A failure of left temporal cortex to specialize for language is an early emerging and fundamental property of autism

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Failure to develop normal language comprehension is an early warning sign of autism, but the neural mechanisms underlying this signature deficit are unknown. This is because of an almost complete absence of functional studies of the autistic brain during early development. Using functional magnetic resonance imaging, we previously observed a trend for abnormally lateralized temporal responses to language (i.e. greater activation on the right, rather than the expected left) in a small sample (n = 12) of sleeping 2–3 year olds with autism in contrast to typically developing children, a finding also reported in autistic adults and adolescents. It was unclear, however, if findings of atypical laterality would be observed in a larger sample, and at even earlier ages in autism, such as around the first birthday. Answers to these questions would provide the foundation for understanding how neurofunctional defects of autism unfold, and provide a foundation for studies using patterns of brain activation as a functional early biomarker of autism. To begin to examine these issues, a prospective, cross-sectional design was used in which brain activity was measured in a large sample of toddlers (n = 80) during the presentation of a bedtime story during natural sleep. Forty toddlers with autism spectrum disorder and 40 typically developing toddlers ranging in age between 12–48 months participated. Any toddler with autism who participated in the imaging experiment prior to final diagnosis was tracked and diagnoses confirmed at a later age. Results indicated that at-risk toddlers later diagnosed as autistic display deficient left hemisphere response to speech sounds and have abnormally right-lateralized temporal cortex response to language; this defect worsens with age, becoming most severe in autistic 3- and 4-year-olds. Typically developing children show opposite developmental trends with a tendency towards greater temporal cortex response with increasing age and maintenance of left-lateralized activation with age. We have now demonstrated lateralized abnormalities of temporal cortex processing of language in autism across two separate samples, including a large sample of young infants who later are diagnosed with autism, suggesting that this pattern may reflect a fundamental early neural developmental pathology in autism.

Keywords: autism; language; functional magnetic resonance imaging; sleep; temporal cortex
Abbreviation: ADOS = autism diagnostic observation schedule; ASD = autism spectrum disorder
Introduction

Early subclinical behavioural signs of autism appear in the first years of life (Palomo et al., 2006; Zwaigenbaum et al., 2009; Pierce et al., 2011) but the underlying brain dysfunction remains a mystery. This is because of a virtual absence of functional imaging studies of the autistic brain during this crucial developmental period.

One of the earliest appearing signs of autism is a failure to develop sophisticated expressive and receptive language. While most typical toddlers begin to say words around their first birthday (Fenson et al., 1994), some toddlers who will eventually develop autism miss or are slow to achieve this important milestone. In fact, delay in language is reported as the strongest and earliest red flag by most experts (De Giacomo and Fombonne, 1998; Wetherby et al., 2004). Later in development, children with autism often, but not always, display unusual patterns of language that may include the presence of neologisms (i.e. made up words (Volden and Lord, 1991)), speech that is highly repetitive, ritualized or echolalic (McEvoy et al., 1988) or characterized by high rates of pronoun reversals (Rapin and Dunn, 2003). Early studies on the topic found that comprehension of speech was more severely compromised in children with autism than those with global developmental delay (Bartak et al., 1975). Studies examining samples of individuals from the 1980s reported that ~50% of the children with autism spectrum disorder have little to no functional language (Sigman and McGovern, 2005), although more recent studies suggest better language outcomes with at least 75% of the individuals acquiring some functional language (Luyster et al., 2008). Non-verbal social communication engagement, including the use of gestures such as pointing and waving, are reduced in autism (Loveland et al., 1988). Not surprisingly, the pragmatic use of language (e.g. conversational speech) is severely impaired in affected children (Loveland et al., 1988; Philofsky et al., 2007). Overall, brain systems that support language development must be abnormal in autism. Moreover, given the early emergence of language capacity in humans, with typical infants demonstrating the ability to discriminate phonemes within the first weeks of life (Eimas et al., 1971; Werker and Lalonde, 1988), recognize their own name at 4 months (Mandel et al., 1995) and discriminate familiar from unfamiliar words by 7 months (Tincoff and Jusczyk, 1999), this neural abnormality must begin very early in development in babies that will eventually show autism.

