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The Effects of the X Chromosome on Intrinsic Functional Connectivity in the Human Brain: Evidence from Turner Syndrome Patients

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Abstract

Turner syndrome (TS), a disorder caused by the congenital absence of one of the 2 X chromosomes in female humans, provides a valuable human "knockout model" for studying the functions of the X chromosome. At present, it remains unknown whether and how the loss of the X chromosome influences intrinsic functional connectivity (FC), a fundamental phenotype of the human brain. To address this, we performed resting-state functional magnetic resonance imaging and specific cognitive assessments on 22 TS patients and 17 age-matched control girls. A novel data-driven approach was applied to identify the disrupted patterns of intrinsic FC in TS. The TS girls exhibited significantly reduced whole-brain FC strength within the bilateral postcentral gyrus/intraparietal sulcus, angular gyrus, and cuneus and the right cerebellum. Furthermore, a specific functional subnetwork was identified in which the intrinsic FC between nodes was mostly reduced in TS patients. Particularly, this subnetwork is composed of 3 functional modules, and the disruption of intrinsic FC within one of these modules was associated with the deficits of TS patients in math-related cognition. Taken together, these findings provide novel insight into how the X chromosome affects the human brain and cognition, and emphasize an important role of X-linked genes in intrinsic neural coupling.

Key words: functional module, intrinsic functional connectivity, resting state functional MRI, the X chromosome, Turner syndrome

Introduction

In addition to sex determination, the X chromosome is believed to play a crucial role in the development of the human brain and intelligence (Lehrke 1972; Turner 1996; Johnson et al. 2009). X-linked gene defects have been disproportionately observed in various neuropsychiatric disorders, particularly mental retardation (Ropers and Hamel 2005; Skuse 2005). Empirically ascertaining the patterns in which the X chromosome influences and for elucidating sex-specific incidences and symptom presentations for the majority of neuropsychiatric disorders.

In recent years, intrinsic functional connectivity (FC), a measure derived from resting-state functional magnetic resonance imaging (rs-fMRI), has emerged as an effective tool for exploring large-scale human brain organization (Biswal et al. 1995; Buckner et al. 2013). Using this powerful tool, landmark observations such as the default mode network (DMN) have been revealed (Greicius et al. 2003; Raichle 2010). Developmental processes and brain diseases can alter the patterns of intrinsic FC (Greicius 2008; Di Martino et al. 2014). Moreover, intrinsic FC is strongly associated with individual differences in cognitive performance, suggesting a critical role of intrinsic FC in cognition (Hampson et al. 2006; Baldassarre et al. 2012; Wei, Liang, et al. 2012). However, it remains unknown whether and how the X chromosome affects the intrinsic FC of the human brain, and this understanding may provide critical insight into the neural mechanisms underlying X-linked cognitive profiles in health or disease.

Turner syndrome (TS), a disorder in female humans that is characterized by the absence of a normal second X chromosome, serves as a unique human "knockout model" to study X chromosome function in the nervous system (Sybert and McCauley 2004). In TS patients, specific cognitive deficits, such as inferior visuospatial, math, and social cognitive abilities, caused by the loss of the X chromosome have been well documented (Rovet 2004; Hong and Reiss 2012). Neuroanatomically, TS patients have been shown to have reduced parieto-occipital gray matter (GM) volume (Murphy et al. 1993; Reiss et al. 1995; Molko et al. 2004; Marzelli et al. 2011), aberrant thickness and/or surface area of temporal-parieto-occipital cortical regions (Raznahan et al. 2010; Lepage, Clouchoux, et al. 2013; Lepage, Hong, et al. 2013; Lepage, Mazaika, et al. 2013), and impaired microstructural integrity of white matter (WM) tracts such as the superior longitudinal fasciculus (Holzapfel et al. 2006; Yamagata et al. 2012). Furthermore, functional MRI studies have revealed that during visuospatial (Kesler et al. 2004; Bray et al. 2013), executive function (Tamm et al. 2003), working memory (Haberecht et al. 2001; Hart et al. 2006; Bray et al. 2011), or arithmetic tasks (Molko et al. 2003; Kesler et al. 2006), TS patients show abnormal profiles of functional activation or functional coupling/connectivity, predominantly in the frontal/parietal cortices and in subcortical regions such as the caudate. However, currently, the patterns of intrinsic FC in TS patients remain unexplored.

