junctions in the epidermis may be accompanied by disrupted barrier function in the airway epithelium or intestinal mucosa, easing the transcutaneous invasion of allergens and inducing a maladaptive immune response [15]. Notably, this systemic and hyper-sensitive atopic response has been observed to reduce the integrity of the blood-brain-barrier, facilitating excess cytokine entry into the brain and potentiating neuroinflammation [16–18]. Accordingly, previous studies have reported positive correlations between elevated plasma and serum concentrations of atopic inflammatory markers, including interleukin (IL)-4 and IL-17A, and increasingly pronounced behavioural impairments in children with ASD [19].

Therefore, despite acknowledging a likely skin-brain connection and appreciating the influence of atopic barrier dysfunction on neurodevelopment, it remains unclear whether atopic eczema may influence neurodevelopment in a manner distinct from related respiratory and gastrointestinal atopies.

In this study, we explored whether atopic comorbidities (asthma, allergies, eczema and hay fever) were associated with increased symptom severity in a cohort of children with ASD. Specifically, we aimed to determine whether symptom severity differed between children with ASD, with and without atopic complaints, including overall symptoms as well as social affect (SA) and restricted and repetitive behaviours (RRB). Additionally, we aimed to identify whether reporting an atopic condition increased a child’s risk of displaying symptoms classified within the severe range and determined whether ASD-related behavioural impairments varied as a function of atopic disease type for each domain. We hypothesised that, relative to those without an atopic complaint, children with ASD and an atopic comorbidity would present with increased ASD symptomatology overall and across both SA and RRB domains. Further, we proposed that atopic children would be more likely to display symptoms classified as severe, compared to children without an atopic comorbidity. Additionally, given the proposed role of cutaneous syndromes as precursors to related atopic conditions, we hypothesised that children with comorbid eczema would show increased symptom severity in each of these domains relative to both non-atopic children and those reporting other prevalent atopic conditions.

METHOD
Participant recruitment
Participants included children with ASD who were recruited from the Clinic for Autism and Neurodevelopment (CAN) Research, located at the Brain and Mind Centre, University of Sydney (n = 105) and the Child Development Unit, located in Westmead’s Children’s Hospital (n = 35). Overall, 140 children were included in the study (n = 110 males, 30 females; mean age 6.14 [standard deviation = 2.62]). All participants were ambulatory with no suspected vision or hearing impairments. The Sydney Children’s Hospitals Network (SCHN) Human Research Ethics Committee (HREC) (LNR/17/SCHN/293), University of Sydney HREC (2013/502) or Sydney Local Health District (SLHD) HREC (18/RPAH/157) provided approval. Informed consent was obtained from all parents or caregivers before participation.

Participant diagnoses of ASD were confirmed using the Autism Diagnostic Observation Schedule, Second Edition (ADOS-2) [20]. All participants were assessed using either ADOS-2 Module 1 (for children with few-to-no words, n = 70), Module 2 (for those with phrase speech, n = 40) or Module 3 (for verbally fluent children and young adolescents, n = 28). The presence of any auto-immune, neurological (e.g., seizures) or medical conditions were also considered exclusion criteria for all participants [6, 19]. Six children were excluded due to reports of systemic lupus erythematosus (n = 1), an unspecified thyroid disease (n = 1), Klinefelter syndrome (n = 1), Phelan–McDermid syndrome (n = 1), Prader–Willi syndrome (n = 1) and a seizure disorder (n = 1).

Statistical analyses
Shapiro–Wilk normality tests indicated that several variables were non-normally distributed, as shown in Supplementary Table 1. Consequently, all analyses were conducted using bootstrapping (5000 resamples), enabling parametric statistical procedures to be applied. Demographic characteristics of children in the atopic and non-atopic cohort, as well as children reporting eczema relative to other atopic conditions, were compared using Welch two independent-sample t-tests and χ² tests, as appropriate. Additionally, Fisher’s exact tests were used to compare sleep behaviours between the atopic and non-atopic groups due to the reduced sample of available data, with sleep history only collected at one study site.