Integrity of temporal and frontal cortices is essential for normal language development, but in autistic infants, toddlers and young children these structures display pathological overgrowth (Carper et al., 2002; Carper and Courchesne, 2005; Schumann et al., 2010; Hazlett et al., 2011). For example, Carper and Courchesne (2002) showed that frontal and temporal cortices were ~13% greater in volume in children with autism spectrum disorder relative to typically developing children, whereas other brain regions, such as occipital cortex, were not significantly enlarged. When brain systems directly engaged in language comprehension were challenged in studies of older autistic children, adolescents and adults, abnormal functional lateralization in frontal and temporal cortices has consistently been observed, including increased right hemisphere (Boddaert et al., 2004; Wang et al., 2006; Kleinhans et al., 2008; Mason et al., 2008; Tesink et al., 2009; Knaus et al., 2010) and decreased left hemisphere (Just et al., 2004; Gendry Meresse et al., 2005; Harris et al., 2006; Gaffrey et al., 2007; Groen et al., 2009) responsiveness. When differences in brain response between hemispheres are measured directly, individuals with autism generally display less functional lateralization (Muller et al., 1999; Kleinhans et al., 2008; Minagawa-Kawai et al., 2009; Anderson et al., 2010), or a tendency towards abnormal right hemisphere lateralization (Dawson et al., 1989; Bruneau et al., 1999; Flagg et al., 2005). Whether or not such abnormal response and rightward functional lateralization occurs at the beginning stages of autism has never been tested. This is largely due to substantial practical and technical difficulties in measuring brain function in infants and toddlers with disabilities using functional MRI, a technique that traditionally requires subjects to remain still for extended periods of time. Fortunately, several recent studies have demonstrated that it is possible to obtain functional MRI data in response to language sounds from infants during natural sleep, thus eliminating the requirement for infants and toddlers to hold still (for review, see Pierce, 2011). Moreover, the ‘sleep functional MRI’ method has demonstrated responsiveness of superior temporal gyri to auditory stimuli in non-autistic infants as young as 1 week to 3 months old (Anderson et al., 2001; Dehaene-Lambertz et al., 2002, 2006). More recently, our laboratory examined brain response to language stimuli during natural sleep in two age groups of typically developing children (mean ages of 21 and 39 months; Redcay et al., 2008). Both age groups showed superior temporal gyri response to language stimuli, and the younger group’s response included additional frontal, occipital and cerebellar sites of activation, demonstrating that expected cortical responses during language comprehension can be measured in very young typically developing children using functional MRI during natural sleep.

Sleep functional MRI provides many scientific as well as safety advantages for examining brain function in very young autistic and typically developing subjects including the ability to test subjects from all functioning levels. Despite its strengths, only one study has examined differences between very young children with autism and typically developing children using this technology. In response to a bedtime story during natural sleep, Redcay and Courchesne (2008) found reduced functional activity in temporal cortices in a small sample (n = 12) of 2–3 year olds children with autism compared with both chronological and mental age-matched groups and a non-statistically significant, but interesting trend towards abnormal rightward laterality of brain response to language stimuli in the autism group.

There are several natural questions that follow: can abnormal functional activation in response to language be detected even at the first signs of autism? Is abnormally right lateralized brain response present at that at-risk time-point or does it emerge gradually with age after full symptom onset? What is the developmental trajectory of normal left lateralization for language in the first several years of life in autism when the explosion in language abilities normally takes place in typical infants? Given the relatively
late age of diagnosis, which remains around the age of 4 years nationally (Network, 2009), it has historically been challenging to identify and study toddlers with autism spectrum disorder prospectively before the onset of full-blown clinical symptoms. However, new advances in the early detection of autism, such as the 1-Year Well-Baby Check-Up Approach, have made it possible to study autism prospectively from the first year of life in the general population (Pierce et al., 2011).

Here, we report results of the first functional MRI study of autism in the earliest years of life that addresses these foundational questions. We analysed functional MRI activation to language stimuli, a baby’s bedtime story, measured during natural sleep from the largest sample collected to date of infants and toddlers at-risk for autism spectrum disorder whose diagnosis was confirmed at older ages, as well as from 3- to 4-year-old children with autism spectrum disorder. Functional MRI activation patterns were compared with those in typically developing infants, toddlers and very young children. We predicted abnormalities of functional response and lateralization at the earliest ages in autism spectrum disorder, with right instead of left hemisphere specialization, and a failure to develop normal left lateralization patterns across the years from infancy to young childhood.

Subjects and methods

Participants

As part of an ongoing study (www.autism-center.ucsd.edu) toddlers ranging in ages between 12 and 48 months were brought to a scanning facility on one or more nights. Toddlers at-risk for autism spectrum disorder whose diagnosis was confirmed at older ages, as well as from 3- to 4-year-old children with autism spectrum disorder. Functional MRI activation patterns were compared with those in typically developing infants, toddlers and very young children. We predicted abnormalities of functional response and lateralization at the earliest ages in autism spectrum disorder, with right instead of left hemisphere specialization, and a failure to develop normal left lateralization patterns across the years from infancy to young childhood.

A total of 21 participants were younger than 30 months (an age at which autism spectrum disorder diagnoses can be reliably made) at the time of scanning and were judged to be at-risk for autism spectrum disorder at that time. Five of these children were later judged to no longer warrant a spectrum diagnosis and were not included in the analyses [revised diagnoses were: typical (n = 2), language delayed (n = 1), developmental delay/borderline IQ (n = 1) and autistic features of insufficient severity to warrant a diagnosis (n = 1)]. Of the remaining 16 children judged at-risk prior to 30 months of age, all but one (who moved out of the area) have participated in at least one follow-up clinical evaluation 6 months to 1 year later [at a mean (SD) age of 31.9 (5.2) months], and all of these have had their provisional diagnoses confirmed by a psychologist (one currently has autism diagnostic features with diagnosis pending a second follow-up). Total Autism Diagnostic Observation Schedule-Toddler Module (ADOS-T) scores [social and communication plus repetitive scores; mean (SD)] at follow-up were 17.6 (5.0).