Given the previously observed abnormalities in brain structure and function, we hypothesized that the loss of the X chromosome would result in severe disruption of intrinsic FC patterns in TS, likely affecting a specific functional circuit/network that accounts for the particular cognitive deficits of TS patients. To test this hypothesis, we performed rs-fMRI and obtained a set of cognitive assessments from female TS patients and controls. Using a novel data-driven approach, intrinsic FC was comprehensively analyzed to characterize the disrupted patterns of FC in TS.

Methods and Materials

Participants

Twenty-two TS girls with a nonmosaic 45XO karyotype (age range: 9–18 years) were recruited from the China-Japan Friendship Hospital (CJFH) and Peking Union Medical College Hospital (PUMCH). Age-matched healthy controls (HCs) (21 girls; age range: 10–18 years) were recruited from the local community and parent networks. For each TS patient, the nonmosaic 45XO karyotype was confirmed using a standard cytogenetic assessment of peripheral blood. All of the patients showed defective ovarian development, which was verified via pelvic ultramatrix, 64×64 ; 3.1×3.1 mm in-plane resolution; and 200 volumes in total.

High-resolution T₁-weighted Image

High-resolution 3D T_1 -weighted images were sagittally acquired by using a magnetization prepared rapid gradient echo (MPRAGE) sequence: 144 slices; TE, 3.39 ms; TR, 2530 ms; inversion time (TI), 1100 ms; 1.33 mm slice thickness with no gap; acquisition matrix, 256 × 256; and 1 × 1 mm in-plane resolution.

rs-fMRI Preprocessing

rs-fMRI preprocessing was performed using the pipeline tool Data Processing Assistant for Resting-State fMRI (DPARSF) (Yan and Zang 2010). Briefly, the first 10 volumes were removed due to T_1 equilibration effects and the participants' adaption to the scanner. The remaining volumes were then corrected for slice time differences due to interleaved acquisition, realigned to the first volume to correct for head motion, and normalized to Montreal Neurological Institute (MNI) space. To improve the normalization accuracy, we used a pediatric T₁ template (http://www.bic. mni.mcgill.ca/ServicesAtlases). The age range of this template is 13.0-18.5 years (Fonov et al. 2011), which fits well with the age range of our participants. Specifically, for each participant, the rs-fMRI scan was first coregistered with the native T₁ image that was subsequently normalized to the pediatric template. The rs-fMRI scans were accordingly transformed into the MNI space, such that the image was resampled into 3 mm isotropic voxels. This procedure was implemented in SPM8 (http://www. fil.ion.ucl.ac.uk/spm, Friston et al. 1994). Next, the normalized rs-fMRI scans were spatially smoothed with a 6 mm full-width at half-maximum Gaussian kernel, linearly detrended, and temporally band-pass filtered (0.01-0.08 Hz). As generally recommended, 9 common nuisance variables were regressed out, including 6 head motion parameters (3 translations and 3 rotations along the x, y, and z axes), the global mean signal, the WM signal and the cerebrospinal fluid signal (Fox et al. 2005).

Notably, the data from 2 HCs were excluded because their head motion exceeded 2 mm or 2° in a specific direction. In addition, recent rs-fMRI studies regarding head motion have suggested that mean frame-wise displacement (FD) can be used to effectively estimate micromovement (Power et al. 2012). Individuals with a mean FD of more than 2 SDs from the mean of all of the participants should be excluded (Di Martino et al. 2013). We applied this exclusion criterion in the present study. Accordingly, the imaging data from one HC were discarded. Finally, the rs-fMRI data for 22 TS patients (mean age: 14.2 years; SD: 2.7 years) and 17 HCs (mean age: 14.1 years; SD: 2.3 years) were used for subsequent analysis. Moreover, we used the FD to determine the micromovements of the brain volumes during rs-fMRI scanning. Specifically, volumes with an FD >0.5 mm, along with the immediately preceding volume and 2 subsequent volumes, were considered micromovement-containing volumes (Power et al. 2012; Abrams et al. 2013).

