Wolff et al. Molecular Autism (2017) 8:8 DOI 10.1186/s13229-017-0126-z

Molecular Autism

RESEARCH Open Access

CrossMark

Neural circuitry at age 6 months associated with later repetitive behavior and sensory responsiveness in autism

Jason J. Wolff^{1*}, Meghan R. Swanson², Jed T. Elison³, Guido Gerig⁴, John R. Pruett Jr.⁵, Martin A. Styner⁶, Clement Vachet⁷, Kelly N. Botteron⁵, Stephen R. Dager⁸, Annette M. Estes⁹, Heather C. Hazlett^{2,6}, Robert T. Schultz¹⁰, Mark D. Shen², Lonnie Zwaigenbaum¹¹, Joseph Piven^{2,6} and The IBIS Network

Abstract

Background: Restricted and repetitive behaviors are defining features of autism spectrum disorder (ASD). Under revised diagnostic criteria for ASD, this behavioral domain now includes atypical responses to sensory stimuli. To date, little is known about the neural circuitry underlying these features of ASD early in life.

Methods: Longitudinal diffusion tensor imaging data were collected from 217 infants at high familial risk for ASD. Forty-four of these infants were diagnosed with ASD at age 2. Targeted cortical, cerebellar, and striatal white matter pathways were defined and measured at ages 6, 12, and 24 months. Dependent variables included the Repetitive Behavior Scale-Revised and the Sensory Experiences Questionnaire.

Results: Among children diagnosed with ASD, repetitive behaviors and sensory response patterns were strongly correlated, even when accounting for developmental level or social impairment. Longitudinal analyses indicated that the genu and cerebellar pathways were significantly associated with both repetitive behaviors and sensory responsiveness but not social deficits. At age 6 months, fractional anisotropy in the genu significantly predicted repetitive behaviors and sensory responsiveness at age 2. Cerebellar pathways significantly predicted later sensory responsiveness. Exploratory analyses suggested a possible disordinal interaction based on diagnostic status for the association between fractional anisotropy and repetitive behavior.

Conclusions: Our findings suggest that restricted and repetitive behaviors contributing to a diagnosis of ASD at age 2 years are associated with structural properties of callosal and cerebellar white matter pathways measured during infancy and toddlerhood. We further identified that repetitive behaviors and unusual sensory response patterns co-occur and share common brain-behavior relationships. These results were strikingly specific given the absence of association between targeted pathways and social deficits.

Keywords: Infant, Diffusion tensor imaging, Autism, Repetitive behavior, White matter, Longitudinal

Background

Restricted and repetitive behaviors (RRBs) are defining characteristics of autism spectrum disorder (ASD). Behaviors comprising this domain range from relatively simple topographies—such as motor stereotypies—to more complex forms including inflexible adherence to routines and intense, circumscribed interests. There is

evidence that RRBs in toddlerhood are early emerging, prognostic features [1] that may differentiate infants who do and do not later develop ASD [2, 3]. Separately, neuroimaging studies of infant siblings of children with ASD, who are themselves at elevated risk for the disorder, indicate that atypical features and trajectories of brain development may be evident as early as 6 months of age in children who later receive a diagnosis [4–7]. Although such brain changes occur in parallel with emerging patterns of atypical RRBs, the specific aspects

Full list of author information is available at the end of the article



^{*} Correspondence: jjwolff@umn.edu

¹Department of Educational Psychology, University of Minnesota, Minneapolis, MN, USA

Page 2 of 12

of brain development underlying their emergence in infancy is unknown.

Preclinical work has implicated components of cortico-striatal-thalamo-cortical circuitry [8] as the neurobiological basis of RRBs [9–11]. Specific to individuals with ASD, there is supporting evidence linking RRBs with connectivity and morphology of this system in children [12–15] and adults [16–18]. Despite some consistency across studies, the direction of effects within the same structures or circuits is mixed. For example, striatal volumes have been reported to be positively [13, 14] and negatively [12, 19] correlated with repetitive behavior. What appear as inconsistent findings may instead reflect phenotypic heterogeneity or developmental effects, wherein the role of striatal circuits and structures in the etiology and maintenance of RRBs is not static across subpopulations or over time [13, 14, 20].

Sensorimotor processing and motor control are also supported by the cerebellum [21–23]. Studies implicating the cerebellum in ASD extend back over two decades [24–26], and this structure has been linked to RRBs associated with the disorder in humans and nonhuman animal models [18, 27–29]. Repetitive behavior has been inversely correlated with cerebellar volume among adults with ASD [18], and cerebellar hypoplasia has been linked to stereotypy and decreased environmental exploration in children [28]. Similar results have been reported more recently with the addition of significant positive associations between RRBs and vermis grey matter [27].

Recently updated nosology of ASD under DSM 5 includes for the first time symptoms related to unusual behavioral responses to, or interests in, sensory stimuli [30] as part of restricted and repetitive behaviors. While this change reflects clinical consensus, the conceptual grouping of RRBs with unusual responses to sensory stimuli in ASD is supported by a limited body of empirical and theoretical work addressing the relationship between these features of ASD [30–33]. There is evidence that as with RRBs, atypical sensory responses are evident in toddlerhood [34, 35] and are linked to patterns of neural connectivity in adolescence [36, 37]. While not yet empirically tested, it has been hypothesized that early cerebellar dysfunction may also explain the range of sensory response patterns observed in ASD [38, 39].

Our primary aim was to examine the structural properties of select neural circuits in relation to RRBs and sensory responsiveness in familial high-risk infants who developed ASD. Our analyses focused on pathways connecting brain regions implicated by previous studies of RRBs, including: (1) thalamo-cortical and cortico-striatal circuitry, (2) ponto-cerebellar and cerebello-thalamic circuitry, and (3) anterior corpus callosum [5, 12–19, 27–29]. We posited that RRBs and sensory response

patterns would covary in toddlers with ASD. Proceeding from this, we investigated: (1) whether development of targeted white matter pathways, measured from infancy through toddlerhood using diffusion tensor magnetic resonance imaging (DT-MRI), would be associated with RRBs and sensory responsiveness at age 2 years; (2) whether variation in white matter pathways at age 6 months would be associated with later RRBs or sensory responsiveness at age 2 years; and (3) whether classes of behavior (i.e., repetitive behaviors and sensory responsiveness) were characterized by overlapping versus distinct brain-behavior relationships. As a follow-up to these aims, we examined whether observed effects extended to high-risk infants who did not receive a diagnosis of ASD.

