# Atypical frontal-posterior synchronization of Theory of Mind regions in autism during mental state attribution

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This study used fMRI to investigate the functioning of the Theory of Mind (ToM) cortical network in autism during the viewing of animations that in some conditions entailed the attribution of a mental state to animated geometric figures. At the cortical level, mentalizing (attribution of metal states) is underpinned by the coordination and integration of the components of the ToM network, which include the medial frontal gyrus, the anterior paracingulate, and the right temporoparietal junction. The pivotal new finding was a functional underconnectivity (a lower degree of synchronization) in autism, especially in the connections between frontal and posterior areas during the attribution of mental states. In addition, the frontal ToM regions activated less in participants with autism relative to control participants. In the autism group, an independent psychometric assessment of ToM ability and the activation in the right temporoparietal junction were reliably correlated. The results together provide new evidence for the biological basis of atypical processing of ToM in autism, implicating the underconnectivity between frontal regions and more posterior areas.

Keywords: Autism, Mentalizing, Theory of Mind (ToM), Mental state, fMRI, Underconnectivity.

#### INTRODUCTION

Even though difficulty with Theory of Mind (ToM) processing in autism has been demonstrated in many studies (e.g., Baron-Cohen, Leslie, & Frith,

1985; Baron-Cohen, Campbell, Karmiloff-Smith, Grant, & Walker, 1995; Leekam & Perner, 1991; Perner, Frith, Leslie, & Leekam, 1989; Reed & Peterson, 1990; Swettenham, 1996; Swettenham, Baron-Cohen, Gomez, & Walsh, 1996), only a few

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neuroimaging studies have explored this domain in autism (e.g. Happe et al., 1996; Castelli, Frith, Happé, & Frith, 2002; see Frith, 2001 for a review). The impairment in ToM (processing the mental states of others) in autism often results in difficulties with the communications and interactions that underpin everyday life. Attributing mental states (or mentalizing) is perhaps one of the most complex forms of reasoning in which humans engage. It involves the interplay of a set of subprocesses, such as representation of reality, understanding one's own beliefs and the beliefs of others, and decoupling beliefs from reality. Disruption of such complex information processing is a fundamental impairment in people with autism (Minshew, Goldstein, & Siegel, 1997). At the cortical level, such complex information-processing requires the integration of the activity of multiple brain regions. According to the underconnectivity theory of autism (Just, Cherkassky, Keller, Kana, & Minshew, 2007), whenever interregion connectivity and coordination come into play, an underconnected system can manifest impairments, particularly when there is a large load on the system. ToM processing is supported by a network of several discrete neural structures that require connectivity among themselves to provide the integrated functioning. The purpose of the study reported here was to establish whether the differences in ToM processing in autism are related to a disruption of processing at the cortical network level and not just at the level of individual cortical areas.

The previously proposed theory of cortical underconnectivity (Just, Cherkassky, Keller, & Minshew, 2004) starts with the propositions that interregional connective circuitry in the brain is disrupted in autism, and that patterns of thought that are particularly dependent on integration of frontal and more posterior contributions are disrupted (Just et al., 2007). Furthermore, the theory attributes ToM deficits and executive function disruption to such underconnectivity. It proposes a causal link between the anatomical, physiological (brain activity), and psychological phenomena. Specifically, the theory posits that the communication bandwidth among cortical areas, particularly between frontal and posterior areas, is lower in autism than in typical participants. Bandwidth, which refers here to the amount of information that can be transmitted between cortical centers per unit time, is a critical factor in the performance of a computational network. Brain imaging research has been very

clear in showing that human thought involves a network of cortical areas whose activity is coordinated (synchronized), and that coordination has to be based on interregional communication, using the white matter tracts that provide the anatomical connectivity. The functional magnetic resonance imaging (fMRI) findings emerging in the past few years indicate that the synchronization between frontal and posterior areas is lower in autism (Just et al., 2004; Just et al., 2007; Kana, Keller, Cherkassky, Minshew, & Just, 2006; Koshino et al., 2005; Villalobos, Mizuno, Dahl, Kemmotsu, & Muller, 2005). For example, in a Tower of London (TOL) problem-solving task, which entails activation of both frontal and parietal areas, the synchronization between the frontal and parietal areas is lower in autism than in a control group.

Neuroimaging studies of ToM processing have identified a frontal-posterior network of brain regions that include the medial prefrontal cortex, the temporal-parietal junction at the top of the superior temporal sulcus, and the temporal poles adjacent to the amygdala (Brunet, Sarfati, Hardy-Bayle, & Decety, 2000; Castelli, Frith, Happé, & Frith, 2000; Castelli, Happé, Frith, & Frith, 2002; Fletcher et al., 1995; Gallagher et al., 2000; Vogeley et al., 2001). There has been some debate concerning the differential roles of frontal versus posterior regions in ToM processing. Although the study reported below does not focus on differentiating the contributions of various brain areas to ToM processing, it is nevertheless useful to acknowledge that different areas are very likely playing different roles. Some researchers argue that while the medial frontal cortex is directly involved in mental state reasoning and decisions, the right temporoparietal junction collates the cues that are the input to mentalizing (Gallagher & Frith, 2003). Others argue that the right temporoparietal junction is the central player in mental state reasoning, not only collating but also processing the cues to compute a mental state (Saxe & Kanwisher, 2003). Regardless of the precise computation that each area performs in ToM processing, it appears that the two regions function in concert with each other. It is this coordination among the contributing areas that is the focus of this article.

The communications with the frontal components of the ToM network are of special interest here, because of the hypothesis of lower frontalposterior functional connectivity in autism. A number of studies indicate that the role of frontal lobe regions in processing ToM is major. Evidence for this view comes in large part from lesion research. Patients with damage to frontal cortex are frequently impaired in the performance of ToM tasks (Stuss, Gallup, & Alexander, 2001). Damage to the anterior prefrontal regions has been associated with impaired self-awareness for the appropriateness of social interactions, judgment and planning difficulties (Stuss, 1991), as well as impaired awareness of the mental states of others (Stone, Baron-Cohen, & Knight, 1998; Stuss et al., 2001).