All toddlers participated in a series of psychometric tests across multiple 2-h sessions that included the Autism Diagnostic Observation Schedule-Toddler Module newly validated for use with infants as young as 12 months (Luyster et al., 2009), the Mullen Scales of Early Learning (Mullen, 1995) and the McArthur-Bates Communicative Development Inventories (Fenson et al., 1993). Toddlers participated in additional behavioural (e.g. examinations of play behaviour) and biological tests as part of a larger study. All standardized assessments were administered by three highly experienced doctoral-level psychologists with over 15 years combined experience in the field of autism.

Demographic and clinical summary data for the groups at the time of scanning are presented in Table 1. As expected, the typically developing group had significantly higher receptive language and visual T-scores and significantly lower Autism Diagnostic Observation Schedule scores than the autism spectrum disorder group. In addition, the typically developing group was significantly younger than the autism spectrum disorder group (P = 0.005); however, the difference in mean age was only 6 months. This study was approved by the University of California San Diego Institutional Review Board. Parents provided written informed consent according to the Declaration of Helsinki and were paid for their participation.

Stimuli and task design

The speech task was identical to that used in our previous pilot study (Redcay et al., 2008) and consisted of three types of stimuli, presented in 20-s blocks: complex forward speech, simple forward speech and backward speech. All speech conditions were created using the same female speaker. During the complex forward speech condition, toddlers were exposed to segments of a children’s story that was written at a comprehension level of over 48 months. During the simple forward speech condition, toddlers were exposed to excerpts from a child’s story written at a comprehension level between 12 and 36 months. Finally, during the backward speech condition, toddlers were exposed to the simple story segments played backwards. There were also 20-s rest blocks (no presented stimuli) between each stimulus type. Each stimulus type was repeated three times in a pseudorandom order. The total task length was 6 min 25 s. The stimuli were presented using commercially available music presentation software with maximum volume set both for the software and the computer’s speakers. Stimulus presentation was through pneumatic headphones (Confon, Inc.) set to a volume attenuation of −40 dB for all participants.

A total of 111 toddlers attempted participation in the scanning session and a total of 80 infants were included in final analyses (40 autism spectrum disorder and 40 typically developing). Infants and toddlers were included in the final sample if both a structural MRI scan and a functional MRI scan examining speech processing were acquired before the child awoke. A total of 12 participants with autism spectrum disorder and 10 typically developing participants woke during the functional scan before two-thirds of the time-points had been collected, so their data were not included in analyses. One typically developing participant was included who woke after completing more than two-thirds of the scan; all other participants slept through the entire scan. Data from an additional four participants were excluded for other imaging-related reasons [given an over-the-counter medication by the family (n = 1), given earplugs to aid in staying asleep (n = 1), primary language in the home was German (n = 1) and found to have a temporal lobe cyst (n = 1)].
Scanning procedure

Infants and toddlers were imaged in a 1.5 Tesla General Electric MRI scanner during natural sleep; no sedation was used. Parents were encouraged to forgo any usual naps by the child and engage the child in rigorous physical activity during the day. Families arrived at the scanning facility 1 h after the child’s typical bedtime and most children had been asleep in the car for ~15 min prior to arrival. If not already asleep or if awakened by placement on the scanner bed, the child was allowed to fall asleep in the waiting or scanning room and was placed on the scanner bed after ~15 min of sleep. The time that the child fell asleep was recorded for every participant. After placement of the headphones, padding of the head for comfort and motion reduction and covering the child with a weighted blanket for warmth and motion reduction, the scan was started (~15 min later). A research assistant was present in the room next to the child during the entire scan and stopped the scan if the child woke up or made a large movement.

The order of scans varied somewhat between individuals, but all participants first received a high-resolution, T1-weighted anatomical scan (repetition time = 6.5, flip angle = 12°, bandwidth = 31.25, field of view = 24 cm, in-plane resolution = 1 × 1 mm, slice thickness = 1.2 mm, 170 slices, scan length = 7 min 24 s) for localization of functional signals and warping into standard atlas space. On average, the speech task was presented 18.3 ± 13.3 min after the onset of scanning. Other functional and anatomical scans were acquired before and after the speech task; data from these will not be presented in this article. If the child remained asleep for the entire procedure, the total scan time was 1 h 15 min.

Blood oxygenation level-dependent signal was measured across the whole brain with echoplanar imaging during the language paradigm. Scan parameters were: echo time = 30 ms, repetition time = 2500 ms, flip angle = 90°, bandwidth = 70 kHz, field of view = 25.6 cm, in-plane resolution = 4 × 4 mm, slice thickness = 4 mm, 31 slices.

