Archival Report

Aberrant Emotional Prosody Circuitry Predicts Social Communication Impairments in Children With Autism

Simon Leipold, Daniel A. Abrams, Shelby Karraker, Jennifer M. Phillips, and Vinod Menon

ABSTRACT

BACKGROUND: Emotional prosody provides acoustical cues that reflect a communication partner’s emotional state and is crucial for successful social interactions. Many children with autism have deficits in recognizing emotions from voices; however, the neural basis for these impairments is unknown. We examined brain circuit features underlying emotional prosody processing deficits and their relationship to clinical symptoms of autism.

METHODS: We used an event-related functional magnetic resonance imaging task to measure neural activity and connectivity during processing of sad and happy emotional prosody and neutral speech in 22 children with autism and 21 matched control children (7–12 years old). We employed functional connectivity analyses to test competing theoretical accounts that attribute emotional prosody impairments to either sensory processing deficits in auditory cortex or theory of mind deficits instantiated in the temporoparietal junction (TPJ).

RESULTS: Children with autism showed specific behavioral impairments for recognizing emotions from voices. They also showed aberrant functional connectivity between voice-sensitive auditory cortex and the bilateral TPJ during emotional prosody processing. Neural activity in the bilateral TPJ during processing of both sad and happy emotional prosody stimuli was associated with social communication impairments in children with autism. In contrast, activity and decoding of emotional prosody in auditory cortex was comparable between autism and control groups and did not predict social communication impairments.

CONCLUSIONS: Our findings support a social-cognitive deficit model of autism by identifying a role for TPJ dysfunction during emotional prosody processing. Our study underscores the importance of tuning in to vocal-emotional cues for building social connections in children with autism.

https://doi.org/10.1016/j.bpsc.2022.09.016

Understanding the emotional state of a communication partner is a crucial component of meaningful human interactions (1–3). The emotional state of a speaker is provided by emotional prosody and is expressed with distinct patterns in a speaker’s tone of voice, intonation, emphasis, rhythm, and speech rate (4–6). Learning to identify these acoustical features and map them to a speaker’s mental state is an important aspect of social development and neurodevelopment in children (7). Children with autism spectrum disorder (ASD) have pronounced deficits in social interaction and communication (8). Difficulties in the perception and production of prosody have been identified as a prominent component of the autism behavioral phenotype since Kanner’s initial description of childhood autism (9–13), and subsequent studies have identified deficits in interpreting emotional prosody cues in ASD (14–24). However, it is unclear why individuals with ASD may have deficits in emotional prosody processing. This question is critical, as emotional prosody represents a unique interface between language, emotion, and social function, and understanding why children struggle with prosody perception may provide novel insights into improving verbal communication in affected individuals.

Two hypotheses have emerged to explain deficits in interpreting vocal-emotional cues in ASD. The first hypothesis proposes that children with ASD have a sensory processing deficit in which abnormal sensory processing of emotional prosody cues precludes access to downstream systems that evaluate and categorize these acoustical features (25,26). Sensory processing of prosodic stimuli is thought to be performed by auditory cortical brain regions in superior temporal cortex in both adults (5,27) and children (28). Thus, the sensory deficit model predicts aberrant brain function at the level of auditory cortical regions that subserve acoustical analyses of vocal-emotional cues (29,30). A second hypothesis posits that acoustical processing of emotional prosody cues is largely intact in individuals with ASD; however, impairments in social cognitive and evaluative processes preclude the accurate interpretation of emotional information from these vocal cues. For example, one prominent theory proposes that the core social deficits of ASD are caused by a deficit in theory of mind, the ability to reason about other people’s mental states and
emotions (31–33). A key brain region that has consistently been implicated in the context of theory of mind processing is the bilateral temporoparietal junction (TPJ) (34–38), a central hub of the social brain network (39). Thus, the social cognition model would predict that aberrant TPJ function contributes to deficits in inferring mental states based on vocal-emotional cues (40,41). A significant gap in our knowledge of social communication deficits in children with ASD is that most studies have focused on visual stimuli (42,43), with relatively few studies conducted in the auditory modality, particularly in children. Understanding the relative contribution of auditory sensory and social cognitive systems to emotional prosody perception deficits in children with ASD would provide important information regarding the nature of this communication impairment and may provide insight into its remediation (44,45).