It may not be just the frontal activation that is critical; the synchronization of frontal activity with activity in posterior ToM areas, as well as other areas, may be important in accomplishing mentalizing. From an anatomical perspective, major afferents to the medial frontal regions come from the dorsolateral prefrontal cortex, temporal pole, anterior superior temporal gyrus, parietotemporal cortex, and posterior cingulate cortex (Barbas, Ghashghaei, Dombrowski, & Rempel-Clower, 1999; Carmichael & Price, 1996; see Amodio & Frith, 2006 for a review). Therefore, the disruption of connections from and to the frontal lobe may affect higher cognitive functions, such as ToM processing.

The present study investigates the synchronization of brain activation in autism using a ToM task involving an animation of two interacting triangles. ToM processing in autism using these animations has been studied previously using PET (Castelli et al., 2002). The rationale for the present study is that: (1) it used the higher temporal resolution of fMRI to examine the synchronization of brain activation (functional connectivity) in autism in a ToM task for the first time, and (2) the current task targeted the explicit online assessment of mental state attribution in people with autism. We predicted reduced brain activation in autism in ToM processing regions

such as medial frontal cortex and right temporoparietal junction. This hypothesis is based on the findings from studies of impairment in ToM (Castelli et al., 2002), and perception of biological motion (Blake, Turner, Smoski, Pozdol, & Stone, 2003) in autism. The novel portion is our prediction of functional underconnectivity in autism during mentalizing, especially in connections involving the frontal lobe. This prediction is based on previous findings of functional underconnectivity in autism in a wide variety of higher cognitive tasks in autism (Just et al., 2007; Kana et al., 2006; Castelli et al., 2002). The underconnectivity prediction is also based on evidence from structural brain imaging studies which indicate abnormalities in autism of the white matter that provides the communication channels between brain areas (Herbert et al., 2003, 2004; Keller, Kana, & Just, 2007; Barnea-Goraly et al., 2004).

#### **METHOD**

# **Participants**

Twelve high-functioning adult individuals with autism (mean age 24.6 years) and 12 typical control participants (mean age 24.4 years) were included in the analyses (full-scale and verbal IQ scores of 75 or above). Participants were matched on the basis of age and IQ (see Table 1). Among the 12 participants in the autism group, two were females and one was left-handed. In the control group, all participants were males and one was left-handed. Data from eight other participants (five with autism and three controls) were discarded due to excessive head motion. The diagnosis of autism was established using two structured research diagnostic instruments, the ADI-R (Autism Diagnostic Interview—Revised;

 TABLE 1

 Age, IQ, handedness, and gender of participants

	Autism	Control	
Age (years: range; mean $\pm SD$ )	15.8-35.7; 24.6±6.9	16.6–31.4; 24.4±3.7	
Verbal IQ (range; mean $\pm SD$ )	81–135; 104.1±14.4	89–126; 110.6±9.9	
Performance IQ (range; mean $\pm SD$ )	75–129; 102.8±13.6	$100-127; 115.2\pm7.4$	
Full-scale IQ (range; mean $\pm SD$ )	83–136; 104.3±14.4	99–129; 114.3±8.6	
Handedness (right: left)	11: 11	1:1	
Gender (male: female)	10: 2	12: 0	

Notes: IQ, intelligence quotient; SD, standard deviation.

Lord, Rutter, & LeCouteur, 1994) and the ADOS-G (Autism Diagnostic Observation Schedule—Generic, Lord et al., 2000), supplemented with confirmation by expert opinion. None of the participants were diagnosed with Asperger syndrome. Potential participants with autism were excluded on the basis of an associated infectious or genetic disorder, such as fragile-X syndrome or tuberous sclerosis. Potential control participants and participants with autism were also excluded if found to have evidence of birth asphyxia, head injury, or a seizure disorder. Exclusionary criteria were based on neurologic history and examination, and chromosomal analysis.

The control participants were medically healthy community volunteers recruited to match the participants with autism on age, full-scale IQ, gender, race, and family of origin socioeconomic status, as measured by the Hollingshead method. Potential control participants were screened by questionnaire, telephone, face-to-face interview, and observation during screening psychometric testing to determine eligibility. Exclusionary criteria, evaluated through these procedures, included current or past psychiatric and neurologic disorders, birth injury, developmental delay, school problems, acquired brain injury, learning disabilities, and medical disorders with implications for the central nervous system or those requiring regular medication. Potential control participants were also screened to exclude those with medical illnesses or a family history of autism, developmental cognitive disorder, affective disorder, anxiety disorder, schizophrenia, obsessive compulsive disorder, or other neurologic or psychiatric disorders thought to have a genetic component (in first-degree relatives or self), medications that affect the CNS, hypertension, diabetes, substance abuse (self, or firstdegree relative), steroid use (extreme use such as steroids used in inhalers for asthmatics) and autism in first-, second- or third-degree relatives. Each participant signed an informed consent that had been approved by the University of Pittsburgh and Carnegie Mellon University Institutional Review Boards.

# **Experimental paradigm**

This experiment compared autism and control participants on attributing mental states to the movement of geometrical figures. The animations

were generously provided by Dr Fulvia Castelli, who performed the original PET study of this task (Castelli et al., 2002). All animations featured two "characters"—a large red triangle and a small blue triangle—moving about on a framed white background.

Three types of animations were used in this study: the ToM, the Goal-Directed (GD), and the Random (RD) formed three experimental conditions. In the ToM condition, the geometrical figures engaged each other with intentional action/interaction involving thoughts and feelings. For example, a character's actions were determined by what the other character thought. In the GD condition, the geometrical figures engaged in an interaction with each other in a simple purposeful level. For example, one character's actions were determined by what the other character did. In the RD condition, the geometrical figures did not engage with each other at all. For example, all character actions were random and unassimilated. There were three trials each of ToM, GD, and RD stimuli, each trial containing one animation. The basic visual characteristics depicted in the three types of animations were similar in terms of shape, overall speed of motion, orientation changes, and total duration (between 26 and 47 s).