Whole-Brain FC Strength

First, we measured whole-brain functional connectivity strength (wFCS). A GM mask was generated by applying a threshold of 0.1 to the GM probability map of the pediatric template. Then, Pearson correlations of the blood oxygen level-dependent (BOLD) series between every pair of voxels within the GM mask were calculated and converted to Fisher's Z-values, which represent the strength of voxel pair-wise FC in the resting state. For a given voxel, its Z-values with every other voxel were summed together, which was defined as the wFCS value of that voxel (Buckner et al. 2009; Tomasi and Volkow 2010; Wang et al. 2014). Notably, because removing the global BOLD signal during preprocessing can introduce ambiguous negative functional correlations, only positive correlations (i.e., Z-value > 0) were counted for the wFCS, as described previously (Buckner et al. 2009; Liang et al. 2013).

The wFCS values for each voxel were statistically compared between the TS and HC groups, using 2-sample t-tests (two-tailed). Age was included as a covariate. To correct for multiple comparisons, the Monte Carlo simulation method was applied using the "3dClustSim" function of the AFNI package (Cox 1996). Family-wise error (FWE) corrected P values that were <0.05 at the cluster level were considered significant. To evaluate the influence of weak positive correlations (i.e., potential noise) on the results, we further compared the 2 groups using 2 other wFCS maps. Specifically, the 2 additional wFCS maps were calculated after eliminating weak positive correlations, that is, thresholding the voxel pair-wise correlation at 0.1 and 0.2 (R > 0.1 and R > 0.2), respectively.

Identifying Relevant Subnetwork and Functional Modules

For brain clusters showing a significant wFCS group difference, the TS-associated FC alterations were indicated in a wholebrain manner. To further identify specific functional circuits/subnetworks, additional seed-based FC analyses were conducted. Specifically, each of the clusters identified above (6 in total) was taken as a seed. For each seed cluster, the representative BOLD series was extracted as the mean series from a sphere with a radius of 6 mm that was centered on the statistically determined peak voxel (Fox et al. 2005). Pearson correlations of the BOLD series between the seed and every other voxel within the GM mask were computed and converted into Z-values. For each seed cluster, a Z-map of FCS was generated for each subject.

Similarly, a 2-sample t-test that also considered age as a covariate was applied to detect intergroup differences in the Z-map for each seed region. Because only positive correlations were considered in the present study, the group comparison was constrained by a mask based on the voxels that showed a positive mean FC (i.e., Z-value > 0) in the HC group. To correct for multiple comparisons, the method of Monte Carlo simulation was performed using "3dClustSim".

A set of clusters (referred to as target clusters) showed significantly reduced FCS with the seed clusters in the TS patients compared with the HCs. The intrinsic FC of all of these target and seed clusters was putatively affected by the loss of the X chromosome in the TS patients. We therefore selected these clusters to establish a functional subnetwork in which each seed or target cluster represented a node (a total of 31 nodes). A similar approach was applied in a recent study (Gotts et al. 2012). Again, the representative BOLD series for each node/cluster was extracted by averaging the series within a sphere with a radius of 6 mm that was centered on the calculated peak voxel. For each subject, we generated a node-by-node correlation matrix (also converted to a Z-matrix) that represented the functional subnetwork. Within each group, a one-sample t-test (one-tailed) was first applied to all of the node pairs (595 in total) to determine whether the converted Z-value was significantly greater than zero (i.e., significantly positive at the group level). Here, the false discovery rate (FDR) approach was used to correct for multiple comparisons, and q < 0.05 was considered significant. For the between-node

connections that were significantly positive at the group level in either the HC or TS group, we performed a 2-sample t-test to evaluate intergroup differences, and the FDR method was applied to correct for multiple comparisons.

For each group, the average functional subnetwork (i.e., the average Z-matrix) across individuals was obtained, and the Z-values of the node pairs that showed non-significantly positive correlations at the group level were removed (i.e., set to zero). A modular analysis was then performed on the average Z-matrix to explore whether this functional subnetwork could be subdivided into different functional modules. Specifically, a spectral reordering algorithm was applied (Newman 2006).