Little is known regarding the brain basis of emotional prosody processing deficits in children with ASD. Examining this question in children is particularly important given that remediation efforts are generally more beneficial in children compared with adults (46). Previous studies in adults with ASD have failed to identify significant group differences in neural measures during emotional prosody processing when contrasted with neutral speech (47–49). In the only controlled study of emotional prosody in children with ASD, results showed that angry speech compared with neutral speech elicited greater activation in a distributed brain network in children with ASD than in typically developing (TD) children (50). However, this sample of children with ASD had lower cognitive and language abilities than TD children (50), and therefore it is not clear whether the observed neural effects reflect deficits in emotional prosody perception or differences in general cognitive abilities. Thus, it remains unknown whether children with ASD have differential brain activation, connectivity, and decoding of emotional prosody stimuli compared with TD children.

Theoretical models have proposed a link between emotional prosody recognition and social abilities (2,3). However, it is unknown whether neural processing of emotional prosody in children with ASD is related to individual differences in clinical measures of social communication, a key diagnostic symptom domain of ASD. Identifying sources of heterogeneity in key symptom domains remains an important challenge for autism research that has not been addressed in the emotional prosody literature.

To address these questions, we used event-related functional magnetic resonance imaging (fMRI) to measure neural responses in children with ASD and closely matched TD children 7 to 12 years of age while they listened to emotional prosody and neutral speech (Figure 1). We had 3 major goals. First, we investigated whether neural responses and neural decoding of emotional prosody decoding are altered in children with ASD in brain regions serving auditory function, including voice-sensitive auditory cortex. Second, we investigated task-related functional connectivity during emotional prosody processing to examine the integrity of brain circuitry linking voice-sensitive auditory regions with social brain regions, including the TPJ. Third, we assessed whether heterogeneity in neural activation measured during emotional prosody processing is related to individual differences in key clinical domains of social communication and interaction impairments.

METHODS AND MATERIALS

Participants

The Stanford University Institutional Review Board approved the study protocol. Parental consent and children’s assent were obtained for all evaluation procedures, and participants were paid for their participation in the study. Our sample included 43 children between 7 and 12 years of age (22 children with ASD, 21 TD children). This age range is the youngest that can be practically included in multirun task-based fMRI studies in ASD. Children in the ASD group were diagnosed based on a classification algorithm (51) that combines information from both module 3 of the Autism Diagnostic Observation Schedule, Second Edition (ADOS-II) (52) and the Autism Diagnostic Interview–Revised (ADI-R) (53). Children with ASD and TD children underwent extensive neuropsychological testing to assess their cognitive abilities, language function, and emotion recognition abilities. Groups were matched for age, sex, cognitive and language abilities, and motion during functional imaging. Demographic and neuropsychological characteristics are listed in Table 1. Details are provided in Supplemental Methods.

fMRI Stimuli

The stimuli presented during fMRI scanning consisted of acoustic sentences spoken in emotional and neutral prosody as well as nonspeech environmental sounds. Emotional and neutral prosodic stimuli were recorded in a recording studio by a professional actress who produced vocal samples of 2 sentences, “A bag is in the room.” (sentence 1) and “My spoon is on the table.” (sentence 2), using sad, happy, and neutral emotions (Figure 1B). Emotional and neutral prosody stimuli for the fMRI experiment were selected based on results from a behavioral experiment conducted in an independent cohort of 27 school-age TD children who did not participate in the fMRI study and 9 adults (Figure 1C and Supplemental Methods). Stimuli can be downloaded from the Open Science Framework (https://dx.doi.org/10.17605/OSF.IO/TYFXS).