Differences in meanADOS CSS-SA, CSS-RRB and CSS scores for children in the atopic and non-atopic cohort were determined using Welch two independent-sample t-tests. The distribution of participants’ ADOS scores, including mean ± standard deviation (SD), were subsequently graphed using a truncated violin plot. The same statistical procedures were applied to determine differences in mean ADOS CSS-SA, CSS-RRB and CSS scores for children with comorbid eczema relative to age- and sex-matched non-atopic and asthma/allergies cohorts. Additionally, binomial logistic regressions were used to investigate whether ≥1 atopic comorbidity increased the likelihood of a participant presenting with ASD symptomatology exclusively defined within the severe range (score ≥7 on the CSS-SA, CSS-RRB and overall CSS scales), generating odds ratios (ORs) and 95% confidence intervals (CIs). Participants with missing data for a particular outcome were excluded from the analysis for that measure.

C. Jameson et al.
Table 1. Demographic characteristics of children with ASD in the atopic and non-atopic cohorts.

|                        | Full cohorta | Atopic cohort | Non-atopic cohort | p value |
|------------------------|--------------|---------------|-------------------|---------|
| Number of participants, n (%) | 140 (100)    | 47 (33.57)    | 93 (66.43)       | –       |
| Asthma                  | 15 (10.70)   | –             | –                 | –       |
| Allergies               | 23 (16.40)   | –             | –                 | –       |
| Eczema                  | 26 (18.60)   | –             | –                 | –       |
| Hay fever               | 2 (1.40)     | –             | –                 | –       |
| Sex, n (%)              |              |               |                   | 0.366b  |
| Male                   | 110 (78.60)  | 39 (82.98)    | 71 (76.34)       | –       |
| Female                 | 30 (21.40)   | 8 (17.02)     | 22 (23.66)       | –       |
| Age (years), mean ± SD  | 6.15 ± 2.62  | 6.13 ± 2.69   | 6.14 ± 2.61      | 0.983c  |
| Sleep behaviours, n (%) |              |               |                   |         |
| Poor sleeping pattern   | 22 (33.85)   | 6 (35.30)     | 16 (33.33)       | 0.883d  |
| Snoring                | 10 (15.38)   | 4 (23.53)     | 6 (12.50)        | 0.434d  |
| Trouble falling asleep | 13 (20.00)   | 4 (23.53)     | 9 (18.75)        | 0.729d  |
| Dislikes sleeping alone | 15 (23.08)   | 6 (35.30)     | 9 (18.75)        | 0.191d  |
| Waking overnight        | 17 (26.15)   | 3 (17.65)     | 14 (29.17)       | 0.523d  |
| Maternal infection, n (%) | 21 (15.22) | 8 (17.02) | 13 (13.98) | 0.560d |

SD standard deviation.

aPercentage of missing data in each measure for full cohort: maternal infection 1.43%. Sleep behaviours 47.14%. Percentages are expressed as a fraction of the number of participants in each group and may not total 100 due to missing data for some variables. p values are reported for comparisons between the atopic and non-atopic cohort using the χ² test.

bBootstrapped Welch two-sample t-test.
cOr Fisher’s exact tests.
dAs appropriate.

Statistical analyses were performed using R for macOS 10.13, version 4.1.0 (RStudio, Boston) and graphs were created using GraphPad Prism version 9.0.0 for macOS (GraphPad Software, California). A two-sided value of p < 0.05 was considered statistically significant. Sample size was based on a sample of convenience but was powered to a similar degree to past studies examining related variables [26].

RESULTS

Participant characteristics

Table 1 presents the demographic characteristics of the sample. Of the total 140 children with ASD included in the study, atopic conditions of any type were found in 47 (33.57%) participants (atopic cohort, n = 39 males, 8 females; mean age 6.13 [standard deviation = 2.69]). The remaining 93 (66.43%) participants did not report any atopic conditions, serving as a comparison group (non-atopic cohort, n = 71 males, 22 females; mean age 6.14 [standard deviation = 2.61]). There were no statistically significant differences between the atopic and non-atopic cohorts in sex, age, or reports of maternal infection (p > 0.05). Additionally, of the 66 participants with available sleep disturbance data (n = 17 atopic, n = 49 non-atopic), no significant differences were detected between the atopic and non-atopic groups in the occurrence of any undesirable sleep behaviours.

Scale scores and assessments of normality

Supplementary Table 1 shows descriptive data for the sample, including scale scores and assessments of normality. The Shapiro–Wilk normality test on all participants revealed a normal distribution for age, with the variable showing a positive skew towards early childhood. Similarly, ADOS-2 CSS total, CSS-SA and CSS-RRB scores showed non-parametric distributions, with a negative skew towards scores indicative of a high level of impairment.