Methods

Participants

Participants were from the Infant Brain Imaging Study, a prospective, longitudinal study of infants at high and low familial risk for ASD. Familial high-risk status was defined by having an older sibling with a community diagnosis of the ASD confirmed by the Autism Diagnostic Interview-(ADI-R) [40] and Social Communication Questionnaire [41]. Infants were enrolled at one of four clinical data collection sites: the Children's Hospital of Philadelphia, University of North Carolina, University of Washington, and Washington University in St. Louis. Exclusion criteria were (1) evidence of a genetic condition or condition affecting development; (2) significant vision or hearing impairment; (3) birth weight < 2000 g or gestational age < 36 weeks; (4) significant perinatal adversity or prenatal exposure to neurotoxins, (5) contraindication for MRI, (6) predominant home language other than English, (7) adopted, half siblings, or twins and (8) first degree relative with psychosis, schizophrenia, or bipolar disorder.

The present sample included high-risk children who met the following criteria: (1) at least one complete DT-MRI scan and (2) complete cognitive and behavioral assessment battery at age 2 years including diagnostic evaluation and assessment of RRBs. Our primary sample of interest were high-risk infants who received a clinical best-estimate diagnosis of ASD at age 2 years (HR-ASD; n = 44). We also included high-risk infants who did not receive a diagnosis of ASD at age 2 (HR-Neg; n = 173) to discern whether effects observed among HR-ASD extended to unaffected children with shared familial liability for ASD. Clinical best-estimate diagnoses were made based upon DSM-IV-TR criteria using all available assessment data including the Autism Diagnostic Observation Schedule (ADOS) [42], ADI-R, Mullen Scales of Early Learning (MSEL) [43], and the Vineland Adaptive Behavior Scales II [44]. Reliability for these standardized instruments was initially established and maintained

Page 3 of 12

between sites through monthly case reviews. Clinical best-estimate diagnoses made at age 2 years were independently confirmed by a second senior clinician blind to risk status and diagnosis made by the first clinician. All participants in our study sample had a completed ADOS administration. MSEL data were incomplete for two participants (1 HR-ASD, 1 HR-Neg), and composite scores from age 12 months were substituted. Although the parent study also collects data on low-familial risk infants, this group was excluded from the present study given lack of variance in RRB scores due to floor effect [3]. Written informed consent was obtained for all participants from their parent or guardian, and all study procedures were approved by institutional review boards at each clinical site (Children's Hospital of Philadelphia, University of North Carolina, University of Washington at St. Louis, and Washington University).

Clinical measures

Assessment data included the Repetitive Behavior Scale-Revised (RBS-R) [45] and the Sensory Experiences Questionnaire v2.1 (SEQ) [46] administered at age 2 years. The RBS-R is a measure of RRBs consisting of 43 items, each of which represents a discrete behavioral topography. The RBS-R is sensitive to individual differences among toddlers at high-risk for ASD [3], and the measure has been independently validated for use in children as young as age 2 years [42]. We partitioned total inventory scores from the RBS-R into two categories: *lower-order* (combining stereotypical, self-injurious, and restricted behaviors), and *higher-order* (combining compulsive, ritualistic, and sameness behaviors). This grouping was made upon the basis of conceptual and factor analytic work [47, 48].

The SEQ consists of 38 items assessing responses to sensory stimuli across modalities. In addition to a total score, the SEQ yields summary scores for items representing hypo- and hyper-responsivity and sensory seeking. The SEQ has been shown to have strong psychometric properties [46, 49]. Complete SEQ data was available for 89% (n = 39) of the HR-ASD sample and 83% (n = 144) of the HR-Neg sample. The MSEL Early Learning Composite (MSEL composite) [43] was used to characterize general developmental level. Social affect, RRB, and total severity scores were derived from the Autism Diagnostic Observation Schedule (ADOS) [42]. Descriptive data for the study sample are provided in Table 1.

MRI data acquisition and processing

MRI scans were acquired on 3T Siemens TIM Trio scanners equipped with 12-channel head coils during natural sleep. The imaging protocol included: sagittal T1 MP-Rage (TR = 2400 ms, TE = 3.16 ms, slice thickness = 1 mm, FOV = 256 mm, 256×160 matrix), 3D T2 fast

Table 1 Descriptive data for study sample

Variable	HR-ASD	HR-Neg	P^{a}
Six-month sample	32	106	
Total longitudinal sample	44	173	
Longitudinal scan complemen	nt		
6, 12, and 24 m scan	15	54	
6 and 12 m scan	7	28	
6 and 24 m scan	6	11	
12 and 24 m scan	7	37	
6 m scan	4	13	
12 m scan	3	22	
24 m scan	2	8	
Mean age time 1	6.4 (0.4)	6.7 (0.7)	.08
Mean age time 2	12.8 (0.7)	12.6 (0.6)	.16
Mean age time 3	24.5 (0.7)	24.7 (0.7)	.92
Sex (% male)	89	58	<.001
ADOS repetitive behavior	2.7 (1.8)	0.7 (0.9)	<.001
ADOS social affect	12.0 (3.8)	2.3 (2.2)	<.001
ADOS severity score	6.0 (1.9)	1.6 (1.0)	<.001
MSEL composite	79.3 (16.9)	102.1 (15.8)	<.001
RBS-R total items endorsed	6 (0–38) ^b	2 (0-27) ^b	<.001
SEQ total raw score	51.7 (12.6)	39.8 (7.6)	<.001

^aTest statistic for independent samples t test except for sex (Fisher's exact test) and RBS-R total items endorsed (Mann-Whitney U test)

^bMedian and range

ADOS Autism Diagnostic Observation Schedule, MSEL Mullen Scales of Early Learning, RBS-R Repetitive Behavior Scale-Revised, SEQ Sensory Experiences Questionnaire

spin echo (TR = 3200 ms, TE = 499 ms, slice thickness = 1 mm, FOV = 256 mm, 256×160 matrix), and 25-direction ep2d_diff with FoV = 190 mm (6 and 12 months) or FoV = 209 mm (24 months), 75–81 transversal slices, slice thickness = 2 mm isotropic, $2 \times 2 \times 2$ mm3 voxel resolution, TR = 12,800–13,300 ms, TE = 102 ms, variable *B* value 0–1000 s/mm². Intra- and inter-site reliability was established and maintained across sites and time [50].