After the presentation of each animation, four words were presented on the screen, and participants were asked to make a forced-choice judgment about which of the words best described the action depicted in each animation. (Note the dissimilarities to Castelli et al.'s (2002) procedure, which required only passive viewing of the animations and reporting after the scan what had happened in the animation.) The correct response was always an accurate description of the animation (determined as the most frequently generated description in a norming study). Another response alternative was an inaccurate description of the animation, but of the appropriate category (e.g., in the ToM condition, this incorrect response referred to a mental state, but not the correct mental state). The other two answers were inaccurate descriptions of the animation that could have applied to animations in the other two conditions (i.e., in the ToM condition, these incorrect response alternatives referred to a random motion and a goal-directed motion, in the GD condition they referred to a ToM motion and a random motion, and in the RD condition they referred to a ToM motion and a goal-directed motion). For example, for a ToM animation that depicted "coaxing", the foils were pushing (GD), surprising (ToM), and spinning (RD). Participants made their responses using two two-button mice, one held in each hand. Each button corresponded to one of the four multiplechoice answers. Responses were accepted for 15 s from the end of each animation. The presentation of each animation constituted a separate event in the experimental design. The animations were presented in blocks of three, one from each condition, with a separation of 6 s between trials within a block. The onset of each animation was synchronized with the beginning of a TR. A 30-s fixation condition was presented between each block to provide a baseline measure of brain activation with which to compare each experimental condition. In this fixation condition, participants fixated on a centered asterisk without performing any task. A practice animation belonging to the RD condition was presented at the beginning of the experiment to familiarize the participant with the task.

We also administered a separate neuropsychological test outside the scanner to get a different measure of participants' ToM abilities. The Happe Strange Stories Test for the assessment of theory of mind was administered as a neuropsychological test just to get an independent outof-scanner measure of the ToM abilities of the participants with autism.

Prior to testing in the scanner, participants were familiarized with the task using one example of each of the three conditions that was not presented again during the fMRI study. Familiarization also entailed the use of a scanner simulator to acclimate the participants and to attain motion quality standards.

# fMRI procedures

All imaging data were acquired at the Brain Imaging Research Center (BIRC) co-administered by Carnegie Mellon University and the University of Pittsburgh on a 3-T Siemens Allegra scanner. The stimuli were rear projected onto a semi-translucent plastic screen and participants viewed the screen through a mirror attached to the head coil. For the functional imaging, a gradient echo, echo-planar pulse sequence was used with TR = 1000 ms, TE = 30 ms, and a flip angle of  $60^{\circ}$ . Sixteen adjacent oblique-axial slices were acquired in an interleaved sequence, with 5mm slice thickness, 1-mm slice gap, a  $20 \times 20$  cm FOV, and a  $64 \times 64$  matrix, resulting in in-plane resolution of  $3.125 \times 3.125$  mm. A 160-slice 3D MPRAGE volume scan with TR = 200 ms, TE = 3.34 ms, flip angle = 7°, FOV = 25.6 cm, 256 × 256 matrix size, and 1-mm slice thickness was acquired at the same orientation as the obliqueaxial functional images for each participant. This structural scan was used for making measurements of corpus callosum size.

# **Distribution of activation**

To compare the participating groups in terms of the distribution of activation, the data were analyzed using SPM99. Images were corrected for slice acquisition timing and head motion, and they were normalized to the Montreal Neurological Institute (MNI) template, resampled to  $2 \times$  $2 \times 2$  mm voxels, and smoothed with an 8-mm Gaussian kernel to decrease spatial noise. Highpass filtering and global scaling were performed on each participant's data. Statistical analysis was performed on individual and group data by using the general linear model as implemented in SPM99 (Friston et al., 1995). Group analyses were performed using a random-effects model. Contrasts reflecting the group differences in the distribution of activation relative to fixation, and group-by-condition interactions were computed. For the group difference contrasts, possible differences in deactivation (relative to fixation condition) were excluded because previous studies have shown that medial prefrontal areas that activate in response to ToM tasks also show deactivation relative to a resting baseline for a variety of other cognitive tasks (e.g., Cherkassky, Kana, Keller, & Just, 2006). This masking procedure ensured that group differences were not due to greater activation during fixation in one of the groups relative to the other. An uncorrected height threshold of p < .005 and an extent threshold of six voxels were used.

# **Functional connectivity**

The functional connectivity was computed (separately for each participant) as a correlation between the average time course of signal intensity (for a region of interest (ROI) this was calculated separately within each participant by averaging the signal intensity separately for each time point (image) across activated voxels within the ROI) of all the activated voxels in each member of a pair of ROIs. Fifteen functional ROIs were defined to encompass the main clusters of activation in the group activation map for each group in the ToM-Fixation contrast. Labels for these 15 ROIs (the superior medial frontal gyrus (SMedFG), left and right inferior orbital frontal cortex (LIOFC and RIOFC), left precentral gyrus (LPrecen), right inferior frontal gyrus (RIFG), left and right inferior parietal lobes (LIPL and RIPL), left and right superior parietal lobes (LSPL and RSPL), right middle temporal gyrus (RMTG), right superior temporal gyrus (RSTG), left and right fusiform gyrus (LFFG and RFFG), and left and right inferior occipital gyrus (LIOG and RIOG)) were assigned with reference to the parcellation of the Montreal Neurological Institute (MNI) single-subject T1weighted dataset carried out by Tzourio-Mazover and colleagues (2002). For each of these ROIs, a sphere was defined for each cluster (with a radius ranging from 4 to 8 mm) that best captured the cluster of activation in the map for each group. The ROIs used in the analysis were each the union of the two spheres defined for the two groups (a complete table of the location and size of each of the ROIs within each group can be found in the supplementary material available online). Within these ROIs data were extracted from voxels that exceeded a *t*-threshold of 4.5 (corresponding approximately to a significance level of p < .10, corrected for multiple comparisons) in at least one contrast between an animation condition and the fixation baseline from the individual participant's first-level analysis. In addition, we required that at least 12 voxels were active within a ROI for an individual participant to ensure a stable estimate of the time-course. The activation time-course extracted for each participant over the activated voxels within the ROI originated from the normalized and smoothed images that were high-pass filtered and had the linear trend removed. The correlation was computed on the images belonging to the random, goal-directed, and ToM conditions, so it reflects the interaction between the activation in two areas while the participant is performing the task, and not during the baseline condition. Fisher's r to z transformation was applied to the correlation coefficients for each participant prior to averaging and statistical comparison of the two groups.