ASD phenotypic outcomes

We then looked at differences in ADOS CSS and CSS-SA scores as a function of atopic disease type. In these analyses, the full cohort was stratified into three subgroups: eczematic (n = 26), non-atopic (n = 24) and asthma/allergies (n = 17), individually matched on
Table 2. Generalised linear models evaluating the presence of at least one atopic condition as a predictor of moderately high to severe ASD-related symptomatology in children.

|                          | β   | SE  | z-value | OR  | 95% CI       | P (Wald’s test) |
|--------------------------|-----|-----|---------|-----|--------------|-----------------|
| ADOS CSS ≥7              | 0.81| 0.41| 1.98    | 2.25| 1.01–5.04    | 0.047*          |
| ADOS CSS-SA ≥7           | 1.08| 0.43| 2.49    | 2.93| 1.26–6.84    | 0.013*          |
| ADOS CSS-RRB ≥7          | 0.17| 0.45| 0.38    | 1.19| 0.49–2.86    | 0.704           |

CSS calibrated severity scores, SA social affect, RRB restricted and repetitive behaviour.
ADOS-2 autism diagnostic observation schedule, second edition; CSS, SA, RRB. Percentage of missing data in each measure for full cohort: maternal infection 1.45%, ADOS-2 all scales 3.6%. *p < 0.05.

Table 3. Demographic characteristics of children with ASD and comorbid eczema, non-atopic or comorbid asthma/allergies (n = 67).

|                          | Eczema cohort | Non-atopic cohort | Asthma/allergies cohort |
|--------------------------|---------------|-------------------|-------------------------|
| Number of participants   | 26            | 24                | 17                      |
| Sex, n (%)               |               |                   |                         |
| Male                     | 22 (84.6)     | 21 (87.5)         | 14 (82.4)               |
| Female                   | 4 (15.4)      | 3 (12.5)          | 3 (17.6)                |
| Age (years), mean ± SD   | 6.10 ± 2.84   | 6.07 ± 2.85       | 5.64 ± 2.30             |

SD standard deviation.

age and sex where possible. Table 3 presents the demographic characteristics for each of these cohorts. There were no significant differences between the eczema and non-atopic or asthma/allergies groups in terms of sex (p = 0.769 and p = 0.844 respectively) or age (p = 0.971 and p = 0.557).

As expected, children with ASD and comorbid eczema received significantly higher mean scores on the ADOS CSS total scale (7.95 ± 1.62) relative to those in the non-atopic cohort (5.71 ± 1.27; t = 5.35, df = 39.29, p < 0.001). The comparison between the eczema and asthma/allergies subgroups was only significant in a one-tailed test (p = 0.034; two-tailed test: t = 1.88, df = 36.86, p = 0.069), as shown in Fig. 2A. As depicted in Fig. 2B, children with ASD and comorbid eczema also displayed significantly higher mean scores on the ADOS CSS-SA (7.73 ± 1.58) compared to those...
in the non-atopic cohort (5.75 ± 1.96; \( t = 3.78, \text{df} = 43.31, p = 0.001 \)), yet not those in the asthma/allergies cohort (7.12 ± 1.32; \( p = 0.197 \)).

**DISCUSSION**

To our knowledge, this is the first study to examine whether children with ASD and atopic conditions display increased autism-related symptom severity, with a particular focus on comorbid eczema. Findings showed that children with ASD and atopic disease experienced more pronounced symptoms overall and on the ADOS social domain, relative to those children with ASD without atopic complaints. That is, relative to the non-atopic cohort, atopic children were 2.3 times more likely to experience overall symptoms classified within the ADOS-2 highest-level severity bracket and 2.9 times more likely to show social difficulties in this range. Furthermore, the current findings suggested that ASD symptom severity may vary as a function of atopic disease type, with eczematic children displaying a greater degree of overall impairment relative to both age- and sex-matched non-atopic participants and those reporting other prevalent atopies (asthma/allergies). These findings support further investigations of a possible early life link between cortical and epidermal development, underpinned by a possible skin-brain-immune connection. This research may lead to the identification of sub-types of children that may have different contributing factors to neurodevelopment and, as has been highlighted recently [27, 28], opportunities for novel therapies to improve outcomes.