Correlating Intrinsic FC Changes to Math Ability Deficits in TS

To evaluate whether the observed FC abnormalities could account for the math ability deficits of TS girls, we examined the correlations between the composite math score and the FC parameters across subjects in both the TS and HC groups, adjusting for age. The FC parameters included the voxel-wise wFCS across the entire GM, the wFCS of the identified clusters, the FCS of the connections in the identified subnetwork, and the mean FCS of the identified functional modules.

Results

Demographics and Cognitive Assessment

The demographic information and the results of the cognitive assessment are summarized in Table 1. No significant difference in age was observed between the groups (P = 0.87). The TS patients had significantly lower values for the 5 IQ scores: FSIQ (P = 0.0004), WMI (P = 0.03), VCI (P = 0.01), PRI (P = 0.0004), and PSI (P = 0.0004). The TS patients also scored lower than the HCs on the 3 math tasks (simple subtraction: P = 0.03; number comparison: P = 0.007; and numerosity comparison: P = 0.02) as well as on overall math ability (composite score: P = 0.003).

wFCS Differences

First, several head motion parameters were compared between the TS and HC groups, but no significant difference was observed (maximum head displacement, P = 0.51; maximum head rotation, P = 0.54; mean FD, P = 0.80). Moreover, the number of rs-fMRI brain volumes with micromovement also did not differ significantly between the 2 groups (HC: 9.4 ± 13.5 , range 0–40; TS: 5.4 ± 7.2 , range 0–27; P = 0.24).

On the GM mask, the total wFCS was significantly smaller in the TS group than in the HC group (HC: $0.36 \pm 0.03 \times 10^9$; TS: $0.34 \pm 0.02 \times 10^9$; P = 0.02). The mean wFCS maps for both groups are illustrated at the voxel level in Figure 1. The 2 maps exhibited very similar spatial patterns of voxel-wise wFCS; the highest wFCS values occurred within several DMN regions, including the posterior cingulate gyrus (PCC)/precuneus, the medial prefrontal cortex (MPFC) and the inferior parietal lobule (IPL). The insula and the postcentral gyrus (POCG) also showed relatively high wFCS values. The observed spatial distribution of wFCS was highly consistent with that reported in previous studies (Tomasi and Volkow 2010; Wang et al. 2014).

Visual inspection indicated reduced wFCS values for the TS group in multiple brain regions. The statistical comparison revealed 6 clusters that exhibited significantly reduced wFCS in the TS group compared with the HC group (FWE-corrected P < 0.05, Fig. 1). These 6 clusters were located in (1) the cuneus, (2) the right cerebellum, (3) the right PoCG/intraparietal sulcus (IPS), (4) the left PoCG/IPS, (5) the right angular gyrus (ANG), and (6) the left ANG. The details are summarized in Table 2. The statistical map was closely replicated when the wFCS that eliminated weak correlations was used (Supplementary Figure 1), suggesting that the wFCS thresholding value had a limited effect on our results. Moreover, to evaluate the effect of head micromovements on this result, we removed the micromovement rs-fMRI volumes from the entire time series for each individual and reran the analysis. Statistically, the results remained almost the same (data not shown), indicating a limited effect of head micromovement on our results.

Relevant Subnetworks and Modules

By considering each significant wFCS cluster as a seed, we next performed seed-based FC analysis. For each of the seeds, a set of clusters (referred to as a target cluster) was observed, and the FCS between the seed and the target cluster was significantly reduced in the TS group compared with the HC group (i.e., FWE-corrected P < 0.05). In total, we observed 26 significant target clusters for the 6 seed clusters. We did not find any significant target clusters with increased FCS in the TS patients. The identified target

Table 1 Demographic characteristics and cognitive assessment performance

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clusters were primarily observed within the superior parietal lobule (SPL)/IPL and surrounding areas, the PCC/precuneus, the MPFC/lateral prefrontal cortex (LPFC), the middle/inferior temporal cortex (MTC/ITC), the pericalcarine visual cortex, and the cerebellum (Fig. 2 and Table 2). Overall, the identified target clusters exhibited a bilaterally symmetrical pattern despite a few unilaterally distributed target clusters. The 2 seed clusters in the left and right ANG exhibited a very similar spatial distribution of target clusters. Notably, some target clusters from one seed cluster spatially overlapped with other seed clusters or with target clusters from another seed cluster, indicating circuit-like relationships between these clusters (Fig. 2 and Table 2).