Functional connectivity was measured for each participant in each group for the ToM, GD, and RD conditions using the 15 functional ROIs described above. On the basis of the prediction that frontal-posterior networks would show significant underconnectivity in autism (Just et al., 2007; Kana et al., 2006), these 15 ROIs were grouped into larger regions on the basis of lobe (frontal, parietal, temporal, or occipital), and then functional connectivity measures for these groups of ROIs were obtained for each participant by averaging the connectivities of all of the relevant ROI pairs. This resulted in 10 "networks" for which connectivities were aggregated: six interlobe connectivities (frontal-parietal, frontal-temporal, temporal-parietal, frontal-occipital, temporal-occipital, and parietal-occipital), and four intralobe connectivities (within frontal, temporal, parietal, and occipital). In addition, we calculated functional connectivity between the frontal centers of the ToM network (medial frontal, orbitofrontal) and the posterior centers (right middle and superior temporal gyri, temporoparietal junction).

# Corpus callosum morphometry

The cross-sectional area of the midsagittal slice of the corpus callosum was measured using the parcellation scheme described by Witelson (1989). The seven regions of the corpus callosum defined in this scheme include the rostrum, genu, rostral body, anterior midbody, posterior midbody, isthmus, and splenium. The corpus callosum size was normalized (divided by) by the total grey and white matter volume for each participant. The grey matter, white matter, and cerebrospinal fluid (CSF) volumes were measured for each subject by segmenting the T1-weighted structural brain image into three masks. The segmentation was performed by SPM2 routines. The outer contour of the corpus callosum was manually traced (with an interrater reliability of .87), and then interior segmentation, area, and length computations were performed by image processing software.

TABL	.E 2
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Brain activation within autism and control groups when theory of mind processing was contrasted with processing of random animations.

Location of peak activation	Brodmann's area	t(11)		MNI coordinates		
			Cluster size	x	у	z
Participants with autism						
R inferior frontal gyrus	47	25	7.63	52	30	0
L inferior frontal-orbital	47	51	7.46	-48	38	-14
R fusiform gyrus	39	49	6.17	42	-54	-24
R inferior frontal gyrus	9	46	5.42	46	8	24
R middle temporal gyrus	22	28	4.97	58	-52	6
Typical control participants						
L superior medial frontal gyrus	9	1037	12.37	-10	56	42
L angular gyrus	39	1009	11.16	-50	-72	28
L inferior frontal gyrus	45	228	8.18	-54	20	8
R middle temporal gyrus	13	1006	7.70	44	-48	18
R precuneus	7	407	7.08	0	-62	32
L middle frontal gyrus	8	43	7.01	-42	18	48
R middle temporal gyrus	21	172	6.91	60	-10	-18
R fusiform gyrus	37	25	6.22	44	-46	-22
L middle temporal gyrus	21	113	6.17	-54	-12	-24
L fusiform gyrus	37	125	6.01	-42	-56	-14
L middle temporal gyrus	20	33	5.35	-54	-34	-10
R inferior frontal gyrus	45	87	5.30	60	24	22

*Notes*: L, left; R, right. Entries in **bold** type are significant at p < .05 corrected for multiple comparisons on the basis of the cluster size. Other entries are significant at p < .001, uncorrected. Region labels, *t*-values, and MNI coordinates are for the peak activated voxel in each cluster only.

# RESULTS

#### Overview

The central findings indicate that during the attribution of mental states to animated figures, participants with autism (relative to control participants): (1) showed reduced functional connectivity within the ToM network as well as in the connections between the frontal lobe and more posterior regions (to our knowledge, this is the first finding linking ToM processing in autism to the lowered synchronization of activation across relevant brain areas); (2) showed reduced levels of brain activation in the frontal components of the ToM network (medial prefrontal cortex, anterior cingulate cortex, and orbitofrontal gyrus); (3) showed a positive correlation between ToM ability (measured psychometrically) and the activation in right superior temporal gyrus; and (4) showed a smaller midsagittal area of the rostrum of the corpus callosum than in controls. As described in more detail below, the findings are consistent with disrupted ToM processing in autism being underpinned by frontal-posterior functional underconnectivity.

#### **Behavioral results**

The behavioral results reported here are based on performance in the fMRI task. The autism and control groups performed similarly. There was no statistically reliable difference between the autism and control groups in either reaction time, F(1, 22) = 1.20, p = .29, or error rate, F(1, 22) = 2.50, p = .13, nor was there a statistically reliable interaction between the groups and conditions for either response measure, F(2, 44) = 0.82, p = .45, and F(2, 44) = 1.85, p = .17. There was an effect of condition overall, such that the ToM condition showed significantly longer reaction times, F(2, 44) = 25.12, p < .001, and significantly higher error rates, F(2, 44) = 14.63, p < .001.

# Contrast between ToM and random motion for both groups

In the ToM condition, both the autism and the control group showed activation in regions associated with ToM, such as the medial frontal gyrus, orbitofrontal gyrus and right middle temporal gyrus. However, the autism group clearly showed

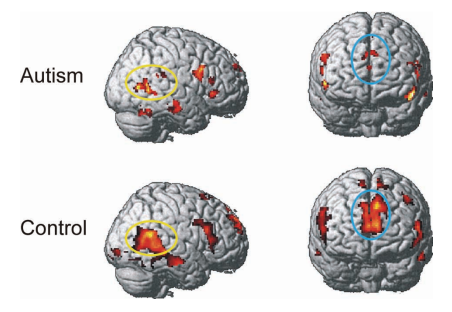


Figure 1. Activation in right superior temporal (yellow ovals) and superior medial frontal (blue ovals) areas in autism and control groups for the contrast of ToM processing with Random animation processing. The t-maps are thresholded at p < .005 and an extent threshold of 6 voxels here for display.

reliably less activation in medial frontal regions compared to control participants (see Figure 1 and Table 2), as further described in the next section. the superior temporal sulcus, t(22) = 3.41, p = .001, centered at x = 56, y = -28, z = 18). Overall, the autism group showed lower activation in frontal ToM regions while showing no difference in activation in the posterior ToM region.