These findings broadly align with previous research in paediatric populations that have shown pro-inflammatory mediators, which are hallmarks of the atopic response, to correlate with the severity of behavioural outcomes in ASD [19, 29]. One plausible explanation for these findings may relate to the epithelial barrier dysfunction associated with atopic conditions [15, 19]. For example, aberrant tight junction formation in the airway, intestinal and/or cutaneous epithelium of children with atopic diseases may have eased the invasion of external irritants, inducing a hypersensitive immune response [4, 15, 30]. Given that the initial phases of this response are believed to be predominately mediated by Th2 cells, interleukin (IL)-4 and IL-13 are released, promoting isotypic class switching and immunoglobulin-E production in B cells [4]. Consequently, Th1 immuno-active cytokines and chemokines are released, and mast cells are recruited to synthesise additional prostaglandins, tumour necrosis factor and vascular endothelial growth factor [4, 30]. Interestingly, previous research indicates that, when chronic, this atopic response is associated with reduced integrity of the BBB, likely increasing cytokine entry into the brain, and facilitating cortical disruption in circuits regulating early life development, relative to non-atopic participants [16, 17]. Overall, the convergence between our findings and prior studies offers support for the notion that atopic conditions could potentially influence social outcomes in children with ASD, however further prospective, longitudinal research is needed to establish directional and causative pathways.

Alternatively, the current findings may be underpinned by a model whereby severe ASD-related symptomatology, including higher sensory sensitivity, may exacerbate cutaneous affictions. For instance, given that children with autism regularly report hypo- and hyper-responsivity to touch, the observed difference in symptom severity between eczematic and asthmatic/allergic participants may be underpinned by ASD-related differences in sensory processing [31, 32]. In typically developing eczematic children, excessive scratching and severe pruritis have been correlated with intense irritation and seclusion [33]. It seems plausible that both cutaneous symptoms and their associated adverse behavioural outcomes may be exacerbated in children with ASD who present with severe tactile sensitivity. Results may indicate that severe sensory sensitivity may be associated with intensified cutaneous discomfort and act as a mediator of worsened behavioural outcomes in eczematic children with ASD. Future research is warranted to examine the prevalence of tactile processing difficulties amongst children with ASD and eczema relative to other atopic conditions, and whether this relates to ASD symptom severity.

Our study further explored possible alternative determinants proposed in previous literature to mediate variation in ASD-related symptom severity. For instance, prenatal exposure to pro-inflammatory mediators, such as that stemming from maternal immune activation, has been associated with an increase in the severity of emotional and behavioural impairments in children with ASD [26, 34]. The current study, however, detected no significant differences in reports of maternal infection during pregnancy between the atopic and non-atopic cohorts. Consequently, gestational immune activation was unlikely a key factor.
underlying the observed differences in ASD symptom severity between these groups. Similarly, prior literature has pointed to an association between sleep disorders and adverse behavioural outcomes in children with atopic conditions [4, 14, 35]. In our study, however, no significant differences in sleep pattern were determined between our atopic and non-atopic cohorts, indicating that abnormal sleep behaviours were unlikely a confounder in the observed difference in symptom severity between the two groups.

By contrast, the results of this study did not support a link between non-social symptoms, namely restricted and repetitive behaviours, and reports of atopic disease. These findings are consistent with the difficulties of previous research to detect an effect in the RRB domain, with a distinct heterogeneity in the prevalence and severity of repetitive and stereotyped actions possibly attributable to these behaviours being less naturally evoked [25]. Further, prior research has established ceiling effects in this domain for young children displaying lower-percentile language ability, which resembles the current sample [23]. Overall, our study points to the need for future research into alternative severity metrics for evaluating restricted and repetitive interests and behaviours in paediatric populations.

The present study has several limitations. Our sample was one of convenience, consisting of children attending assessment services at university and hospital clinics. Accordingly, this population may not be representative of a broader community sample of children with ASD. Second, the current study is cross-sectional. As a result of the relatively small sample size, some findings, such as the difference in ASD symptom severity between children with eczema and children with other atopic conditions, only reached statistical significance [4, 14, 35]. In our study points to the need for future research into alternative severity metrics for evaluating restricted and repetitive interests and behaviours in paediatric populations.