Taken together, we identified 31 separate nodes (6 seed clusters and 25 target clusters). The seed cluster in the left PoCG/IPS highly overlapped with the target cluster from the seed cluster in the right PoCG/IPS (and mapped to identical MNI coordinates); therefore, they were counted as a single node. For each individual, a 31×31 Pearson correlation matrix was generated. The average Z-matrices for both groups are illustrated in Figure 3.

Furthermore, the spectral reordering algorithm revealed 3 modules for the average Z-matrix in the HC group. Within each module, the nodes were densely connected. However, very few connections were detected between modules. Notably, the modular organization was apparent only in the reordered matrices, as illustrated in Figure 3. To simplify interpretation, the average Z-matrix for the TS group was reordered in terms of the modules that were identified in the HC group.

Specifically, Module I was composed of 3 vision-related nodes (i.e., the seed cluster in the cuneus and 2 related target clusters). Module II contained 12 nodes, including 3 seed clusters (the right cerebellum and the left and right PoCG/IPS) and their related target clusters. The remaining 16 nodes formed Module III, which consisted of 2 seed clusters (i.e., the left and right ANG) and their related target clusters. These details are presented in Table 2.

As illustrated in Figure 3, the TS group visually showed less or weaker FC in each of the 3 modules. Given the node selection criteria described above, the reduction of FC between the seed cluster and its corresponding target clusters was expected, but it remains unknown whether there are FCS differences in the connections between different seed or target clusters. The results presented here also revealed significant reductions in FC between many different seed or target clusters. The reduced FCS values were evenly distributed across the 3 modules, without showing a preference for any specific module. Notably, a few increases in FC between modules were observed.

Functional Module Disruption versus Math Ability Deficits in TS

The voxel-wise correlation between the wFCS and composite math score across the entire GM showed no significant cluster after multiple comparison correction (i.e., FWE-corrected P

Table 2 Clusters/ROIs showir	g differences	in resting-state	functional co	onnectivity in 🕻	ΓS girls
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Cluster/ROIs		MNI coordinates of the cluster peak			Volume (mm ³)	Modular index
		Х	Y	Ζ		
1	Cuneus	-6	-78	18	2403	1
2	Pericalcarine occipital cortex	-27	-63	-18	26 568	1
3	Superior/middle occipital cortex	27	-78	18	2916	1
4	Right cerebellum	21	-72	-54	3132	2
5	Left supramarginal gyrus	-63	-36	39	4239	2
6	Right anterior fusiform gyrus	27	-6	-39	3105	2
7	Left anterior fusiform gyrus/inferior temporal gyrus	-33	-6	-42	2565	2
8	Right postcentral gyrus/Intraparietal sulcus	39	-30	42	12 123	2
9	Left postcentral gyrus/Intraparietal sulcus	-45	-36	39	11 151	2
9	Left postcentral gyrus/Intraparietal sulcus	-45	-36	39	2538	2
10	Right postcentral gyrus/intraparietal sulcus	30	-39	48	15 093	2
11	Right posterior superior/middle frontal gyrus	30	0	66	8613	2
12	Right SPL	27	-57	69	6831	2
13	Right posterior inferior frontal gyrus	51	15	33	6183	2
14	Left posterior superior/middle frontal gyrus	-27	0	48	4158	2
15	Left anterior supramarginal gyrus	-63	-27	39	4023	2
16	Right angular gyrus	51	-54	33	1998	3
17	Medial/LPFC	-33	66	0	57 537	3
18	Left middle/inferior temporal gyrus	-63	-21	15	17 955	3
19	Right cerebellum-crus	18	-90	-27	12 096	3
20	Precuneus/PCC	9	-42	36	10 125	3
21	Left angular gyrus/intraparietal sulcus	-45	-66	57	9018	3
22	Right middle/inferior temporal gyrus	72	-30	-9	8532	3
23	Right ventrolateral prefrontal cortex	39	54	-15	6507	3
24	Left angular gyrus	-45	-57	33	2592	3
25	Medial/lateral prefrontal cortex	6	51	21	38 853	3
26	Left middle temporal gyrus	-54	15	-33	13 554	3
27	Posterior cingulate gyrus/precuneus	-3	-30	42	10 179	3
28	Right temporal pole	45	21	-30	6804	3
29	Caudate	-12	9	12	4617	3
30	Right angular gyrus	54	-51	27	4401	3
31	Cerebellum-vermis	6	-57	-42	3510	3