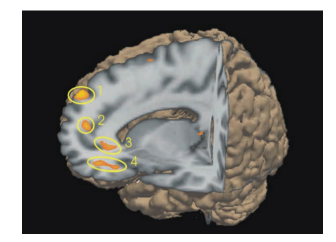
# Group differences in brain activation

There were statistically reliable group differences in the contrast between the ToM condition and the random animations. The autism group showed reliably lower brain activation compared to controls in several regions in the frontal cortex, including the medial frontal gyrus, anterior paracingulate cortex, anterior cingulate gyrus, and inferior orbital frontal gyrus, as shown in Figure 2. All of these regions have been found to coactivate in a "social" processing brain network in typical populations (Adolphs, 2003; Blakemore, Winston & Frith, 2004; Brothers, 1990; Castelli et al., 2002; Gallagher & Frith, 2003; Winston, Strange, O'Doherty, & Dolan, 2002).

In contrast to the lower activation shown by the autism group in frontal parts of the ToM network, there was no statistically reliable group difference in activation in the right posterior superior temporal sulcus region itself (a key posterior ToM area centered at x = 52, y = -46, z = 24; see Castelli et al., 2002). However, the autism group showed more activation than the control group in a region slightly more anterior to

#### Functional connectivity

The key hypothesis regarding functional connectivity was that in the ToM condition, synchronization would be reduced in the group with autism relative to controls between the frontal ToM areas (medial frontal, orbitofrontal) and posterior ToM areas (right middle and superior temporal gyri, temporoparietal junction). A planned contrast between the two groups comparing this frontal-posterior within-ToM-network functional connectivity within the ToM condition confirmed this prediction (autism mean = 0.20; control mean = 0.48; t(19) = 2.65, p < .05). Note that this result is not due to a failure of the group with autism to activate either frontal or right temporoparietal areas in the ToM task. Both groups showed activation in these regions for the contrast between ToM and the fixation baseline, and this contrast was used to define the ROIs for extracting the time course of signal intensity (see Table 3). In addition, the procedure for extracting the time-course of signal intensity from an individual participant included only voxels found



**Figure 2.** Participants with autism showed reduced levels of activation relative to control participants (p < .005, uncorrected) in four regions in the frontal ToM network: (1) left superior medial frontal gyrus, (2) left anterior paracingulate cortex, (3) bilateral anterior cingulate cortex, and 4) left inferior orbitofrontal cortex. The graphic was created using Slicer software.

to be activated for the participant in at least one of the contrasts of an animation condition with this baseline condition. This procedure eliminated consideration of functional connectivity in the frontal-posterior ToM network for three of the participants with autism (all normal controls had activation within the defined ROIs), but prevents differences in the response of frontal areas to ToM task from biasing the group estimate of functional connectivity by taking into account data from only those participants that activated the regions. Note also that functional connectivity measures the degree of synchronization of signal intensity among regions across time rather than

28

54

60

42

-46

42

-42

-64

-58

-40

-50

-64

-76

-80

54

8

18

-24

-18

-10

 $^{-4}$ 

RSPL

RMTG

RSTG

LFFG

RFFG

LIOG

RIOG

the amplitude of the response during the presentation of the animation.

Because reduced frontal-posterior functional connectivity in autism has previously been established in a variety of cognitive tasks and even in a resting baseline, we also tested the specificity of the difference to the requirement for ToM processing by examining the functional connectivity in the RD and GD conditions. Participants with autism showed lower functional connectivity than controls in all three conditions, although this group difference was smallest in the GD condition, as shown in Figure 3. A 3 (Condition)  $\times$  2 (Group) mixed ANOVA revealed a reliable main

54

2

20

-16

-18

-10

2

6

8

4

5

3

6

4

the autism and control groups. x, y, and z represent the MNI coordinates for the center of each ROI cluster. Control group Autism group ROI х Z, radius х z radius y v SmedFG 4 54 36 4 0 56 36 4 LIOFC -50 30 -65 -5038 -63 36 RIOFC 52 26 -84 48 -143 Lprecen -462 40 4 -484 36 5 18 RIFG 52 24 8 52 18 24 6 LIPL -40 44 48 -40 42 48 7 6 RIPL 42 -44 52 6 40 -4050 5 56 LSPL -2256 -24-666 -646

7

6

6

4

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50

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-38

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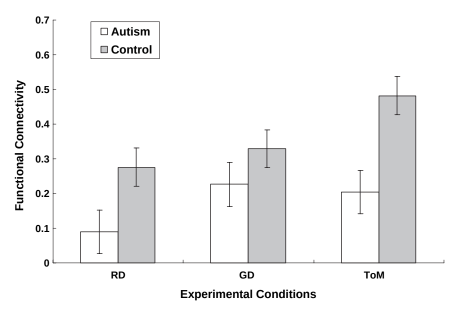
-50

-72

-74

TABLE 3

Regions of interest for the functional connectivity analysis based on within-group regions of activation for ToM-Fixation contrast for the autism and control groups, *x*, *y*, and *z* represent the MNI coordinates for the center of each ROI cluster.