fMRI Task

Stimuli were presented in 10 separate fMRI runs, each lasting approximately 3.5 minutes. One run consisted of 39 trials of acoustic sentence stimuli spoken in sad (high and low intensity), happy (high and low intensity), and neutral prosody as well as environmental sounds and catch trials. Across the 10 runs, a total of 60 exemplars per stimulus condition were presented to each participant. Acoustic stimuli were presented during silent intervals between volume acquisitions to eliminate the effects of scanner noise on auditory perception (Figure 1D). Functional images were subjected to preprocessing procedures, which included realignment, slice timing correction, spatial normalization, and smoothing. Details are provided in Supplemental Methods.
**Figure 1.** Overview of participants, stimuli, experimental design, analysis strategy, and behavioral data. (A) Our sample included children with autism spectrum disorder (ASD) and typically developing (TD) children between 7 and 12 years of age. (B) Functional magnetic resonance imaging (fMRI) stimuli consisted of acoustic sentences spoken in emotional and neutral prosody. Spectrograms of sentence 2, “My spoon is on the table,” spoken in sad prosody (left panel), in neutral prosody (middle panel), and in happy prosody (right panel). (C) Stimuli were selected based on results from a behavioral experiment conducted in an independent cohort of 27 school-age TD children and 9 adults, who provided ratings on a 5-point scale (“How sad or happy is this voice?”). Stimuli that were consistently rated 1 and 5 were identified as the high-intensity sad and happy stimuli for the fMRI experiment; stimuli rated 2 and 4 were identified as the low-intensity sad and happy stimuli, and the stimulus consistently rated 3 was identified as the neutral control stimulus. (D) A sparse sampling fMRI scanning protocol with a repetition time (TR) larger than the acquisition time (TA) was used to present acoustic stimuli during silent intervals between volume acquisitions to eliminate the effects of scanner noise on auditory perception. (E) Schematic of the analyses employed in the study. (i) Neural decoding of emotional prosody within auditory cortical regions of interest, (ii) neural connectivity during emotional prosody, and (iii) neural activation during emotional prosody were compared between children with ASD and TD children. (iv) Relationships between neural measures and clinical measures of social function were assessed in the group of children with ASD. (F) ASD (green) and TD (orange) groups were closely matched regarding measures of intelligence, as quantified by the Wechsler Abbreviated Scale of Intelligence (WASI), language abilities measured with the Wechsler Individual Achievement Test, Second Edition (WIAT-II), and scanner motion. ADI-R, Autism Diagnostic Interview–Revised; aSTS, anterior superior temporal sulcus; FIQ, Full Scale IQ; HG, Heschl's gyrus; mSTS, middle STS; PIQ, performance IQ; PP, planum polare; pSTS, posterior STS; PT, planum temporale; VIQ, verbal IQ.
The goal of voxelwise analysis of fMRI activation was to identify brain regions that showed differential activity levels in children with ASD and TD children in response to emotional prosody stimuli, neutral prosody, and environmental sounds. A second-level analysis used a two-sample t test to compare activation in children with ASD and TD children on the contrasts of interest [neutral prosody > environmental sounds], [sad prosody > neutral speech], and [happy prosody > neutral speech]. Details are provided in Supplemental Methods.
Association Between Neural and Clinical Measures of Autism Symptom Severity

Whole-brain regression analysis was used to examine the relationship between neural activation to emotional prosody and social communication impairments in children with ASD, as quantified using the ADI-R communication and language subscale (53). This ADI-R subscale was chosen based on prior evidence showing associations between social information processing and social communication deficits in adults with ASD (40). Details are provided in Supplemental Methods.