Diffusion-weighted images were processed with DTI-prep, which detects common artifacts, corrects for motion and eddy current deformations, and flags bad gradients for manual removal by expert raters [51, 52]. Data sets with fewer than 18 (of 25 total) gradients were excluded from further processing to ensure consistent signal-to-noise ratio. Overall, approximately 10.5% of acquired DTI datasets were excluded following quality control procedures, with about one quarter of excluded cases due to subject motion. An additional 4% of all DWI scans were excluded for incomplete acquisition (e.g., child awoke during scan). There were no differences between HR children with and without a diagnosis of ASD in proportion of scans excluded. Group analysis of diffusion weighted data utilized an atlas-based

processing pipeline providing consistent spatial parameterization within and between individual datasets across ages [53, 54].

Deterministic fiber tractography was performed by the first and second author in the study specific, average atlas space using 3D Slicer (https://www.slicer.org/) and refined via FiberViewerLight. Fiber tract definitions followed anatomically informed methods [55]. Pathways of interest (Fig. 1) included the cortical-spinal tract (CST), which passes through the basal ganglia to the motor-sensory cortex; anterior thalamic radiation (ATR), which passes from thalamus to pre-frontal cortex; genu of the corpus callosum, which connects frontal hemispheres; mid-cerebellar peduncle (MCP), which passes through the pons and connects cerebellar hemispheres; and the superior cerebellar peduncles (SCP), which connect the cerebellum and thalamus.

Fractional anisotropy (FA) values were obtained via DTIAtlasFiberAnalyzer[53]. FA represents degree of diffusion along the primary fiber orientation (axial diffusivity) relative to transverse diffusion (radial diffusivity). Following previous infant DTI studies [56, 57], bilateral pathways were averaged to yield a single estimate. All components of this processing pipeline are freely available as part of the UNC/Utah NAMIC DTI Fiber Analysis Framework [53].

Statistical analysis

Because RBS-R and SEQ scores were non-normally distributed, Spearman's rank order correlations were used to examine strength of association between RRBs and sensory responsiveness. Correlation analyses were performed using bootstrap sampling with replacement (B = 1000) to generate confidence interval estimates. Follow-up analyses were conducted with control for possible confounding

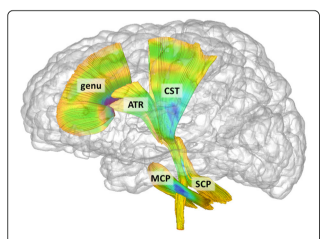


Fig. 1 White matter fiber pathways defined through tractography of DTI data. Targeted pathways include *ATR* anterior thalamic radiation, *CST* cortico-spinal tract, *genu* genu of corpus callosum, *MCP* mid-cerebellar peduncle, *SCP* superior cerebellar peduncle

variables: ADOS social affect and MSEL composite. Correlation analyses were uncorrected for multiple comparisons with $\alpha = .05$.

We next examined longitudinal imaging data across ages 6, 12, and 24 months in relation to these behaviors using generalized estimating equations (GEE). Predictors included fractional anisotropy of target white matter pathways, sex, and age. MSEL composite score was assessed as a potential covariate and ultimately excluded from our analytic models as this variable did not contribute to model fit and was not itself significantly associated RBS-R or SEQ scores. Sex and age were included as covariates in all models given potential effects on DTI data [54]. The interaction of FA and age was tested for each model to determine whether differences in slope explained the relationship between FA and dependent variables. Secondary analysis was performed using subscales derived from the RBS-R and SEQ and all targeted fiber pathways to fully explore possible brain-behavior relationships.

Models were fit based upon response distribution and measurement type of the primary dependent variables. For analyses of RRBs, a negative binomial regression model with log link function was fit with total RBS-R inventory as the dependent variable. This approach was selected given that the RBS-R inventories constituted integer count data. Goodness-of-fit statistics indicated that a negative binomial distribution was superior to a Poisson distribution. For sensory responsiveness, a gamma regression model with log link function was fit with SEQ total raw score as the dependent variable. This approach was selected given that SEQ data followed a log-normal distribution. Goodnessof-fit statistics indicated that this was superior to regression based on a Gaussian distribution. Secondary analyses were performed on subscale measures derived from the RBS-R and SEQ and included higher- and lower-order repetitive behaviors (from RBS-R) and hypo- and hyper-responsiveness and sensory seeking (from SEQ) as dependent variables following significant primary model results. Exponentiated beta (e^{B}) coefficients were generated to provide an estimate of effect size (incident rate ratio).

Following longitudinal analyses, we next employed generalized linear modeling (GLM) to focus specifically on the predictive relationship of white matter pathways measured at age 6 months with RRBs and sensory responsiveness measured at age 2 years. These models were controlled for the effects of sex and age and were fit based on the response distributions and link functions described above. In a final set of exploratory analyses, we replicated the steps described above in high-risk infants who did not meet diagnostic criteria for ASD at age 2 years (HR-Neg) to determine if effects observed in

Page 5 of 12

HR-ASD extended to unaffected siblings. An adaptive false-discovery rate procedure was applied at each stage of primary and secondary GEE and GLM analyses to account for multiple comparisons [58]. Adjusted q values are presented with the significance threshold at q < 0.05. Supplementary analyses were conducted to explore the differential roles of axial and radial diffusivities, as well as possible lateralization effects (see Additional files). Results from supplementary and exploratory analyses are presented without correction for multiple comparisons.

Results

Concurrence of repetitive behavior and sensory response patterns

At 24 months, RBS-R scores correlated strongly with total SEQ, $r_{\rm s}$ = 0.74, p < 0.001, 95% CI [0.55, 0.87], supporting our hypothesis that RRBs and sensory responsiveness would covary in the HR-ASD group. This correlation held up to partial models controlling for MSEL composite (pr_s = 0.75, p < 0.001) and ADOS social affect (pr_s = 0.74, p < 0.001). Correlations of lower- and higher-order RRBs derived from the RBS-R with patterns of sensory responsiveness from the SEQ (hypo, hyper, and seeking) were all statistically significant with the exception of the association between higher-order RRB and sensory seeking behavior (see Table 2).