**Figure 3.** Mean functional connectivity (z') in the frontal-temporal ToM network for each group and each condition. Participants with autism showed significantly reduced functional connectivity relative to control participants in the ToM condition and the Random condition, but not in the Goal-directed condition.

effect of Group, F(1, 19) = 7.56, p < .05, indicating reduced overall functional connectivity in the group with autism, and a reliable main effect of condition, F(2, 38) = 7.75, p < .005, indicating that across groups the functional connectivity was modulated by the type of animation. There was a trend toward a Condition × Group interaction, F(2, 38) = 2.27, p = .12. Tests of the simple main effects of group within each condition indicated that in addition to expected reduction in functional connectivity in the ToM condition, there was also a reliable group difference in the RD condition, t(19) = 2.58, p < .05, but not in the GD condition, t(19) = 1.49, p = .15. Within the autism group there was only a marginal effect of Condition on functional connectivity in the frontaltemporal ToM network, F(2, 16) = 3.63, p = .05, resulting from lower connectivity in the RD condition than in either the GD condition, t(16) = 2.51, p < .05, or the ToM condition, t(16) = 2.11, p = .05. In contrast, within the control group there was a reliable Condition effect, F(2, 22) = 6.82, p < .005, with significantly greater functional connectivity in the ToM condition than either the RD condition, t(22) = 3.56, p < .05, or the GD condition, t(22) = 2.64. p < .05. Thus, the overall pattern shown in Figure 3 suggests that while normal control participants show increased functional connectivity between frontal and right posterior temporal-parietal areas only when the task demanded ToM, the group with autism

showed increased functional connectivity in this network for both the Goal-directed and ToM conditions, but did not approach the degree of synchronization of activation between regions in the ToM condition as control participants.

In order to fully characterize the functional connectivity patterns involving all the regions with substantial activation in the ToM condition, an ANOVA of the mean interlobe and intralobe connectivities compared the mean functional connectivities of the two groups averaged within each group over the pairs of functional ROIs that were in different cortical lobes. Thus groups were compared across 10 possible pairings (networks) of the four lobes (six interlobe networks and six intralobe networks): frontal-parietal, frontal-temporal, frontal-occipital, temporal-parietal, temporal-occipital, parietal-occipital, frontal-frontal, temporal-temporal, parietal-parietal, and occipital-occipital. This 2 (Group)  $\times 10$  (Network) mixed ANOVA revealed a reliable main effect of network, F(9, 153) = 26.46, p < .0001, a reliable main effect of Group, F(1, 17) = 6.76, p < .05, and no reliable Network × Group interaction, indicating that even in a task targeting ToM processing, widespread reductions in functional connectivity are found in autism. Means and simple main effects of Group are shown in Table 4. As the table indicates, the group with autism showed reliably lower functional connectivity between frontal regions and posterior regions (temporal,

	Autism	Control	Group difference	F value	p value
Interlobe networks					
Frontal-parietal	0.21	0.51	0.30	8.60	0.008
Frontal-occipital	0.23	0.56	0.33	11.89	0.003
Frontal-temporal	0.32	0.57	0.25	7.75	0.012
Parietal-temporal	0.45	0.68	0.23	5.68	0.026
Parietal-occipital	0.48	0.76	0.28	6.95	0.015
Occipital-temporal	0.70	0.91	0.21	4.35	0.049
Intralobe networks					
Frontal-frontal	0.28	0.49	0.21	3.74	0.068
Temporal-temporal	0.58	0.83	0.25	3.27	0.086
Parietal-parietal	0.53	0.82	0.29	5.65	0.027
Occipital-occipital	0.67	0.95	0.28	5.81	0.025

 TABLE 4

 Functional connectivity differences in interlobe and intralobe networks between autism and control groups for the ToM condition.

parietal and occipital) compared to control participants. In addition, reliably lower functional connectivity was found in the autism group in the functional connectivities between temporal and occipital areas. In the intralobe networks, the autism participants showed reliably lower connectivity in parietal-parietal as well as occipitaloccipital connections.

# Correlation between brain activation and mentalizing ability

The brain activation in two regions of interest (the medial frontal gyrus, and the right posterior superior temporal gyrus), which are associated with processing of ToM, was correlated with the ToM score of the autism participants from the Happé strange stories test (Happé, 1994). The activation measure used in the correlation was the contrast of parameter estimates (beta-values) from the general linear model extracted from individual participants (anatomical region of interest) for the ToM vs. Random contrast. The ToM score was reliably positively correlated, r =.57, t(10) = 2.21, p < .05, with the activation in right posterior superior temporal gyrus (see Figure 4). Although the correlation was positive (r = .26) for the medial frontal gyrus ROI, the effect was not statistically significant. (The ToM scores were not available for the control participants, but one might expect scores near ceiling without much variation.)

# Corpus callosum size

To investigate a possible anatomical basis of frontal-posterior functional underconnectivity and frontal underactivation among the group with autism, we compared the size of the corpus callosum in anterior regions (the rostrum and genu) between the groups. The results showed that the rostrum was reliably smaller in participants with autism compared to control participants, t(22) = 2.547, p < .02, although no such effect was found for the genu. The rostrum of the corpus callosum connects orbital prefrontal regions in the two hemispheres, regions previously found to be involved in ToM processing (Sabbagh, 2004; Stone et al., 1998). Moreover, the present study found reduced activation in a left orbital prefrontal region in the group with autism relative to controls, providing a link between lowered activation in ToM in autism and connective white matter. Unlike several previously reported studies (Cherkassky et al., 2006; Just et al., 2007; Kana et al., 2006), this analysis found no significant correlation between the size of any corpus callosum segment and the functional connectivities between the areas that the segment connects.

#### DISCUSSION

The most important finding of the current study is that the participants with autism showed functional underconnectivity among several brain

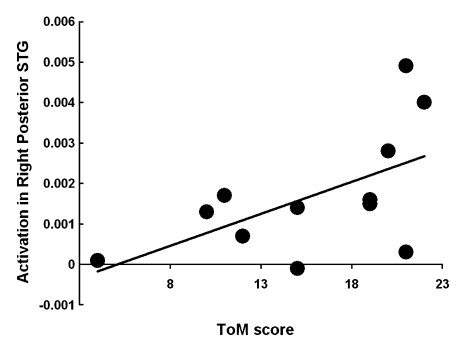


Figure 4. Participants with autism (r = 0.57) showed a significant positive correlation between ToM score and the activation in the right posterior superior temporal gyrus.

areas involved in ToM processing. As predicted, the frontal ToM regions (medial frontal gyrus, anterior paracingulate, orbital frontal gyrus) were found to be functionally underconnected with the posterior ToM regions (right middle and superior temporal gyrus) in autism during mentalizing. In addition, underconnectivity was found in the connections between the frontal lobe and other more posterior regions, and within the occipital and parietal lobes. Castelli et al. (2002) reported a correlation across autistic and control participants between the level of activation (averaged over the entire course of the task) in occipital and temporal regions in a PET study. The current study, using the same stimuli, provides new evidence of a lower degree of synchronization (with observations obtained once per second) between frontal and posterior areas in autism during the viewing of these animations. The fMRI measures provide critical detail that speaks to the underlying mechanisms that distinguish the two groups, namely the lower degree of synchronization and hence coordination among brain areas in autism.