RESULTS

Emotional Face and Prosody Recognition Accuracy in Children With ASD and TD Children

We first assessed behavioral emotion recognition abilities in children with ASD and TD children using vocal and facial emotional cues. Emotion recognition was assessed using the Diagnostic Analysis of Nonverbal Accuracy 2 (56,57) (Supplemental Methods). Children with ASD showed lower accuracy compared with TD children in recognizing emotions across voices and faces ($F_{1,38} = 5.12, \ p = .03$) (Figure 2). Follow-up analyses revealed that children with ASD had lower accuracies than TD children in recognizing emotions from voices ($t_{26.75} = 2.35, \ p = .03$), but not faces ($t_{32.6} = -1.39, \ p = .17$). The group difference in emotion recognition from voices had a moderate to large effect size (Hedges’ $g = 0.74$). These results suggest selective deficits for recognizing emotions from vocal cues in children with ASD.

Neural Activation During Emotional Prosody in Auditory Cortex

We next assessed neural activation during emotional prosody processing in children with ASD and TD children within distinct subregions of auditory cortex (28). Results showed no group differences in auditory cortical activation to emotional prosody using either frequentist (all $p > .05$) or Bayesian statistical analyses (Figure S1B and Supplemental Results). See Supplemental Results for additional analysis of group differences in activation at the whole-brain level.

Multivariate Decoding of Emotional Prosody in Auditory Cortex

As multivariate decoding provides a more sensitive measure of differential response to stimuli (58), we assessed whether decoding of emotional prosody differed between children with ASD and TD children within distinct subregions of auditory cortex (28,59). Results from ROI-based multivariate pattern analysis showed striking similarity of emotional prosody decoding for children with ASD and TD children across all investigated auditory cortical ROIs as revealed by both frequentist (all $p > .05$) and Bayesian statistical analyses (Figure S1C and Supplemental Results). Notably, children with ASD showed the same pattern of emotional prosody decoding in auditory cortex as previously reported in TD children (28) with middle STS and posterior STS showing the highest classification accuracies for differentiating emotional prosody from neutral speech (Figure S1C). Across both groups, all auditory ROIs decoded emotional prosody from neutral speech above chance level for both sad and happy emotions (all $p < 0.05$).

Functional Connectivity of Voice-Sensitive Auditory Cortex During Emotional Prosody Processing

We next examined the integrity of voice-sensitive functional circuitry in children with ASD. We used whole-brain generalized psychophysiological interaction analysis (55) to examine group differences between TD children and children with ASD with regard to functional connectivity of voice-sensitive auditory cortex during the processing of emotional prosody stimuli.

Figure 2. Reduced emotion recognition accuracy in children with autism spectrum disorder (ASD). Selective impairments in recognizing emotion for voices on the Diagnostic Analysis of Nonverbal Accuracy 2 (DANVA2), a standardized emotion recognition test. * $p < .05$. ns, not significant; TD, typically developing.
For the [sad prosody > neutral speech] contrast, group comparisons of functional connectivity revealed a pronounced pattern of hyperconnectivity in children with ASD. Specifically, results revealed greater task-based functional connectivity between voice-sensitive cortex and the bilateral TPJ in children with ASD compared with TD children [left TPJ: 1520 mm$^3$, familywise error-corrected $p_{FWE} < .01$; right TPJ: 1320 mm$^3$, $p_{FWE} < .01$] (see Table S1 for Montreal Neurological Institute coordinates of all significant clusters). Children with ASD further showed hyperconnectivity between voice-sensitive cortex and right fusiform gyrus (944 mm$^3$, $p_{FWE} < .01$), primary visual cortex (800 mm$^3$, $p_{FWE} < .01$), and left middle temporal gyrus (792 mm$^3$, $p_{FWE} < .01$). For the [happy prosody > neutral speech] contrast, children with ASD again showed hyperconnectivity between voice-sensitive cortex and left middle temporal gyrus compared to TD children (1216 mm$^3$, $p_{FWE} < .01$) (Figure 3). Across both sad and happy prosody, no brain areas showed reduced emotional prosody-related connectivity with voice-sensitive cortex in children with ASD compared with TD children. These results demonstrated hyperconnectivity of voice-sensitive auditory cortex during sad prosody processing in children with ASD.