Data analyses

Analysis of Functional Neuroimages software was used to detect and correct for head motion. Head motion in the analysed participants, who slept through the entire scan (or, for one participant, more than two-thirds of the scan), was minimal: no participant was excluded for having visually apparent residual motion following motion correction on more than one-third of time-points. In addition, there were no group differences in the amount of motion as determined by t-tests comparing the average across the scan of the squared movement parameters in each in-plane direction and each axis of rotation (all \( P > 0.19 \)). Analysis of Functional NeuroImages software was also used to carry out regression analyses in which we compared blood oxygenation level-dependent response between speech blocks and periods of no stimulation while controlling for linear trends and residual motion. Specifically, we examined the contrast between blood oxygenation level-dependent response during periods of both complex and simple forward speech and backward speech to the response during periods of non-stimulation (AllSpeech versus Rest). Individual subject activation maps were then spatially blurred with an 8-mm full-width at half-maximum Gaussian kernel and normalized to standard atlas space (Talairach and Tournoux, 1988) by rigid-body transformation based on manual placement of markers at the anterior and posterior commissures and at the extremes of the brain in each plane. We have found in previous investigations (Redcay et al., 2008) that group averaging following Talairach transformation is at least as accurate for toddlers with and without autism spectrum disorder as it is for adults. Group maps of significant language-related response were then created for the typical and autism spectrum disorder groups individually and for the direct comparison between them. For all whole-brain analyses, we retained clusters with an individual-voxel \( P < 0.01 \) (for whole group; \( t > 2.86 \) for within group; \( t > 2.42 \) for a one-tailed between-group comparison) and a volume of at least 32 voxels (2048 μl). This threshold and cluster size combination protects a whole-brain rate of false-positive clusters less than \( P = 0.05 \) as determined by Monte–Carlo simulation. Whole-brain analyses were repeated with age or sleep latency as a covariate in the model using the Analysis of Functional Neurimages program 3dttest +. To examine the generality of the voxel-wise findings based on AllSpeech to each different speech condition, we separately compared the mean response to Simple Forward, Complex Forward and Backward speech between groups in the observed region of group difference. Furthermore, follow-up analyses of the relationship of age to the amplitude and extent of activation within the identified region of differential speech response were conducted. We tested for a group × age interaction and also calculated the within-group age correlations. Correlation analyses were also conducted for the clinical variables within the region of differential response to speech.

We were also interested in examining the laterality of response to speech stimuli within the superior temporal gyrus. We confined our analysis to voxels falling within the boundaries of the superior temporal gyrus as defined by the Talairach Daemon (Lancaster et al.,

### Table 1 Participant characteristics

<table>
<thead>
<tr>
<th>Measure</th>
<th>Autism spectrum disorder</th>
<th>Typically developing</th>
<th>Group comparison</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Mean (SD), range or number (%)</td>
<td>n</td>
</tr>
<tr>
<td>Age (months)</td>
<td>40</td>
<td>32.0 (12.5–47.6)</td>
<td>40</td>
</tr>
<tr>
<td>Sex (no. of males) (%)</td>
<td>40</td>
<td>29 (73)</td>
<td>40</td>
</tr>
<tr>
<td>Receptive language age (months)</td>
<td>37</td>
<td>20.1 (9.8)</td>
<td>36</td>
</tr>
<tr>
<td>Receptive language T-score</td>
<td>37</td>
<td>30.5 (11.9)</td>
<td>36</td>
</tr>
<tr>
<td>Receptive visual age (months)</td>
<td>37</td>
<td>27.1 (10.7)</td>
<td>36</td>
</tr>
<tr>
<td>Receptive visual T-score</td>
<td>37</td>
<td>40.9 (15.8)</td>
<td>36</td>
</tr>
<tr>
<td>ADOS social and communication score</td>
<td>40</td>
<td>14.2 (3.7)</td>
<td>40</td>
</tr>
<tr>
<td>ADOS repetitive and restrictive</td>
<td>40</td>
<td>3.5 (1.6)</td>
<td>40</td>
</tr>
<tr>
<td>Number of words produced</td>
<td>39</td>
<td>126.6 (153.0)</td>
<td>39</td>
</tr>
<tr>
<td>Minutes asleep before task</td>
<td>40</td>
<td>53.6 (22.3)</td>
<td>40</td>
</tr>
</tbody>
</table>

ADOS = Autism Diagnostic Observation Schedule; SD = standard deviation.
2000). For the region of interest analysis, we divided bilateral superior temporal gyrus into three sections (anterior: left: 279 voxels, right: 277 voxels), middle (left: 157 voxels, right: 156 voxels) and posterior (left: 219 voxels, right: 218 voxels) with equal anterior-posterior extents. For each speech condition, we used a repeated-measures general linear model to test for main effects of group (a between-subjects factor), hemisphere (a within-subjects factor), region (a within-subjects factor) and two- and three-way interactions between them on the number of voxels that showed a positive response (t > 2.0).

Follow-up analyses within each group examined the effects of hemisphere and region and their interaction. To examine how age was related to laterality of response, we added age to the model as a between-subject covariate for any condition that had shown hemisphere effects.

To further examine the localization of possible group laterality differences in the superior temporal gyrus and to replicate our previously published findings, we conducted a voxel-wise analysis in which the magnitude of response to forward speech in each voxel of the right superior temporal gyrus was subtracted from the value in the corresponding voxel in the left hemisphere to create a laterality map. These maps were then compared between the autism spectrum disorder and typically developing groups with a two-sample t-test. Clusters were considered to be significant if they were >32 voxels in volume and each voxel t-value was >1.66 (P_{one-tailed} < 0.05). This combination protected a superior temporal gyrus-wide probability of false positives of <0.05. Age and clinical correlations with mean activity in clusters of significant group effects were then examined.