Longitudinal brain development in relation to behavior at age 2

To expand on the results from age 6 months, we next examined development of FA in target pathways across 6, 12, and 24 months in HR-ASD in relation to total RBS-R and SEQ. These results are presented in Table 3. For the RBS-R, there were significant main effects for the genu, MCP, and SCP over the 6 to 24-month interval. These effects were largely consistent across lowerand higher-order RRBs derived from the RBS-R (Table 4). There were no significant effects for the ATR or CST.

Table 2 Nonparametric correlations for subscales derived from the RBS-R and SEQ among HR-ASD

	SEQ total	SEQ hypo	SEQ hyper	SEQ Sensory Seeking		
RBS-R lower-order						
$r_{\rm s}$	0.75***	0.67***	0.57***	0.53**		
95% CI	0.57, 0.86	0.45, 0.80	0.30, 0.74	0.24, 0.74		
RBS-R higher-order						
$r_{\rm s}$	0.53**	0.50**	0.39*	0.21		
95% CI	0.12, 0.74	0.18, 0.71	0.04, 0.65	-0.14, 0.54		

RBS-R Repetitive Behavior Scale-Revised, SEQ Sensory

Table 3 Longitudinal model results for effect of fractional anisotropy on total RBS-R and SEQ among HR-ASD

Variable	χ^2	q	e^{B}	CI
Total RBS-R ¹				
ATR FA	3.2	0.06	1.07	0.99-1.16
CST FA	0.2	0.31	1.02	0.94-1.10
Genu FA	16.3	< 0.001	1.14	1.07-1.22
MCP FA	8.3	0.004	1.07	1.02-1.12
SCP FA	7.0	0.006	1.12	1.03-1.22
Total SEQ ¹				
ATR FA	0.9	0.18	1.01	0.99-1.04
CST FA	0.0	0.38	1.00	0.98-1.02
Genu FA	13.0	< 0.001	1.04	1.02-1.06
MCP FA	9.5	0.003	1.02	1.01-1.04
SCP FA	6.8	0.006	1.03	1.01-1.06

Generalized estimating equation model results

RBS-R Repetitive Behavior Scale-Revised, SEQ Sensory Experiences Questionnaire, ATR anterior thalamic radiation, CST cortico-spinal tract, SCP superior cerebellar peduncle, MCP midcerebellar peduncle

q false-discovery rate adjusted p value, e^{B} exponentiated regression coefficient representing percent of increase in score for every hundredth unit increase in fractional anisotropy, Cl 95% confidence interval

For total SEQ scores, there were significant main effects for the genu, MCP, and SCP (see Table 3). For subscales derived from the SEQ, there were significant effects for hypo-responsiveness in relation to the genu and MCP (Table 4). While initial analyses suggested that these pathways were likewise significantly associated with hyper-responsiveness, these results did not survive FDR correction. There were no significant main effects for the CST or ATR, and no significant effects for any pathway with Sensory Seeking. There were no additional significant effects for the interaction of Age X FA for any pathway. See Additional file 1: Table A1 for supplementary longitudinal analyses of axial and radial diffusivities.

Specificity of longitudinal results

To assess the specificity of our findings, we first examined a control behavioral measure of a domain which we reasoned qualitatively differed from repetitive behaviors or sensory responsiveness. Namely, we conducted longitudinal analyses using social affect scores from the ADOS as our dependent variable. This revealed no significant relationships with any pathways associated with the RBS-R or SEQ (Genu FA: $\chi^2 = 1.7$, p = 0.19; MCP FA: $\chi^2 = 0.0$, p = 0.91; SCP FA: $\chi^2 = 0.3$, p = 0.59).

We next examined FA in pathways not included in our primary hypotheses to assess regional specificity. This analysis included the anterior limb of the internal capsule (ALIC) and splenium of the corpus callosum. Both have been previously implicated in imaging studies

Experiences Questionnaire

^{***}p < 0.001, **p < 0.01, *p < .05

Page 6 of 12

Table 4 Secondary longitudinal analysis results for fractional anisotropy and behavior for HR-ASD

Variable	χ^2	9	e^{B}	Cl
RBS-R				
Lower order				
ATR FA	1.2	0.33	1.05	0.96-1.14
CST FA	0.0	0.84	1.01	0.93-1.09
Genu FA	5.7	0.05	1.09	1.02-1.17
MCP FA	6.2	0.048	1.07	1.01-1.12
SCP FA	6.7	0.04	1.13	1.03-1.23
Higher order				
ATR FA	4.2	0.08	1.10	1.00-1.20
CST FA	0.6	0.42	1.04	0.95-1.13
Genu FA	25.3	< 0.001	1.21	1.12-1.30
MCP FA	6.9	0.04	1.08	1.02-1.14
SCP FA	4.4	0.07	1.11	1.01-1.23
SEQ				
Нуро				
ATR FA	0.7	0.42	1.01	0.98-1.05
CST FA	0.0	0.86	1.00	0.97-1.03
Genu FA	14.3	0.002	1.05	1.02-1.07
MCP FA	11.5	0.007	1.03	1.01-1.05
SCP FA	5.0	0.07	1.04	1.01-1.07
Hyper				
ATR FA	0.7	0.42	1.01	0.99-1.04
CST FA	0.1	0.75	1.00	0.98-1.03
Genu FA	4.5	0.07	1.02	1.00-1.05
MCP FA	2.1	0.24	1.01	1.00-1.03
SCP FA	4.9	0.07	1.03	1.00-1.06
Seeking				
ATR FA	1.3	0.33	0.98	0.93-1.02
CST FA	1.1	0.35	0.98	0.94-1.02
Genu FA	1.6	0.31	1.02	0.99-1.06
MCP FA	1.3	0.33	1.02	0.99-1.04
SCP FA	2.6	0.19	1.04	0.99-1.09

Results for generalized estimating equations

RBS-R Repetitive Behavior Scale-Revised, SEQ Sensory Experiences Questionnaire, ATR anterior thalamic radiation, CST cortico-spinal tract, SCP

Questionnaire, ATR anterior thalamic radiation, CST cortico-spin superior cerebellar peduncle, MCP midcerebellar peduncle

of infants at risk for ASD [6, 59] but not specifically in relation to RRBs or sensory responsiveness. The ALIC was not significantly associated with total RBS-R or SEQ scores (all q > 0.18). We did identify a significant main effect for the splenium and total RBS-R score ($\chi^2 = 6.7$, q = 0.08, $e^{\rm B} = 1.07$) but not for SEQ total score ($\chi^2 = 2.6$, q = 0.17, $e^{\rm B} = 1.02$).