**Figure 3.** Functional connectivity of voice-sensitive auditory cortex. Functional connectivity during emotional prosody processing in children with autism spectrum disorder (ASD) and typically developing (TD) children was examined using a generalized psychophysiological interaction (gPPI) model. (A) The gPPI analysis was seeded in bilateral voice-sensitive auditory cortex. (B) gPPI connectivity was calculated from voice-sensitive auditory cortex to every gray matter voxel in the brain. (C) Group comparisons of emotional prosody-related connectivity revealed hyperconnectivity of voice-sensitive cortex to the bilateral temporoparietal junction (TPJ) in children with ASD compared with TD children during sad prosody. For sad prosody, voice-sensitive cortex was also functionally hyperconnected with left middle temporal gyrus (MTG), primary visual cortex (V1), and right fusiform gyrus in children with ASD. For happy prosody, voice-sensitive cortex was functionally hyperconnected with left MTG in children with ASD. ***$p < .001$. l, left; r, right.
Relation between TPJ activation and social communication impairments

Our final goal was to determine whether neural measures of emotional prosody processing are associated with clinical measures of social communication impairments in children with ASD. We performed a whole-brain linear regression on neural activation using participants’ social communication impairment scores as a regressor (Figure 4). For both [sad prosody > neutral speech] and [happy prosody > neutral speech] contrasts, results from this analysis showed a striking pattern of brain-behavior relationships: children with ASD with more severe social communication impairments, reflected by greater scores on the ADI-R, showed reduced activation during sad prosody in the bilateral TPJ (left TPJ: 1432 mm³, \( P_{\text{FWE}} < .01 \); right TPJ: 1128 mm³, \( P_{\text{FWE}} < .01 \)). Additional clusters where activation was negatively associated with social communication impairments were found in the right lateral occipital cortex (3280 mm³, \( P_{\text{FWE}} < .01 \)) and the prefrontal cortex (2688 mm³, \( P_{\text{FWE}} < .01 \)). Similarly, during processing of happy prosody, children with more severe social communication impairments showed reduced activation in the bilateral TPJ (left TPJ: 5168 mm³, \( P_{\text{FWE}} < .01 \); right TPJ: 12,896 mm³, \( P_{\text{FWE}} < .01 \)). Additional clusters were widely distributed over the cortex and subcortical areas (see Table S3 for a full list and Montreal Neurological Institute coordinates of significant clusters).

DISCUSSION
Understanding a speaker’s emotional state is critical for navigating the social world, and many children with ASD have deficits in interpreting the vocal-acoustical cues that signal a speaker’s emotions, known as emotional prosody. In contrast to studies using emotional faces, investigations of the neural processing of emotional prosody in individuals with ASD, particularly in children, have been few and limited in scope (50). Results from the current study revealed behavioral deficits for recognizing vocal emotions in children with ASD compared with TD children who were matched for sex, age, cognitive abilities, and language function. Behavioral deficits in recognizing emotions from voices in children with ASD were accompanied by aberrant functional coupling between voice-sensitive auditory areas and the bilateral TPJ, a brain region associated with theory of mind (34) and creation of a social context for behavior (60). Crucially, neural activation during emotional prosody processing in the bilateral TPJ predicted the severity of social communication impairments in children with ASD, with reduced brain activity in more severely affected children. In contrast, children with ASD showed accurate neural decoding of emotional prosody stimuli across auditory cortex subdivisions, and decoding accuracies were comparable to decoding accuracies measured in TD children (28). Converging results from brain connectivity and social communication brain-behavior analyses support the hypothesis that TPJ dysfunction in ASD not only is associated with...
deficits in explicit mentalizing and social cognitive tasks, most of which have involved visual stimuli, but also extends to fine-grained analysis of the acoustical features that signal the emotional state of a speaker.