## Results

Among typically developing infants and toddlers, listening to storybook passages resulted in strong bilateral superior temporal gyrus activation, positive response in midline precuneus and deactivation (i.e. less response during speech than rest) in bilateral thalamus and midline cerebellum (Fig. 1 and Table 2). Infants and toddlers in the autism spectrum disorder group showed less extensive, but still significant, response in clusters within bilateral superior temporal gyrus and activation of a region within the right inferior occipital gyrus (Fig. 1 and Table 2). Upon direct comparison, a region of left superior temporal gyrus (in Brodmann’s area 22) was significantly less responsive to speech stimuli in the group with autism spectrum disorder than the typical group (Fig. 1 and Table 2), a difference that remained significant after controlling for age (Supplementary Fig. 1). Our initial analysis found no regions in which the autism spectrum disorder group had significantly greater responsiveness than the typically developing group, although the age-corrected analysis revealed that middle occipital gyrus response was greater in the group with autism spectrum disorder than the typically developing group (Supplementary Fig. 1). Within the region of group difference in the superior temporal gyrus, there was a significantly larger mean brain response in the typical than autism spectrum disorder group when examining simple and complex story passages separately [simple: t(78) = 2.12, P_{one-tailed} = 0.01; complex: t(78) = 3.05, P_{one-tailed} = 0.001] and when examining response to the backward passages [backward: t(78) = 2.06, P_{one-tailed} = 0.02; Fig. 2]. The extent of activation in this region, as indicated by the number of voxels with t > 2.0 (individual voxel P < 0.05), also was significantly different for the forward speech conditions [simple: t(78) = 1.75, P_{one-tailed} = 0.04; complex: t(78) = 1.83, P_{one-tailed} = 0.03] but only a trend for the backward speech condition [(t(78) = 1.3, P_{one-tailed} = 0.09). When we examined age-related changes in responsiveness of this region of differential activation, we found no significant age × group interactions for the amplitude [F (1,78) = 3.1, P = 0.08] or extent [F (1,78) = 1.1, P = 0.30]. However, brain response was significantly negatively correlated with age in the autism spectrum disorder group for both amplitude [r(38) = −0.32, P = 0.04] and extent [r(38) = −0.35, P = 0.03], suggesting that the deficient response in autism spectrum disorder does not normalize with increasing age. Neither the extent nor amplitude of response in this region of the superior temporal gyrus was significantly related to clinical severity in the autism spectrum disorder group (P’s > 0.15) or to our measures of expressive and receptive language ability in either group (all P’s > 0.09).

Next, we tested the degree to which responses were lateralized in each group and examined potential laterality differences between autism spectrum disorder and typical groups within three anatomically defined portions of the superior temporal gyrus in each hemisphere. There were no main effects of group, hemisphere or region for any of the conditions, and none of the interactions with region was significant. Thus, we summed across the three superior temporal gyrus regions to examine laterality effects. Figure 3 presents the means of the two groups for extent of activation (volume of voxels in microlitres with t > 2.0) for all speech and for each individual condition in the entire superior temporal gyrus. There was a trend towards an interaction between group and hemisphere for response to complex [F(1,78) = 2.5, P = 0.06] speech, but differential laterality effects were not significant for all speech [F(1,78) = 1.44, P = 0.15], simple forward speech [F(1,78) = 1.65, P = 0.10] or backward speech [F(1,78) = 0.08, P = 0.77]. In each condition, there was no evidence of laterality of total superior temporal gyrus response within the autism spectrum disorder group (all P’s for within-group hemisphere effect > 0.4). In contrast, there was evidence for leftward laterality in the typically developing group in all but the backward condition [within-group hemisphere effects: all speech: F(1,39) = 5.3, P = 0.03; simple forward: F(1,39) = 3.9, P = 0.05; complex forward: F(1,39) = 3.8, P = 0.06; backward: F(1,39) = 0.54, P = 0.5].

To examine the relationship of age to lateralized brain response patterns, we examined the effects of age, hemisphere and group and their interactions on volume of activation summed across the three superior temporal gyrus subregions (given the lack of regional effects) for response to all speech and simple and complex forward speech (the contrasts that showed hemispheric effects). There were no significant main effects of age or group × age × hemisphere interactions. For both hemispheres, there were group × age interactions for all speech [left: F(1,76) = 3.16, P = 0.04; right: F(1,76) = 3.56, P = 0.03] and simple forward speech [left: F(1,76) = 4.09, P = 0.02; right: F(1,76) = 3.35, P = 0.03], but not for complex speech [left: F(1,76) = 0.05, P = 0.82; right: F(1,76) = 0.47, P = 0.50]. Table 3 shows the correlations between age and extent of response in each hemisphere of the entire superior temporal gyrus during all speech and simple and complex forward speech. It can be seen

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Figure 1 Clusters of significant response to all speech stimuli in the typically developing (n = 40) and autism spectrum disorder (n = 40) groups, and clusters of significant difference in brain response between groups. Colours indicate relative strength and direction of the effect size ($\eta^2$) [for individual group maps, positive values (warm colours) indicate greater brain response to speech compared with non-stimulation; and negative values (cool colours) indicate lower brain response to speech compared with non-stimulation in individual group maps; for group comparison map, positive values (warm colours) indicate greater brain response in the typically developing compared with autism spectrum disorder group]. R = right; L = left.