Additionally, instances of atopic diseases deviated notably from those reported in previous literature, with a possible underestimation observed for hay fever [6, 7]. This may be due to the reliance on self-report within a medical interview for atopic disease, rather than evaluating allergen-specific serum IgE or skin prick testing, possibly enabling some misclassification of atopic and non-atopic participants. Our study further did not collect biological measures of immune activation, which would have improved our understanding of the potential biological mechanisms at play. Additionally, given the proposed role of gut microbiota and metabolites in the pathophysiology of ASD [2] and the development of potential treatments targeting gut-derived metabolites to improve symptoms associated with ASD [27], further investigations should consider the role of gastrointestinal features alongside atopic features.

Despite these limitations, the present study provides valuable insights into possible determinants and predictors of neurodevelopment and ASD symptomatology. Overall, our findings support the notion that atopic conditions, particularly cutaneous syndromes, may be related to the severity of overall and social domain outcomes in children with ASD. This study adds to the growing body of literature examining atopic disorder’s association with ASD by proposing that atopic disease may be a factor in the phenotypic expression of autism symptom severity in a subgroup of children.

REFERENCES

1. American Psychiatric Association. Diagnostic and statistical manual of mental disorders. 5th edn. Arlington, VA: American Psychiatric Association; 2013.
2. Maiuolo J, Giolli M, Musolino V, Carresi C, Scarano F, Nucera S, et al. The contribution of gut microbiota-brain axis in the development of brain disorders. Front Neurosci. 2021;15:616883.
3. Srikantha P, Mohajen MH. The possible role of the microbiota-gut-brain-axis in autism spectrum disorder. Int J Mol Sci. 2019;20:2115.
4. Chen MH, Su TP, Chen YS, Hsu JW, Huang KL, Chang WH, et al. Is atopy in early childhood a risk factor for ADHD and ASD? A longitudinal study. J Psychosom Res. 2014;77:316–21.
5. Liao TC, Lien YT, Wang S, Huang SL, Chen CY. Comorbidity of atopic disorders with autism spectrum disorder and attention deficit/hyperactivity disorder. J Pediatr. 2016;171:248–55.
6. Xu G, Sneteslaar LG, Jing J, Liu B, Strathern L, Bao W. Association of food allergy and other allergic conditions with autism spectrum disorder in children. JAMA Netw Open. 2018;1:e180279.
7. Gurney JG, McPheeters ML, Davis MM. Parental report of health conditions and health care use among children with and without autism: National Survey of Children’s Health. Arch Pediatr Adolesc Med. 2006;160:825–30.
8. Tsai TY, Chao YC, Hsieh CY, Huang YC. Association between atopic dermatitis and autism spectrum disorder: a systematic review and meta-analysis. Acta Derm Venereol. 2020;100:adv00146.
9. Magalhães ES, Pinto-Mari F, Bastos-Pinto S, Pontes AT, Prado EA, deAzevedo LC. Immune allergic response in Asperger syndrome. J Neuroimmunol. 2009;216:108–12.
10. Miyazaki C, Koyama M, Ota E, Swa T, Amiya RM, Milunde LB, et al. Allergies in children with autism spectrum disorder: a systematic review and meta-analysis. Rev J Autism Dev Disord. 2015;2:374–401.
11. Mostafa GA, Hamza RT, El-Shahawi HH. Allergic manifestations in autistic children: relation to disease severity. J Pediatr Neurol. 2015;06:115–23.
12. Donovan MF, Cascella M. Embryology, Weeks 6-8. StatPearls; 2021.
13. Rewane A, Munakorn, S. Embryology, Central Nervous System, Malformations. StatPearls; 2021.
14. Simpson EL. Comorbidity in atopic dermatitis. Curr Dermatol Rep. 2012;2:129–38.
15. Silverberg JI. Selected comorbidities of atopic dermatitis: atopy, neuropsychiatric, and musculoskeletal disorders. Clin Dermatol. 2017;35:360–6.
16. Jackson-Cowan L, Cole EF, Silverberg JI, Lawley LP. Childhood atopic dermatitis is associated with cognitive dysfunction: A National Health Interview Survey study from 2008 to 2018. Ann Allergy Asthma Immunol. 2021;126:661–5.
17. Masi A, Glozier N, Dale R, Guastella AJ. The immune system, cytokines, and biomarkers in autism spectrum disorder. Neurosci Bull. 2017;33:194–204.