FA at age 6 months in relation to behavior at age 2

Generalized linear model results for HR-ASD are presented in Table 5. Genu FA at age 6 months was significantly associated with total RBS-R at age 2, (χ^2 = 17.5, q < 0.001, e^B = 1.24). For the genu, e^B coefficients indicated that each 0.01 unit increase in FA was associated with an estimated 23% increase in total number of RBS-R items endorsed at age 2. Genu FA was significantly associated with lower-order (χ^2 = 5.7, q = 0.02, e^B = 1.16) and higher-order (χ^2 = 32.4, q < 0.001, e^B = 1.36) behaviors from the RBS-R. Each unit increase in genu FA was associated with an estimated 16 and 34% increase in count of lower- and higher-order RRBs. The other four pathways of interest were not significantly associated with total scores on the RBS-R.

FA in the genu ($\chi^2 = 14.0$, q < 0.001, $e^B = 1.06$) at age 6 months was significantly associated with total SEQ score at age 2 years. Each unit increase in genu FA was associated with an estimated 6% increase in SEQ score. The MCP ($\chi^2 = 4.9$, p = 0.02, $e^B = 1.02$) was also associated with later SEQ score, though this result did not survive FDR correction (q = 0.056). Genu FA was significantly associated with sensory hypo-responsivity on the SEQ ($\chi^2 = 10.0$, q = .002, $e^B = 1.08$). Hyperresponsivity scores were also significantly associated with FA in the genu ($\chi^2 = 4.9$, q = 0.014, $e^B = 1.04$). Genu FA was not significantly associated with sensory seeking scores. To provide visualization of RBS-R and SEQ data in relation to FA measures, example scatterplots showing the genu and CST are provided in Additional file 2:

Table 5 Fractional anisotropy at age 6 months predicting behavior at age 2 years among HR-ASD

Variable	χ^2	9	e^{B}	CI
Total RBS-R				
ATR	0.6	0.47	1.05	0.92-1.20
CST	0.2	0.59	1.03	0.91-1.18
Genu	17.5	< 0.001	1.24	1.12-1.37
SCP	0.8	0.47	1.06	0.94-1.19
MCP	1.2	0.41	1.04	0.97-1.11
Total SEQ				
ATR	2.9	0.73	1.03	1.00-1.06
CST	1.1	0.59	1.02	0.98-1.05
Genu	14.0	< 0.001	1.06	1.03-1.09
SCP	2.1	0.31	1.02	0.99-1.06
MCP	4.9	0.06	1.02	1.00-1.04

Generalized linear model results

RBS-R Repetitive Behavior Scale-Revised, SEQ Sensory Experiences Questionnaire, ATR anterior thalamic radiation, CST cortico-spinal tract, SCP superior cerebellar peduncle, MCP midcerebellar peduncle

q false-discovery rate adjusted p value, e^{B} exponentiated regression coefficient representing percent of increase in score for every hundredth unit increase in fractional anisotropy, Cl 95% confidence interval

q false-discovery rate adjusted p value, e^{g} exponentiated regression coefficient representing percent of increase in score for every hundredth unit increase in fractional anisotropy, Cl 95% confidence interval

Page 7 of 12

Figure S1. See Additional file 1: Table A2 for supplementary analyses of axial and radial diffusivity.

Unaffected high-risk siblings

For HR-Neg toddlers (n=137), total RBS-R scores significantly correlated with total SEQ, $r_{\rm s}=0.49$, p<0.001, 95% CI [0.35, 0.60]. This relationship remained when controlling for MSEL composite score (${\rm pr}_{\rm s}=0.47$, p<0.001) or social affect score (${\rm pr}_{\rm s}=0.48$, p<0.001). Correlations of lower- and higher-order RRB scores derived from the RBS-R with patterns of sensory responsiveness from the SEQ (hypo, hyper, and seeking) were all statistically significant (see Additional file 1: Table A3).

Results from longitudinal analyses of HR-Neg participants are presented in Table 6. There were no significant effects observed for FA in relation to total RBS-R or SEQ scores after correction for multiple comparisons. However, the uncorrected model and the effect size estimate for the genu did suggest a possible association $(p = .035, \text{ uncorrected}; e^{B} = 0.94, 95\% \text{ CI } [0.89, 0.99]).$ There were no additional significant effects for the interaction of Age X FA for any pathway. Follow-up analyses conducted with the ALIC and splenium revealed no significant effects for the ALIC in relation to RBS-R or SEQ total scores (all q > 0.25). Splenium FA was not significantly associated with total SEQ scores (q = 0.99). As with the genu, uncorrected model results suggested that splenium FA was significantly associated with total RBS-R, though this result did not survive FDR correction (χ^2 = 7.6, q = 0.08, $e^{B} = 0.95$, 95% CI [0.91, 0.99]).

Table 6 Longitudinal model results for HR siblings not meeting diagnostic criteria for ASD

Variable	χ^2	9	e^{B}	CI
Total RBS-R				
ATR FA	0.2	0.91	0.99	0.92-1.05
CST FA	0.8	0.66	0.97	0.91-1.04
Genu FA	4.5	0.25	0.94	0.89-0.99
MCP FA	1.5	0.46	0.98	0.94-1.01
SCP FA	2.1	0.46	0.95	0.88-1.02
Total SEQ ⁴				
ATR FA	1.8	0.46	1.01	1.00-1.02
CST FA	0.0	0.99	1.00	0.99-1.01
Genu FA	0.3	0.91	1.00	0.99-1.01
MCP FA	0.2	0.91	1.00	0.99-1.01
SCP FA	2.6	0.46	1.01	1.00-1.02

Results for generalized estimating equations

ATR anterior thalamic radiation, CST cortico-spinal tract, SCP superior cerebellar peduncle, MCP midcerebellar peduncle, q false-discovery rate adjusted p value, e^B exponentiated regression coefficient representing percent of increase in score for every hundredth unit increase in fractional anisotropy; CI 95% confidence interval, RBS-R Repetitive Behavior Scale-Revised, SEQ Sensory Experiences Questionnaire

Exploratory analysis of brain and behavior interaction among high-risk infants

Although FDR corrected results for HR-Neg indicated no statistically significant results, we were nonetheless intrigued that the direction of effects for HR-Neg was the reverse of those observed in HR-ASD. We therefore conducted a limited set of exploratory analyses to more fully interrogate the effect size estimates observed for the genu and splenium of the corpus callosum in relation to total RBS-R scores.