Note: The 6 seed clusters are marked in bold (i.e., the 1st, 4th, 8th, 9th, 16th, and 24th clusters), followed by their respective target clusters. Notably, the target cluster for the seed cluster of the right postcentral gyrus/intraparietal sulcus strongly overlapped with the seed cluster of the left postcentral gyrus/intraparietal sulcus (the ninth cluster); therefore, they were considered as a single node. The modular indexes 1, 2, and 3 correspond to Modules I, II, and III in Figure 3. MNI, Montreal Neurological Institute.

FC patterns in the human brain. Compared with the HCs, the TS patients had significantly decreased wFCS within the bilateral PoCG/IPS, ANG, and cuneus as well as the right cerebellum. Furthermore, a specific functional subnetwork was identified in which intrinsic FC between regions was typically reduced in the TS patients. This subnetwork could be divided into 3 functional modules, and the disruption of intrinsic FC within one of these modules was associated with the math-related cognitive deficits of the TS patients in. Together, these results elucidate the effect of the X chromosome on functional coupling in the human brain and provide important information to understand X-linked brain diseases and sex differences in the brain, specifically with respect to cognition.

The X chromosome, which comprises approximately 4% of the human genome, has long been considered to be critical for the development of the human brain and intelligence (Lehrke 1972; Turner 1996; Johnson et al. 2009). TS patients naturally lose one of their 2 normal X chromosomes; therefore, these individuals serve as a valuable human "knockout model" for studying how the X chromosome affects the human brain and cognition. A number of MRI studies have been conducted on TS patients, but they have primarily focused on structural brain anomalies or functional changes during specific cognitive tasks (Mullaney and Murphy 2009). The current investigation is the first study to explore the intrinsic FC patterns in TS, and our results may provide novel understanding to elucidate the function of the X chromosome in the nervous system. Further, the current study provides direct evidence of an X-linked genetic basis for intrinsic FC, which is of particular value for the identification of genes involved in this fundamental phenotype of the human brain.

It is important to note that X-linked genes can affect the brain in at least 2 ways: by directly acting on the brain and by indirectly acting on the gonads to induce differences in specific gonadal secretions (i.e., hormones) that exert specific effects on the brain (Arnold 2004). To isolate the direct genetic effect from the indirect hormonal effect, one possible approach is to ensure identical hormonal levels across individuals with different X-linked genotypes. Although identical hormone levels between adolescent TS patients and healthy controls are difficult to achieve in practice, a suboptimal alternative is to match the pubertal stage, as an approximation of the sex hormone level, between groups. In our cohort, however, pubertal status was not well matched between the



Figure 2. The target clusters showing a significant reduction in intrinsic FC with their seed clusters. (A) The target clusters for the seed region of the cuneus. (B) The target clusters for the seed region of the right cerebellum. (C) The target clusters for the seed region of the right PoCG/IPS. (D) The target clusters for the seed region of the right ANG. (F) The target clusters for the seed region of the left ANG. The seed clusters were indicated by the purple balls and blue arrows. The colors indicate the T values for the intergroup differences. L, left; R, right.



Figure 3. The relevant subnetwork and functional modules associated with the loss of the X chromosome in TS patients. 46XX and 45XO represent the karyotypes of HC and TS girls, respectively. (A) The relevant subnetwork and functional modules for the HC group. (B) The relevant subnetwork and functional modules for the TS group. In the 3D network rendering, the nodes within Modules I, II, and III are shown in yellow, red, and green, respectively. The colors in the matrix represent the mean Z values for the intrinsic FCS across the subjects. Modules I, II, and III are indicated by yellow, red and green rectangles, respectively, in the matrix. (C) The intergroup differences in intrinsic FC within the subnetwork and the modules. The colors in the matrix represent the T-values for the intergroup differences. The number for each row of the matrices correspond to the index of clusters/regions in Table 2. The 3D brain rendering was implemented using the Brainnet Viewer (Xia et al. 2013).