Interestingly, functional connectivity in the frontal-posterior ToM network was modulated by condition in both groups, but in different ways. Normal control participants showed an increase in functional connectivity in this network only when viewing the ToM animations, suggesting an adaptive engagement of processes involving communication among the areas only in the condition in which such processes are necessary. In contrast, the group with autism showed an increase in functional connectivity relative to the RD condition for both the GD and ToM conditions. This differential pattern of modulation resulted in the group with autism showing reliable underconnectivity relative to controls in both the RD condition and the ToM conditions, but not in the GD condition. These results suggest that individuals with autism attempt to carry out processes dependent on communication between frontal and posterior ToM areas for both types of animation involving interactions between the triangles, but that such communication is less efficient overall and less adaptive to the demands of the task.

Although the functional underconnectivity in autism is widespread in this study, affecting not only frontal-posterior connectivity but also occipital-temporal connectivity, the most intriguing facet of this is that the frontal lobe seems to be the epicenter of underconnectivity. This observation emphasizes the significant participation of the frontal lobe and its integration with other areas in a complex task, such as mental state attribution. Lower activation in frontal areas in autism has been found in several different functions, such as language comprehension (Harris et al., 2006; Just et al., 2004), verbal working memory (Koshino et al., 2005), face working memory (Koshino et al., 2008), visuospatial processing (Ring et al., 1999), and understanding emotions of others (Dapretto et al., 2006). In addition, our previous studies of executive functioning (Just et al., 2007), visual imagery (Kana et al., 2006), response inhibition (Kana, Keller, Minshew, & Just, 2007) and discourse processing

(Mason, Williams, Kana, Minshew, & Just, 2008) have found frontal-posterior underconnectivity in autism. These findings from brain activation and functional connectivity, in conjunction with findings of structural abnormalities, such as delayed maturation of frontal lobe (Zilbovicius et al., 1995), increased frontal cortical folding (Hardan, Jou, Keshavan, Varma, & Minshew, 2004), anatomical shifting of major sulci in the frontal lobe (Levitt et al., 2003), and maldevelopment in minicolumns in the frontal cortex (Buxhoeveden, Semendeferi, Schenker, & Courchesne, 2004; Casanova, Buxhoeveden, Switala, & Roy, 2002) suggest impairment in information transfer from and to the frontal lobe in autism.

Although the differences in frontal lobe activation between groups found for the contrast between the ToM condition and the RD condition could conceivably lead to reduced functional connectivity in the group with autism if there was simply no frontal activation in this group, note that the ROIs for which the signal intensity was extracted were not based on areas showing a group difference for this contrast, but rather were defined by the within-group contrasts between the ToM condition and the fixation baseline, a contrast for which both groups showed activation in medial frontal and bilateral inferior orbitofrontal areas. In addition, the procedure for calculating functional connectivity ignored regions within participants that did not show a sufficient volume of activation in this contrast to provide a stable estimate of the time-course of signal intensity. Given a measurable and stable signal from the involved regions of interest, our measure of functional connectivity depends on the correlation of signal change among regions during viewing of the animation, rather than on the amplitude of the signal intensity within individual regions. We also note that in previous work we have found lower functional connectivity in autism despite similar levels of activation between the groups for the regions considered (e.g., Just et al., 2007) and that such differences are found even during a resting baseline condition (Cherkassky et al., 2006). Note also that in the present study we found group differences in functional connectivity in posterior regions that did not show differences in activation between the groups. Although the present study cannot distinguish whether reduced frontal activation in autism is a cause or a consequence of reduced communication between frontal and posterior ToM areas, reduced activation is not a necessary precondition for reduced functional connectivity.

Widespread white matter abnormalities have also been found in autism, including lower fractional anisotropy of the white matter adjacent to the ToM regions (medial frontal gyrus, anterior cingulate cortex, and right temporoparietal junction) (Barnea-Goraly et al., 2004), significantly larger frontal and parietal white matter volume (Carper, Moses, Tigue, & Courchesne, 2002), disturbances (both increases and decreases in volume in different areas) in white matter (Herbert et al., 2004), white matter volume deficits in right middle frontal and left superior frontal regions (Waiter et al., 2005), reduced thickness in orbitofrontal white matter (Chung et al., 2005), increased regional and global white matter (Hendry et al., 2006), and abnormalities in corpus callosum size (Egaas, Courchesne, & Saitoh, 1995; Piven, Bailey, Ranson, & Arndt, 1997; Hardan, Minshew, & Keshavan, 2000). Any or all of these abnormalities could conceivably affect the functional and structural connectivity of the frontal lobe with more posterior regions in autism. These findings taken together point to abnormalities in the neural architecture in autism, which may affect several types of psychological functions, including ToM.

The critical role of the prefrontal cortex in ToM processing is supported by neuropsychological studies that show frontal lobe damage associated with impairments in ToM (Channon & Crawford, 2000; Rowe, Bullock, Polkey, & Morris, 2001; Stone, Baron-Cohen, & Knight, 1998; Stuss et al., 2001), including the medial prefrontal cortex as well as the orbitofrontal cortex. Gallagher and Frith (2003) propose that while the posterior superior temporal sulcus provides the cues for mentalizing, the medial prefrontal cortex does the mental state reasoning. Therefore a communication between these two regions is vital in ToM tasks. In other words, cortical underconnectivity in autism among the components of the network for mentalizing provides a plausible neural basis of impaired ToM in autism. The underconnectivity could simply be at the level of a lack of information coordination

between two areas. However, in autism, structural abnormalities in the white matter connections to the frontal lobe have been reported by several studies, so the functional underconnectivity could well be related to anatomical connectivity properties, as we have demonstrated previously (Cherkassky et al., 2006; Just et al., 2007; Kana et al., 2006; Keller, Kana, & Just, 2007).