First, we explored RBS-R results for the genu and splenium in HR-Neg by examining lower- and higher-order RRBs derived from the RBS-R. For the genu, we found that the relationship was strongest for lower-order ($\chi^2 = 11.7$, p = 0.001, $e^B = 0.90$, 95% CI [0.84, 0.96]) versus higher-order behaviors ($\chi^2 = 1.0$, p = 0.34, $e^B = 0.97$, 95% CI [0.92, 1.03]). Results were similar for the splenium: lower-order ($\chi^2 = 8.3$, p = 0.004, $e^B = 0.93$, 95% CI [0.89, 0.98]) and higher order $\chi^2 = 6.2$, p = 0.01, $e^B = 0.95$, 95% CI [0.91, 0.99].

As a direct test of the inverse relationship of genu/ splenium FA and repetitive behavior scores observed between HR-ASD and HR-Neg groups, we examined the interaction of diagnostic group and genu/splenium FA on total RBS-R scores in generalized estimating equation models that included both HR-ASD and HR-Neg participants controlling for sex and age. The interaction of diagnostic group on FA was significant for the genu ($\chi^2 = 4.3$, p = 0.039) and splenium ($\chi^2 = 4.8$, p = 0.029). The interaction was also significant at the six-month time point in relation to later RBS-R score for the genu ($\chi^2 = 10.0$, p =0.002) but not splenium ($\chi^2 = 3.3$, p = 0.07). Visualizations of these interactions are provided in Fig. 2. To determine whether interaction effects were specific to repetitive behavior in relation to corpus callosum FA, we expanded our analysis to include all targeted pathways as well as SEQ scores in a longitudinal model including both HR-ASD and HR-Neg. For RBS-R score, we observed an additional significant interaction for SCP FA ($\chi^2 = 4.0$, p = 0.046). For the SEQ scores, we observed a significant interaction between diagnostic status and genu FA ($\chi^2 = 4.8$, p = 0.029), with effects for the MCP and SCP just above the critical alpha value (p = 0.067 and 0.076). All other interactions were not statistically significant (see Additional file 1: Table A6). When diagnostic status was not accounted for in the above models, there were no significant predictive effects of brain on behavior.

Given the inverse relationship observed between brain and behavior by diagnostic outcome, as a final step we explored whether FA was predictive of diagnostic status. At age 6 months, FA in the ATR, CST, Genu, and MCP did not significantly predict diagnostic outcome (all p > 0.10). However, diagnostic outcome was significantly predicted

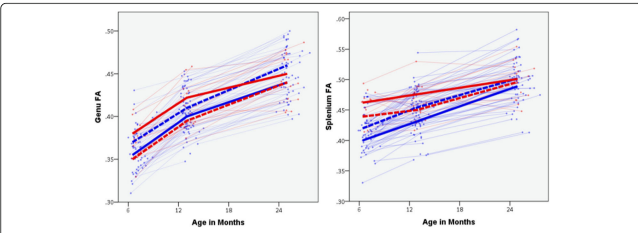


Fig. 2 Trajectories of genu and splenium FA development by ASD diagnosis. *Red* signifies high-risk children diagnosed with ASD at age 2. *Blue* signifies high-risk children without ASD at age 2. *Bolide blue* lines represent *upper* and *lower* quartile groups based on total RBS-R score at age 2. *Bolide blue* lines represent upper quartile of RBS-R scores. Quartile groups are presented solely to provide a visualization of the interaction effect between diagnostic status and FA on repetitive behavior measured at age 2, wherein higher FA was associated with more repetitive behavior for children with ASD, but was associated with less repetitive behavior for children without a diagnosis

by FA at age 6 months for the splenium (χ^2 = 18.4, p < 0.001, e^B = 1.44, 95% CI [1.22, 1.71; PPV = 65%, NPV = 84%]) and SCP (χ^2 = 20.2, p < 0.001, e^B = 1.81, 95% CI [1.40, 2.35]; PPV = 68%, NPV = 85%). Consistent with previous results [6], these findings suggest that higher FA at age 6 months is associated with increased risk for a later diagnosis of ASD.

Discussion

We found that structural properties of cerebellar and corpus callosum white matter in infancy were associated with later-emerging restricted and repetitive behaviors (RRBs) and responsiveness to sensory stimuli in familial high-risk infants who developed ASD. Effects were uniformly characterized by a positive association between FA and symptom severity at age 2. The brain-behavior relationships we observed appeared specific to RRBs and sensory responsiveness given the absence of any association with social deficits. We also identified a strong positive association between RRBs and sensory response patterns measured at age 2 that remained even when controlling for cognitive ability or social symptom severity. Together these results suggest alignment between these classes of behavior in both outward manifestation and in underlying neurobiology early in the course of ASD.

To follow-up our primary results, we extended our analyses to high-risk infants who did not receive a diagnosis of ASD. As with HR-ASD, we observed that RRBs were significantly correlated with patterns of sensory responsiveness. We did not observe any statistically significant brain-behavior relationships after correction for multiple comparisons. However, effect size estimates suggested that genu and splenium FA may be associated with later RRBs in the HR-Neg group. Curiously, the

direction of this effect was reversed, with FA negatively associated with repetitive behaviors at age 2 in HR-Neg. Given this unusual result, we conducted a limited set of exploratory analyses, the results of which suggest that diagnostic status may exert a moderating effect on the relationship between FA and RRB. This adds to previous reports of disordinal interactions in brain and behavior between infants who did and did not receive a diagnosis of ASD [4, 5]. For example, we have previously identified a disordinal interaction between infants who developed ASD and typically developing controls in the relationship of visual orienting latency to structural connectivity in the splenium [4]. Moreover, higher FA associated with a diagnosis of ASD early in life has been reported by several independent groups [5, 6, 60-63]. The preliminary data reported here further suggest that white matter development in infants with ASD may be fundamentally altered relative to unaffected children. As opposed to its connotation in DTI studies of typically developing children, higher FA may reflect pernicious neurodevelopmental effects unique to autism during infancy. It is plausible that higher FA in babies who develop ASD reflects underlying differences in axon structure related to axon refinement, caliber, or packing density. Further work is required to determine the timing of this phenomenon and whether a subgroup of children is driving it. It will also be necessary to explore alternative explanations, such as the possibility that effects observed through DTI are secondary to some unaccounted variable.