Table 2 Significant clusters of brain response to speech in typically developing and autism spectrum disorder infants and toddlers and the difference between them

<table>
<thead>
<tr>
<th>Cluster</th>
<th>Location</th>
<th>Voxel count</th>
<th>Centre of mass coordinates</th>
<th>Peak coordinates</th>
<th>All speech mean effect size ($\eta^2$)</th>
<th>Simple forward speech mean effect size ($\eta^2$)</th>
<th>Complex forward speech mean effect size ($\eta^2$)</th>
<th>Backward speech mean effect size ($\eta^2$)</th>
</tr>
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<tbody>
<tr>
<td>Typically developing</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>1</td>
<td>Left superior temporal gyrus</td>
<td>393</td>
<td>56L, 15P, 4S</td>
<td>54L, 13P, 8S</td>
<td>0.30</td>
<td>0.20</td>
<td>0.24</td>
<td>0.18</td>
</tr>
<tr>
<td>2</td>
<td>Right superior temporal gyrus</td>
<td>285</td>
<td>57R, 13P, 6S</td>
<td>58R, 17P, 4S</td>
<td>0.32</td>
<td>0.24</td>
<td>0.21</td>
<td>0.20</td>
</tr>
<tr>
<td>3</td>
<td>Bilateral thalamus</td>
<td>148</td>
<td>0R, 21P, 5S</td>
<td>14R, 21P, 8S</td>
<td>$-0.23^a$</td>
<td>$-0.05$</td>
<td>$-0.12$</td>
<td>$-0.18$</td>
</tr>
<tr>
<td>4</td>
<td>Midline cerebellum</td>
<td>110</td>
<td>3L, 58P, 31I</td>
<td>6L, 61P, 28I</td>
<td>$-0.22$</td>
<td>$-0.08$</td>
<td>$-0.08$</td>
<td>$-0.17$</td>
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<tr>
<td>5</td>
<td>Midline precuneus</td>
<td>58</td>
<td>1L, 82P, 41S</td>
<td>6L, 61P, 40S</td>
<td>0.22</td>
<td>0.14</td>
<td>0.15</td>
<td>0.11</td>
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<td>Autism spectrum disorder</td>
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<td></td>
</tr>
<tr>
<td>1</td>
<td>Right superior temporal gyrus</td>
<td>181</td>
<td>56R, 14P, 10S</td>
<td>58R, 5P, 8S</td>
<td>0.29</td>
<td>0.22</td>
<td>0.17</td>
<td>0.17</td>
</tr>
<tr>
<td>2</td>
<td>Left superior temporal gyrus</td>
<td>131</td>
<td>54L, 18P, 95</td>
<td>66L, 17P, 85</td>
<td>0.23</td>
<td>0.17</td>
<td>0.12</td>
<td>0.18</td>
</tr>
<tr>
<td>3</td>
<td>Left inferior occipital gyrus</td>
<td>52</td>
<td>40L, 75P, 2I</td>
<td>46L, 81P, 12I</td>
<td>0.19</td>
<td>0.16</td>
<td>0.08</td>
<td>0.10</td>
</tr>
<tr>
<td>Typically developing &gt; autism spectrum disorder</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Left superior temporal gyrus</td>
<td>40</td>
<td>55L, 1A, 6I</td>
<td>54L, 5P, 4I</td>
<td>0.09</td>
<td>0.03</td>
<td>0.11</td>
<td>0.04</td>
</tr>
</tbody>
</table>

$a^2$ values are signed to indicate direction of the effect. Negative values indicate negative response (i.e. less response during speech than during rest). A = anterior; I = inferior; L = left; P = posterior; R = right; S = superior.
that the group × age interactions are driven by opposite developmental trends in the two groups—small positive relationships with age in the typically developing group and negative relationships with age in the autism spectrum disorder group. Extent of response to all speech and simple and complex forward speech in left and right superior temporal gyrus was not significantly related to clinical severity in the autism spectrum disorder group (all P’s > 0.25) or to our measures of expressive and receptive language ability in either group (all P’s > 0.14).