**Aberrant Functional Coupling Between Voice-Sensitive Auditory Cortex and TPJ During Emotional Prosody Processing in Children With ASD**

Results from whole-brain functional circuit analyses revealed hyperconnectivity of voice-sensitive auditory cortex with the bilateral TPJ in children with ASD while processing sad prosody compared with neutral vocal stimuli. This result has several important implications for understanding the neurobiological basis of social information processing in ASD. First, previous studies in individuals with ASD that have highlighted TPJ dysfunction have focused primarily on visual processing tasks that examine complex aspects of social cognition (40,61,62). These previous findings suggest that TPJ dysfunction in ASD is associated with active inferential cognitive processes associated with explicit theory of mind tasks. Results from the current study suggest that TPJ circuit dysfunction extends beyond these explicit mentalizing tasks and impacts the passive processing of sensory-level cues that signal mental states in a communication partner. Therefore, our findings suggest a broader role for TPJ circuitry in social impairments in ASD compared with previous studies.

Our findings also provide important new information regarding voice processing circuitry in children with ASD more generally. An influential model of voice processing circuitry in the brain posits that the STS serves as a hub for the voice processing network (63), and empirical evidence has supported many of the predictions of this model in TD individuals (54,64–66). Importantly, investigations into the brain circuitry underlying voice processing in children with ASD have shown that aberrant STS connectivity with other key brain systems appears to play a central role in voice-related processing deficits in these individuals (67,68). Results from the current study add to this literature by showing that aberrant functional connectivity, specifically hyperconnectivity, of voice-sensitive STS and the TPJ, a core node of the social brain, is a neural signature underlying impaired processing of vocal emotions in children with ASD. Findings support a model of voice processing in ASD that places dysfunction of voice-sensitive STS circuitry at the center of multiple aspects of voice perception deficits in affected individuals.

**TPJ Activation Is Related to Social Communication Symptom Severity in Children With ASD**

Our whole-brain voxelwise regression analysis revealed a striking association between activation within the bilateral TPJ during emotional prosody perception and symptom severity for social communication and social interaction impairments in children with ASD. Specifically, reduced activation in the TPJ during emotional prosody processing was correlated with more severe social communication deficits in these children. Results are consistent with previous studies in adults (40) and adolescents (41) with ASD that revealed a similar relationship between reduced activation of the TPJ during social cognitive theory of mind tasks and symptom severity for social communication deficits. Findings from the current study add to this literature by showing that, in contrast to the active inferential processes highlighted in previous studies (40,41), the link between TPJ activity and severity of autism clinical symptoms extends to social stimuli presented through the auditory modality and passive listening tasks in children with ASD. Importantly, results showing a relationship between bilateral TPJ activation and symptom severity for social communication provide complementary evidence for TPJ dysfunction in children with ASD relative to our finding of group-level STS-TPJ hyperconnectivity during emotional prosody processing (Figure 3). Together, these results provide converging evidence that upregulation of TPJ activity and normalization of TPJ functional circuitry may serve as neurobiological targets for remediation of social communication deficits in individuals with ASD (69).