Our findings align with previous work identifying a role for the cerebellum in repetitive behavior [18, 27–29] and sensory processing [64, 65]. The absence of a strong association between RRBs with striatal pathways overall

Page 9 of 12

was counter to what we expected based upon previous work and supports the specificity of our findings. In our view there, are two plausible explanations. The first involves potential limitations. Our ability to measure more nuanced pathways within the basal ganglia was limited by imaging resolution and immaturity of the infant brain. DTI data is but one means of pursuing brain-behavior relationships, and the absence of signal using this modality does not preclude other approaches revealing clearer striatal involvement.

With regard to the second possibility, it is plausible that cerebellar circuitry supports nascent restricted and repetitive behaviors in early childhood, with striatal circuits coming online as such behaviors become entrenched through experience [19, 66, 67] or as laterdeveloping behaviors emerge. Indeed, much of what is known about the role of the striatum comes from data involving older children [12–15], adults [16–18], or adult animal models [10]. It is less clear what role striatal circuitry plays during infant development or how such circuitry might interact with cerebellar circuits during this period to support emerging function [39], such as the transition from reflexive to instrumental behavior [67]. There is some evidence that the cerebellum and basal ganglia co-develop with motor skill acquisition during infancy [68] and that both share crucial roles in processing output from the cortex —mutually supportive roles which are substantially less segregated than once thought [22, 69].

Cerebellar and striatal circuits innervate similar frontal cortical regions [70, 71], which are in turn connected by the genu of the corpus callosum. The indirect involvement of the genu in both cerebellar and striatal circuits may in part explain our finding that genu FA strongly predicts RRBs and sensory responsiveness. However, we did find some evidence implicating the splenium as well, suggesting a possibly more general role for the corpus callosum in these emerging behaviors. Though less often the focus of such studies, there is precedence for corpus callosum involvement with RRBs associated with ASD [5] and other psychiatric conditions [72, 73]. There is also the strong possibility that our findings reflect biobehavioral mechanisms specific to early development. That is, the corpus callosum may play a unique functional role during infancy, complicating attempts to interpret its involvement through the lens of adult models. A direct test of this premise requires longitudinal follow-up into later childhood to characterize how the brainbehavior relationships observed herein change over time.

Owing to revised diagnostic criteria, we were interested in the co-occurrence of sensory responsiveness and RRBs among toddlers with ASD. We identified strong correlations between these behaviors even with social symptom severity or general developmental level

held constant. This adds to a limited body of evidence supporting a relationship between RRBs and sensory responsiveness [30-32]. As conventionally defined, RRBs outwardly share in common two defining elements: they are relatively invariant in topography and occur repeatedly. The manner in which such behaviors are linked to atypical responses to sensory stimuli is a matter which requires further study. Do measures of these behaviors reflect different aspects of some primary phenomenon? Do repetitive behaviors and unusual responses to sensory stimuli belong together within a plausible, falsifiable theoretical framework [33]? Are existing concepts and terminology related to sensory responsiveness in ASD overly reductionist? While the present results suggest substantial overlap between RRBs and patterns sensory responsiveness in relation to one another and to underlying neurobiology, these critical questions ultimately remain unresolved.

There are several limitations which merit discussion. Our analyses relied on parent report measures, which provide an indirect assessment of behavior. Alternative measures may yield more accurate data and allow inclusion of a low-risk contrast group. Future work might also go beyond course divisions of behavioral domains, such as lower/higher-order RRBs, to examine finegrained brain-behavior relationships. In this study, bilateral fiber pathways were averaged to yield a single estimate. While supplementary analyses indicated a general absence of lateralization, there were some marginal which may warrant closer investigation (Additional file 1: Tables A4-A5). The present analyses were hypothesis-driven and were focused on a specific set of white matter pathways. Follow-up work might elect to further vet regions such as the corpus callosum and cerebellum as well as consider alternate regions. Although diagnoses made at age 2 years show strong stability over time [1, 7], ASD is a heterogeneous disorder and we expect the present findings may be representative only of samples enrolled and assessed with similar procedures [74]. Following infants past early childhood might afford the opportunity to better characterize developmental phenotypic heterogeneity and chart dynamic brain-behavior relationships over time. Lastly, our exploratory analyses into possible interaction effects among all HR participants were limited to a specific set of white matter pathways and behavioral measures. A more comprehensive investigation into potential interactions between diagnostic status and FA development on RRBs and other behaviors core to ASD is needed.

Conclusions

We observed that restricted and repetitive behaviors and unusual responses to sensory stimuli co-occur in toddlers with ASD and that these behaviors are associated

Page 10 of 12

with the structural properties of callosal and cerebellar white matter circuits measured during infancy and toddlerhood. These brain-behavior relationships were remarkably specific, suggesting a possible neurobiological mechanism wherein atypical neural development in infancy precedes the emergence of core autistic features within the domain of restricted and repetitive behaviors. Identifying pre-symptomatic markers of later behavior in ASD also affords the possibility of developing enhanced approaches to screening and preventative interventions.