2 groups. The majority of our TS patients were prepubertal because spontaneous puberty is very rare in TS girls, and most of our TS patients did not undergo ER therapy to artificially induce puberty. Therefore, the intrinsic FC patterns observed in the TS patients may be due to a direct genetic factor, an indirect hormonal factor, or a combination of the 2. Future studies that employ animal models are essential to dissociate these 2 effects (Arnold and Chen 2009; Raznahan et al. 2013).



Figure 4. The correlation of math ability with the mean FCS of Module II in the subnetwork. Module II is indicated by a red rectangle in the matrix.

Hub Regions of Disrupted Intrinsic FC

In the present study, intrinsic FC was measured by analyzing the correlations between BOLD signals in a resting state (Biswal et al. 1995; Buckner et al. 2013). To avoid a priori selection of specific regions of interest (ROIs), we used a data-driven approach to map the wFCS at the voxel level and identified the regions that showed aberrant FCS in the TS patients. Similar approaches have been applied to study mental dysfunction (Gotts et al. 2012; Wang et al. 2014) and to search for connectivity hubs within the brain (Buckner et al. 2009; Tomasi and Volkow 2010).

A set of regions including the bilateral IPS, ANG, cuneus, and cerebellum showed reduced wFCS in the TS patients compared with the HCs. Widespread FCS interruptions between these regions and other region across the entire brain are expected. Clearly, in terms of disrupted FC, these regions represent the most affected, or hub regions, and the FCS of these regions is strongly associated with the loss of the X chromosome. Consistently, these regions have been showed to exhibit structural abnormalities in TS. For instance, aberrant morphology, including abnormal sulcal geometry, cortical thickness, and GM volume, was observed for the IPS in TS females (Raznahan et al. 2010; Marzelli et al. 2011; Lepage, Clouchoux, et al. 2013; Molko et al. 2003). In addition, reduced cortical surface area and GM volume in the ANG and cuneus (Reiss et al. 1995; Marzelli et al. 2011; Xie et al. 2015), as well as increased GM volume in the cerebellum (Cutter et al. 2006; Hong, Hoeft, et al. 2014), have been found in TS. In line with these findings, the ANG and the cuneus also display sex differences in cortical thickness (Sowell et al. 2007), further supporting a function of the X chromosome in these regions.

Aside from the structural anomalies caused by the loss of the X chromosome, these hub regions display abnormal neuronal activity during the performance of cognitive tasks. During specific arithmetic tasks, the IPS, ANG and cuneus showed weaker functional activation in TS patients than in HCs (Molko et al. 2003; Kesler et al. 2006). Moreover, decreases in functional activation or BOLD signal changes were found within the IPS during working memory or visuospatial tasks (Haberecht et al. 2001; Kesler et al. 2004; Hart et al. 2006; Bray et al. 2011, 2013). In particular, during these tasks, abnormal FC seeding from the IPS has been observed (Bray et al. 2011, 2013). It is possible that for the IPS, the disrupted intrinsic FC in the resting state we observed serves

as a baseline abnormality that contributes to the abnormal FC detected during various tasks.

The disrupted intrinsic FC of these regions observed in the TS patients should at least partially correlate with the structural and task-based functional abnormalities described above. However, the relationship between these factors might be very complex, and importantly, it is unclear whether these factors are causally associated with each other. Unraveling these relationships is critical for clarifying the functional pathway by which the X chromosome affects these regions. Future studies using multimodal imaging techniques would be useful for addressing this issue.

Disrupted Functional Modules and Cognition

Based on the observed hub regions, we further identified a subnetwork in which the majority of the region/node pairs displayed reduced intrinsic FC in the TS patients compared with HCs. Importantly, this subnetwork is composed of 3 functional modules. Modular structure is a commonly observed feature of structural and functional brain networks (Chen et al. 2008; He et al. 2009). Such structure provides a balance between the 2 most fundamental principles of brain organization, functional segregation and integration, and facilitates efficient recurrent processing within modules and information exchange between modules (Sporns et al. 2000, 2004; Kotter and Stephan 2003). Within each module, a set of regions/nodes is densely connected, putatively allowing them to work together to perform certain functions. In TS patients, all 3 observed modules showed reduced intrinsic FC.