In addition to showing lower functional connectivity in autism in the ToM task, the present study also demonstrated a reduction in activation during mentalizing in three key frontal regions: the medial frontal gyrus, the orbitofrontal gyrus, and the anterior paracingulate cortex. This finding is consistent with several previous studies of ToM processing in autism (Baron-Cohen et al., 1999; Castelli et al., 2002; Happé et al., 1996). Mental state attribution presupposes self-awareness and it likely involves processing in evolutionarily recent regions of the neocortex, such as the frontal cortex (Gallup, 1982). As mentioned in the earlier part of the discussion, frontal lobe underactivation in autism is not uncommon in other higher cognitive tasks. It may be possible to distinguish the roles of frontal regions and parietal regions in the ToM network, although the differentiation is not critical to the current study, nor do the current results help to distinguish the roles. Three frontal regions have been credited with distinct roles in mental state attribution. The medial prefrontal cortex has been credited with involvement in any kind of thought that uses the self as a referent (Gusnard, Akbudak, Shulman, & Raichle, 2001), the anterior paracingulate cortex is believed to be involved in the decoupling mechanism that distinguishes mental states from reality (Gallagher & Frith, 2003), and the orbitofrontal cortex is believed to play a crucial role in ToM (Baron-Cohen & Ring, 1994; Brothers & Ring, 1992), and may be associated with recognition of mental state terms (Baron-Cohen et al., 1994). The reduced activation in participants with autism in these regions indicates an atypical neural basis for ToM processing.

Contrary to our prediction, there was no reliable group difference in activation in the right superior temporal sulcus (STS) at the temporoparietal junction while processing ToM. This is in contrast with the findings of reduced activation in autism in this region in a previous study (Castelli et al., 2002). However, the present finding raises several interesting questions. First, in contrast to the implicit nature of the task (passively viewing the animations) in the Castelli et al. (2002) study, the present study included instructions that were designed to require participants to explicitly attempt to attribute mental states to the stimuli (watching animations and answering questions related to mental state). Normal or a higher level of activation in autism has been found in explicit tasks of processing mental states (Wang, Lee, Sigman, & Dapretto, 2006). It is possible that the participants with autism might make a more controlled effort to infer a mental state when they are explicitly asked to do so. Second, it is possible that the superior temporal sulcus plays a subordinate role in inferring mental states. Gallagher and Frith (2003) argued that the STS is more generally associated with social cognition but the dominant roles in ToM are played by the medial prefrontal cortex and the anterior paracingulate cortex. Third, superior temporal activation has been found during the viewing of simple interactions of moving objects (Blakemore et al., 2003; Pelphrey, Morris, Michelich, Allison, & McCarthy, 2005), suggesting that STS activation need not be associated with attribution of mental states. Although all regions that are found to be part of the ToM network play different roles, it may be the coordination and communication between these regions that is crucial in accomplishing ToM. As an adaptation in autism to the reduced connectivity with frontal areas, the posterior superior temporal sulcus might function with greater autonomy, assuming a higher than normal degree of control of the task performance. Thus, the present results are consistent with the idea that participants with autism may process the ToM cues in the same way as controls, but if the results of such processing are not communicated efficiently to frontal regions, then the frontal regions may not be able to carry out the mental state attribution normally. This interpretation would also predict reduced synchronization between the activation in anterior and posterior regions involved in the ToM processing in autism, which is what occurred in this study.

An interesting aspect of activation during ToM processing in autism was that the activation in the right posterior superior temporal sulcus in autism was positively correlated with the ToM score. In other words, people with autism who had higher ToM ability also had greater activation in posterior STS. A correlation between a psychometrically measured ability and an activation level in a relevant brain area has previously been found in autism with respect to other abilities, such as the correlation between the psychometric measure of face processing ability measured by a standardized

test of face perception and activation in the fusiform face area in autism (Koshino et al., 2008; Schultz et al., 2005). Schultz et al. (2005) also found the intactness of social functioning in autism (measured by ADOS) to be correlated with the amount of activation in fusiform gyrus in a face processing task in people with autism, with the better socially functioning autism participants having more fusiform activation.

The absence of a behavioral group difference was unexpected, given previous findings of such differences and given our finding of group differences in brain activation. Note, however, that each participant's behavioral data were based on only three items in each condition. Although these few items were apparently sufficient to provide precise enough estimates to detect within-subject effects of condition, they may have been inadequate for detecting betweengroup differences within the ToM condition. Another possible explanation is that the explicit instructions to attend to the thoughts and feelings of the triangles and the forced choice nature of the response prevented the task from being sensitive to the well-established differences in theory of mind processing between the groups.

The present evidence for functional underconnectivity in autism in a ToM task adds to the converging evidence of functional underconnectivity in a wide range of complex information processing tasks, including language comprehension (Just et al., 2004), working memory (Koshino et al., 2005), executive functioning (Just et al., 2007), complex inhibition (Kana et al., 2007), and mental imagery (Kana et al., 2006). What is common across all these studies is a neural pattern in autism characterized by (1) frontalposterior underconnectivity; (2) less frontal activation; (3) equivalent or greater activation in posterior brain areas; and (4) decrease in size of a relevant portion of the corpus callosum. Disruptions in connectivity between frontal and posterior areas in autism could cause the system to adapt in such a way that the posterior regions work more independently to compensate for the impaired communication with the frontal regions. The phenomenon of frontal-posterior underconnectivity and a consequent increased posterior autonomy has been demonstrated in a computational model of a complex executive task (Just et al., in preparation). In summary, the functional connectivity and brain activation results of the present study indicate a systems-level neural impairment in people with autism. The focal point of this impairment seems to be the decreased communication and coordination between frontal regions and others, affecting ToM processing among other forms of thought.

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150 KANA ET AL.

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#### 152 KANA ET AL.

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