To explore the precise localization of group laterality differences and to replicate previously published trends (Redcay and Courchesne, 2008), we undertook a voxel-based analysis in which we directly subtracted response levels between homologous left and right hemisphere voxels during the forward speech
These regions (all ship of the measured clinical variables with speech response in significant sphere, the direction of the interaction was similar but not trum disorder group \( P = 0.96 \), but was negatively related to age in the autism spec-

Table 3 Relationship of age and extent of hemispheric brain response in all speech and simple and complex forward speech conditions in the entire superior temporal gyrus region of interest

<table>
<thead>
<tr>
<th>Measure of extent of response</th>
<th>Group</th>
<th>Typically developing (n = 40)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Autism spectrum disorder (n = 40)</td>
<td></td>
</tr>
<tr>
<td>Response to all speech</td>
<td>( r )</td>
<td>( P )</td>
</tr>
<tr>
<td>Left</td>
<td>(-0.20)</td>
<td>(0.22)</td>
</tr>
<tr>
<td>Right</td>
<td>(-0.26)</td>
<td>(0.11)</td>
</tr>
<tr>
<td>Response to simple forward speech</td>
<td>( r )</td>
<td>( P )</td>
</tr>
<tr>
<td>Left</td>
<td>(-0.32)</td>
<td>(0.04)</td>
</tr>
<tr>
<td>Right</td>
<td>(-0.28)</td>
<td>(0.08)</td>
</tr>
<tr>
<td>Response to complex forward speech</td>
<td>( r )</td>
<td>( P )</td>
</tr>
<tr>
<td>Left</td>
<td>(-0.05)</td>
<td>(0.77)</td>
</tr>
<tr>
<td>Right</td>
<td>(-0.13)</td>
<td>(0.44)</td>
</tr>
</tbody>
</table>

The bold values indicate the one significant correlation (and its \( P \)-value).

Discussion

In response to a simple bedtime story, sleep functional MRI revealed that infants and toddlers with autism display abnormally reduced left temporal cortex activity and that this difference from normal grows more pronounced with age. Furthermore, whereas typical toddlers show the expected pattern of left side dominance in response to stories, toddlers with autism exhibit reversed or absent laterality patterns. Specifically, toddlers with autism display stronger activation on the right relative to the left in the anterior portion of the superior temporal gyrus, the brain region most strongly responsive to language sounds in typically developing toddlers. This abnormal lateralization is consistent with studies of older autistic children and adults (Dawson et al., 1989; Bruneau et al., 1999; Muller et al., 1999; Flagg et al., 2005; Kleinhans et al., 2008; Minagawa-Kawai et al., 2009; Anderson et al., 2010). Thus, one of the earliest appearing signature deficits in autism, failure to develop normal language comprehension, may be due to very early defects in the superior temporal gyrus and these defects may last across the lifetime.

We hypothesize that a failure of left hemisphere specialization during language comprehension in early life may not only delay basic language acquisition in infants and toddlers with autism, but may also impair the development of social language behaviours, such as analysis of prosodic information, which is typically mediated by right hemisphere neural processes (Friederici and Alter, 2004). In our study, the right hemisphere, relative to the left, was disproportionately involved in processing speech sounds in toddlers with autism spectrum disorder. One possibility is that the right superior temporal gyrus is compensating for ineffectual processing by the left superior temporal gyrus, thus crowding out the development of more social language abilities that would normally require activity of that hemisphere. Developmental neuroscientists have long known that language and social development are inextricably linked. Consider the elegant study by Kuhl and colleagues (2003) that revealed that American babies only learned

Figure 4 (A and B) Voxel-based within-group laterality score maps during the forward speech condition (expressed as signed \( \eta^2 \) effect size values; warm colours = left greater than right; cool colours = right greater than left) for typically developing and autism spectrum disorder groups in the superior temporal gyrus (shown at 51 left/right). (C) Group difference map showing the cluster of significant difference in laterality between groups, with more left-lateralized values in the typically developing compared with autism spectrum disorder group [36 voxels (each with \( t = 1.66, P_{\text{one-tailed}} = 0.05 \)], superior temporal gyrus-wide \( P < 0.05 \), with a peak at 50 left, 5 posterior and 4 superior].
Mandarin phonemes within the context of natural play interactions with a Chinese Mandarin speaker. Babies performed at chance levels when exposed to the identical linguistic information delivered via television or audio tape alone.