18. Masi A, Quinlivan DS, Glozier N, Lloyd AR, Hickie IB, Guastella AJ. Cytokine aberrations in autism spectrum disorder: a systematic review and meta-analysis. Mol Psychiatry. 2015;20:440–6.
19. Ashwood P, Krakowiak P, Hertz-Picciotto I, Hansen R, Pessah I, Van de Water J. Elevated plasma cytokines in autism spectrum disorders provide evidence of immune dysfunction and are associated with impaired behavioral outcome. Brain Behav Immun. 2011;25:40–5.
20. Lord C, Rutter, M, DiLavore, P, Risi, S, Gotham, K, Bishop, SL. Autism Diagnostic Observation Schedule. 2nd edn (ADOS-2) Manual (Part 1): Modules I-4. Torrance, CA: Western Psychological Services; 2012.
21. Harris PA, Taylor R, Minor BL, Elliott V, Fernandez M, O’Neal L, et al. The REDCap consortium: Building an international community of software platform partners. J Biomed Inf. 2019;95:103208.
22. Harris PA, Taylor R, Thielke R, Payne J, Gonzalez N, Conde JG. Research electronic data capture (REDCap)—a metadata-driven methodology and workflow process for providing translational research informatics support. J Biomed Inf. 2009;42:377–81.
23. Hus Bal V, Lord C. Replication of Standardized ADOS Domain Scores in the simons simplex collection. Autism. 2015;18:583–92.
24. Gotham K, Pickles A, Lord C. Standardizing ADOS scores for a measure of severity in autism spectrum disorders. J Autism Dev Disord. 2009;39:693–705.
25. Hus V, Gotham K, Lord C. Standardizing ADOS domain scores: separating severity of social affect and restricted and repetitive behaviors. J Autism Dev Disord. 2014;44:2400–12.
26. Patel S, Masi A, Dale RC, Whitehouse AJO, Pokorski I, Alvesa GA, et al. Social impairments in autism spectrum disorder are related to maternal immune history profile. Mol Psychiatry. 2018;23:1794–7.
27. Campbell SA, Needham BD, Meyer CR, Tan J, Conrad M, Preston GM, et al. Safety and target engagement of an oral small molecule sequestrant in adolescents with autism spectrum disorder: an open-label phase 1b/2a trial. Nat Med. 2022;28:528–534.
28. DeMayo MM, Young LJ, Hickie IB, Song YJ, Guastella AJ. Circuits for social learning: a unified model and application to Autism Spectrum Disorder. Neurosci Biobehav Rev. 2019;107:388–98.
29. Al-Ayadhi LY, Mostafa GA. Elevated serum levels of interleukin-17A in children with autism. J Neuroinflammation. 2012;9:158.
30. Theoharides TC, Tsilioni I, Patel AB, Doyle R. Atopic diseases and inflammation of the brain in the pathogenesis of autism spectrum disorders. Transl Psychiatry. 2016;6:e844.
31. Gunderson J, Worthley E, Grzadzinski R, Burrows C, Estes A, Zwingenberg L, et al. Social and non-social sensory responsiveness in toddlers at high-risk for autism spectrum disorder. Autism Res. 2021;14:2143–55.
32. Haggarty CJ, Moore DJ, Trottier PD, Hagan R, McGlone FP, Walker SC. Vicarious ratings of social touch the effect of age and autistic traits. Sci Rep. 2021;11:19336.
ACKNOWLEDGEMENTS

We would like to acknowledge and thank the families and children that participated, and Makana Hilton who provided administrative assistance in the development of this manuscript. We also acknowledge the staff at the Clinic for Autism and Neurodevelopment (CAN) Research, University of Sydney, and the Child Development Unit (CDU) at Westmead Children’s Hospital, who supported data collection for this study.

AUTHOR CONTRIBUTIONS

CJ, KB and AG designed the study. CJ, KB, NS and AG collected data. CJ, KB and AG analysed data. CJ wrote the first draft and authors KB, NS and AG commented on drafts of the manuscript. All authors approved final drafts of the manuscript.

COMPETING INTERESTS

AG and NS declare fee for service support of pharmaceutical clinical trials for autism spectrum disorder in their clinic services (including a current and past trial with Axial Therapeutics of a gut-based therapy). These authors receive no personal financial gain from this contracted work. CJ and KB report no conflict of interests.

ADDITIONAL INFORMATION

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41398-022-02185-5.

Correspondence and requests for materials should be addressed to A. J. Guastella.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.