**Implications for Sensory and Social Cognitive Theories of ASD**

Processing of the subtle acoustical cues in speech that signal a speaker’s emotions may be viewed as a crucial nexus of sensory and social cognitive processing that requires the mapping of brief and fine-grained pitch and timbral features onto representations of distinct emotional state information that is essential for successful and meaningful human interaction. Importantly, autism is closely associated with both auditory sensory and social cognitive deficits, and therefore the current study provides an opportunity to adjudicate models of autism that support primary contributions of sensory and social cognitive deficits in ASD. Results from the current study showing aberrant functional coupling and activity patterns in the TPJ during emotional prosody processing are consistent with social cognitive accounts of ASD, which have consistently implicated the TPJ as a key brain region underlying deficits for theory of mind–related processes (40,42,61). Importantly, results from our decoding and activation analyses showed strikingly similar sensory neural function in ASD and TD groups across a wide range of auditory cortical regions in superior temporal cortex during emotional prosody perception. Our findings are consistent with prior evidence examining auditory social information processing in children and adolescents with ASD (50,67) and in adults with ASD (47,48), which identified only subtle differences, or no differences at all, in neural function at the level of auditory cortex when compared with TD individuals. Our findings do not support a sensory deficit model of autism (25,26) in the context of emotional prosody processing and suggest that auditory sensory processing of vocal-emotional cues is not a primary deficit associated with impaired emotional prosody perception. Rather, findings suggest a crucial role for the TPJ in social cognitive impairments in ASD that encompass sensory elements that contribute to social communication and cognition.

**Task-Based Neuroimaging in Clinical Pediatric Samples**

Reproducibility in neuroimaging research represents a challenge for the study of clinical pediatric populations, in whom
data are more difficult to acquire compared with adult participants (70). While the sample size used in the current study (N = 43) is modest compared with recent task-based brain imaging studies of adult populations, an important consideration is that deep sampling in individual participants improves the precision of the measurements and consequently increases replicability (71). Therefore, a primary goal for our data acquisition was to obtain a large amount (7–10 runs; 60 stimuli/condition) of high-quality brain imaging data per participant to enable robust within-participant sampling.

To keep fMRI runs short and maximize participant engagement, we employed a simple implicit fMRI task with two basic emotions. However, an important future direction for this work is to use prosodic processing tasks with greater complexity to investigate different emotions and processing modes in children with ASD (49).

Conclusions

We identified neural alterations underlying impaired emotional prosody processing in school-age children with ASD, which were predictive of pervasive social communication and interaction difficulties. Children with ASD showed atypical modulation of functional connectivity between voice-sensitive auditory cortex and the bilateral TPJ, which represents a hub of the social brain network. Reduced activation of the bilateral TPJ during emotional prosody processing was associated with greater social communication and interaction impairments. These findings support social-cognitive accounts of social deficits in ASD and highlight the importance of tuning in to vocal-emotional cues for building social connections in children with autism.

ACKNOWLEDGMENTS AND DISCLOSURES

This work was supported by the National Institutes of Health (Grant No. KU1MH102428 [to DAA]), Grant Nos. R21DC017950 and R21DC017950-S1 [to DAA and VM], and Grant No. R01MH084164 [to VMJ]. NARSAD Young Investigator Award from the Brain and Behavior Research Foundation (to DAA). Singer Foundation, Simons Foundation Autism Research Initiative (Grant No. 308939 [to VM]), and Swiss National Science Foundation (Grant No. PZHP1_187704 [to SL]).

We thank all the children and their parents who participated in our study and the staff at the Stanford Lucas Center for Imaging for assistance with data collection. We thank Davlat El-Said and Carlo de los Angeles for help with stimulus production. The authors report no biomedical financial interests or potential conflicts of interest.

ARTICLE INFORMATION

From the Department of Psychiatry and Behavioral Sciences, Stanford University, Stanford, California (SL, DAA, SK, JMP, VM); Department of Neurology and Neurological Sciences, Stanford University, Stanford, California (VM); and Stanford Neurosciences Institute, Stanford University, Stanford, California (VM).

SL and DAA contributed equally to this work as joint first authors.

Address correspondence to Simon Leipold, Ph.D., at simon.leipold@donders.ru.nl, or Vinod Menon, Ph.D., at menon@stanford.edu.

Received Apr 13, 2022; revised Aug 25, 2022; accepted Sep 30, 2022.

Supplementary material cited in this article is available online at https://doi.org/10.1016/j.bpsc.2022.09.016.

REFERENCES