Why is the left superior temporal gyrus hypoactive in autism to begin with and why does it fail to show a steady increase in activity across development (in fact, showing less activity with age)? One possibility may relate to a wider problem of a failure of the superior temporal gyrus to develop effective functional connections during early development. Successful function of brain systems that support sophisticated language depend on the initial coordinated activity of widespread networks. For example, between 6 and 12 months, Imada and colleagues (2006) showed that infants recruit both traditional auditory areas in superior temporal gyrus as well motor areas in the frontal lobes in response to auditory signals. Slightly later in development, Redcay and colleagues (2008) showed that in response to bedtime stories, typical 2- and 3-year olds recruited multiple brain areas in frontal, occipital, cerebellar as well as temporal cortices. In the current study, the typically developing group engaged parietal, subcortical and cerebellar regions in addition to the strong response within the superior temporal gyrus. In autism, functional connectivity studies generally report highly aberrant patterns of functional connectivity usually in the form of underconnectivity (Just et al., 2004; Pierce and Eylar, 2011), although sometimes overconnectivity (Muller et al., 2011). These studies, however, have almost exclusively been conducted with older children, adolescents and adults with the disorder. Connectivity studies during the toddler years in autism are almost non-existent, with the single exception of a study examining inter-hemispheric correlation patterns between brain areas key to language development, such as the superior temporal gyrus and inferior frontal gyrus, in toddlers with autism (Dinstein et al., 2011). Results from that study showed reduced correlations between right and left superior temporal gyrus and inferior frontal gyrus in 2-year olds with autism relative to typically developing as well as language delay contrast groups. The current study demonstrated negative correlations of superior temporal gyrus activations with age in the autism spectrum disorder group, which may suggest that early connectivity problems lead to further failure of functional specialization during early development. One key to answering how frontal and temporal systems connect with each other and mature in autism would be to perform diffusion tensor imaging as well on subjects and in toddlers with autism spectrum disorder (Dinstein et al., 2011). The relationship between brain areas key to language development, such as the left planum temporale, appear to be enlarged even in foetuses and infants (Witelson and Pallie, 1973; Chi et al., 1977). Development of functional lateralization of speech processing has also been examined using functional near-infrared spectroscopy in awake typically developing infants and toddlers. Our sleep functional MRI findings are consistent with near-infrared spectroscopy results showing that activation of temporal cortex is reliably observed in response to hearing language stimuli (Rossi et al., 2011) and showing evidence of early emergence of leftward lateralization, at least for natural speech (Pena et al., 2003; Kotilahti et al., 2010). Rightward lateralization of functional near-infrared spectroscopy response to prosodic versus flattened stimuli (Homae et al., 2007) has also been shown at 10 months of age, but our stimuli were not designed to isolate prosody from other types of speech processing.

Interestingly, there were no statistically significant relationships found between the volume or strength of activation in the left superior temporal gyrus and language ability or symptom severity in toddlers with autism spectrum disorder. This could be due to some instability of language test scores for the youngest toddlers in our study, which were in some cases as young as 12 months in age. Clearer relationships may emerge based on language and symptom profiles at slightly older ages (e.g. 3 years old) as we have found in other studies from our laboratory (Redcay and Courchesne, 2008; Dinstein et al., 2011). The relationship between early functional brain profiles ascertained between 12 and 18 months and outcome at 3–4 years in age will be examined in follow-up studies.

Language development most certainly involves a complex interplay between environmentally driven and genetic factors (Quartz and Sejnowski, 1997; Kuhl, 2010) and disentangling contributions
of each has puzzled scientists for centuries. Considering the influence of environment, studies have shown that exposure to language in the first year of life impacts the brain’s neural circuitry (Kuhl, 2010 for review). For example, while babies are born with a universal ability to detect phoneme contrasts from all the world’s languages (Eimas et al., 1971), by the time they reach 6–10 months in age, they become increasingly more skilled in discriminating phonemes from their native language environment (i.e., what they hear most frequently) and lose the ability to discriminate sounds from other cultures (Kuhl et al., 2006). This critical period, which peaks around 9-months in age, is within a time window before parents become clearly aware of their child’s autism diagnosis, which typically is confirmed sometime around the child’s fourth birthday in the USA (Prevention, 2009). As such, the early language environment between birth and the first birthday for babies destined to develop autism, in contrast to those destined for a typical trajectory, are likely similar. Yet, by the first birthday, our study demonstrates that the brain region highly engaged in language processing, the superior temporal gyrus, is under-responsive in toddlers with autism spectrum disorder and shows reversed laterality. Why is this the case? Although adverse environmental influences that may be occurring selectively in this group during this time period cannot be ruled out, our results of very early superior temporal gyrus abnormality in autism, at face value, suggest a strong genetic component.

While the genetics of language development are far from clear, studies have indeed demonstrated that genes related to language development such as contactin-associated protein-like 2 (CNTNAP2), are dysregulated in autism (Ailarcon et al., 2008; Scott-Van Zeeland et al., 2010). Ongoing studies in our laboratory aim to elucidate the pathways from genes to aberrant functional lateralization in autism spectrum disorder.

Some potential limitations to this study should be noted. First, although this is the largest functional MRI study of very young children with autism spectrum disorder to date, it is still possible that we failed to detect small differences between groups or complicated developmental interactions. In addition, the cross-sectional nature of the presented imaging data limits our ability to make inferences about within-person trajectories of change, although our longitudinal measurement of clinical status is a strength of the design. In the present analysis, we were not able to test the specificity of our findings to those who eventually develop autism compared with those who go on to experience other developmental delays. In our ongoing study, however, we are collecting imaging data from relevant comparison groups and are also conducting follow-up functional scans for future longitudinal analyses. Finally, although our functional MRI results from sleeping babies are similar to those seen in awake typical children using other methods, we have limited information about the sleep stage of the participants in the study. Sleep latency was not different between the groups and did not impact the findings of group differences, however.

In sum, the cortical functional lateralization abnormality observed in this study appears to be early and invariant and may reflect a core, fundamental neurodevelopmental pathology in autism. As such, these aberrant response patterns may serve as the platform for the first neurofunctional biomarker of autism. Given the very early emergence of these functional deficits, our data highlight the importance of very early detection. In this way, affected infants stand the best chance to correct and refine functional defects before they become a permanent hindrance to actualizing their full cognitive and emotional potential.

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Supplementary material

Supplementary material is available at Brain online.

References