Intriguingly, the mean FCS of Module II was correlated with overall math ability, and therefore, the connectivity disruption of this module likely accounts for the math cognitive deficits of the TS patients relative to the controls. Notably, this significant correlation was based on the entire sample, and there was no significant correlation within either the TS (R = -0.11, P = 0.64) or the control group (R = 0.21, P = 0.44), suggesting that the FCS of Module II could not account for the within-group variance in math ability. The lack of significant within-group correlation might be due to the very small sample size, the narrow range, or the small variance of math scores within each group. The significant correlation across the entire sample suggests that the X chromosome might influence math cognition by modulating the

intrinsic FC of a math-specific functional module. Module II may serve as a backbone for such a math-specific functional module. Specifically, Module II consisted of 11 regions predominantly from the parietal cortex, including the IPS, the SPL and the supramarginal gyrus. These regions, particularly the IPS, have been well recognized to be involved in math-related cognitive processing (Molko et al. 2003; Wei et al. 2014), supporting the current results regarding Module II and math ability.

The other 2 modules were not correlated with the math cognitive scores, indicating the specificity of Module II for math cognition. The other 2 modules are likely associated with other cognitive functions. For instance, Module I included 3 visual regions that are located close together in the occipital cortex and that are highly involved in visuospatial cognition. Therefore, the disrupted connectivity of Module I may underlie the deficits of TS patients in visuospatial ability, another well-observed cognitive feature caused by loss of the X chromosome (Rovet 2004). In contrast, Module III consisted of 16 regions that are widely distributed across the brain; these regions are likely involved in multiple functional systems. For example, several regions, including the PCC/precuneus, the MPFC, and the ANG, are considered to be key nodes of the DMN (Fox et al. 2005). These DMN regions, together with other regions in Module III such as the middle/inferior temporal gyrus and the temporal poles, have been recognized as key nodes for social cognitive processing (Kennedy and Adolphs 2012). Accordingly, Module III may be associated with social cognition, and it is possible that the disruption of Module III is related to the social cognitive deficits of TS patients (Hong and Reiss 2012; Hong, Bray, et al. 2014). However, these interpretations of the cognitive functions of Modules I and III are essentially speculative. In the present study, the relevant cognitive tests were not performed on our participants and therefore this study did not provide the data needed to evaluate these speculations. This issue must be addressed in future studies.

To some extent, the reduced FC in our observed modules reflects a decreased efficiency of the underlying anatomical connectivity. This interpretation is supported by previous findings showing a reduction in WM integrity in TS patients (Molko et al. 2004; Holzapfel et al. 2006; Yamagata et al. 2012; Xie et al. 2015). However, some evidence clearly suggests that intrinsic FC is not simply a proxy for the underlying anatomical connectivity (Buckner et al. 2013). To elucidate the biological mechanisms underlying the intrinsic FC finding presented here, specific investigations at the molecular or cellar level are warranted.

Notably, the functional modules that were identified in the present study were extracted using a data-driven approach based on whole-brain FC. Due to our constraint of analyzing only the TS-affected FC, each of the functional modules might be incomplete. Alternatively, a set of ROIs based on previous fMRI studies of math, visuospatial, or social cognition could be applied as modular nodes, which would result relatively complete functional modules for these cognitive functions. However, the challenge of this type of approach is the selection of appropriate ROIs, particularly for cognitive functions such as math that are less studied and for which relatively mixed fMRI results have been obtained in humans.

TS-related Math Deficits in Response Time or Accuracy

Previous studies have suggested that the math deficits in TS are mainly related to slower processing in math-related tasks rather than to overall accuracy (Bruandet et al. 2004; Mazzocco 2006). To evaluate whether the current math-related results were mainly attributed to response time or accuracy in the math-related tasks, we calculated separate composite scores for response time and accuracy. The 2 composite scores showed a significant difference or trend between the TS and control groups (t-test): composite response time, P = 0.043; composite accuracy,1

Supplementary Material

Supplementary material can be found at: http://www.cercor. oxfordjournals.org/.

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