Additional files

Additional file 1: Table A1. Longitudinal model results for axial and radial diffusivities with repetitive behavior and sensory responsiveness, HR-ASD. Table A2 Axial and radial diffusivities at age 6 months predicting repetitive behavior and sensory responsiveness measured at age 2 years, HR-ASD. Table A3 Nonparametric correlations for subscales derived from the RBS-R and SEQ among HR-Neg. Table A4 Longitudinal model results for fractional anisotropy of bilateral pathways with repetitive behavior and sensory responsiveness, HR-ASD. Table A5 Fractional anisotropy of bilateral pathways at age 6 months predicting repetitive behavior and sensory responsiveness measured at age 2 years, HR-ASD. Table A6 Interaction of diagnostic status and fractional anisotropy on behavior. (DOCX 27 kb)

Additional file 2: Figure S1. Scatterplots of FA in select white matter pathways at age 6 months with RBS-R and SEQ scores. Scatterplots show relationship of FA in two pathways measured at age 6 months (genu, cortico-spinal tract) with total Repetitive Behavior Scale-Revised (RBS-R) and Sensory Experiences Questionnaire (SEQ) scores at age 24 months. Panels **a** and **c** were selected to provide visualization of a significant relationship between brain-and behavior in children with ASD (HR-ASD, in red). Panels **b** and **d** provide examples of a non-significant relationship (RBS-R and SEQ, respectively). High-risk children without ASD (HR-Neg) are shown in blue. Note that linear trend lines are for visualization purposes only and were not the basis of statistical modeling. (TIF 191 kb)

Abbreviations

ADOS: Autism Diagnostic Observation Schedule; ASD: Autism spectrum disorder; ATR: Anterior thalamic radiation; CST: Cortical spinal tract; DT-MRI: Diffusion tensor magnetic resonance imaging; FA: Fractional anisotropy; GEE: Generalized estimating equation; GLM: Generalized linear model; HR: High risk; MCP: Midcerebellar peduncle; RBS-R: Repetitive Behavior Scale-Revised; RRBs: Restricted and repetitive behaviors; SCP: Superior cerebellar peduncle; SEQ: Sensory Experiences Questionnaire

Acknowledgements

We wish to express our gratitude to Rachel G. Smith for assisting with data processing and quality control. Thanks to our IBIS families for continued participation in this research. Portions of the data reported here were presented at the 2016 International Meeting for Autism Research. Group author information: the Infant Brain Imaging Study (IBIS) Network is an NIH funded Autism Center of Excellence project and consists of a consortium of eight universities in the U.S. and Canada. Clinical sites: University of North Carolina: J. Piven (IBIS Network PI), H.C. Hazlett, C. Chappell; University of Washington: S. Dager, A. Estes, D. Shaw; Washington University: K. Botteron, R. McKinstry, J. Constantino, J. Pruett; Children's Hospital of Philadelphia: R. Schultz, S. Paterson; University of Alberta: L. Zwaigenbaum; University of Minnesota: J. Elison; Data Coordinating Center: Montreal Neurological Institute: A.C. Evans, D.L. Collins, G.B. Pike, V. Fonov, P. Kostopoulos; S. Das; Image Processing Core: New York University: G. Gerig; University of North Carolina: M. Styner; Statistical Analysis Core: University of North Carolina: H. Gu.

Funding

This research was supported by the National Institutes of Health under awards K01MH101653, P30HD03110, R01HD055741, R01MH093510, T32HD40127, U54EB005149, and U54HD079124; Autism Speaks, and the Simons Foundation. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

Availability of data and materials

Datasets analyzed for the current study are available through the National Database for Autism Research, https://ndar.nih.gov/

Authors' contributions

JW and JP contributed to study design, data processing and analysis, data interpretation, and manuscript preparation. MRS, GG, MAS, and CV contributed to data processing and analysis, data interpretation, and manuscript preparation. JTE, JRP, MDS, and LZ participated in data interpretation and manuscript preparation. KNB, SRD, AME, HCH, and RTS participated in data collection and manuscript preparation. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Consent for publication

Not applicable.

Ethics approval and consent to participate

Study procedures were approved by the institutional review boards at each clinical site (Children's Hospital of Philadelphia, University of North Carolina, University of Washington at St. Louis, and Washington University). Written informed consent was obtained for all participants from their parent or legal quardian.

Author details

¹Department of Educational Psychology, University of Minnesota, Minneapolis, MN, USA. ²Carolina Institute for Developmental Disabilities, University of North Carolina, Chapel Hill, NC, USA. ³Institute of Child Development, University of Minnesota, Minneapolis, MN, USA. ⁴Tandon School of Engineering, New York University, New York City, NY, USA. ⁵Department of Psychiatry, Washington University School of Medicine, St. Louis, MO, USA. ⁶Department of Psychiatry, University of North Carolina, Chapel Hill, NC, USA. ⁷Scientific Computing and Imaging Institute, University of Utah, Salt Lake City, UT, USA. ⁸Department of Radiology, University of Washington, Seattle, WA, USA. ⁹Department of Speech and Hearing Sciences, University of Washington, Seattle, WA, USA. ¹⁰Center for Autism Research, Children's Hospital of Philadelphia, Philadelphia, PA, USA. ¹¹Department of Pediatrics, University of Alberta, Edmonton, AB, Canada.

Received: 25 August 2016 Accepted: 25 February 2017 Published online: 04 March 2017

References

- Lord C, Risi S, DiLavore PS, Shulman C, Thurm A, Pickles A. Autism from 2 to 9 years of age. Arch Gen Psychiatry. 2006;63:694–701.
- Elison JT, Wolff JJ, Reznick JS, Botteron KN, Estes AM, Gu H, Hazlett HC, Meadows AJ, Paterson SJ, Zwaigenbaum L, Piven J. Repetitive behavior in 12-month-olds later classified with autism spectrum disorder. J Am Acad Child Adolesc Psychiatry. 2014;53:1216–24.
- Wolff JJ, Botteron KN, Dager SR, Elison JT, Estes AM, Gu H, Hazlett HC, Pandey J, Paterson SJ, Schultz RT, Zwaigenbaum L, Piven J. Longitudinal patterns of repetitive behavior in toddlers with autism. J Child Psychol Psychiatry. 2014;55:945–53.
- Elison JT, Paterson SJ, Wolff JJ, Reznick JS, Sasson NJ, Gu H, Botteron KN, Dager SR, Estes AM, Evans AC, Gerig G, Hazlett HC, Schultz RT, Styner M, Zwaigenbaum L, Piven J. White matter microstructure and atypical visual orienting in 7-month-olds at risk for autism. Am J Psychiatry. 2013;170:899– 908.
- Wolff JJ, Gerig G, Lewis JD, Soda T, Styner MA, Vachet C, Botteron KN, Elison JT, Dager SR, Estes AM, Hazlett HC, Schultz RT, Zwaigenbaum L, Piven J.

- Altered corpus callosum morphology associated with autism over the first 2 years of life. Brain. 2015;138(Pt 7):2046–58.
- Wolff JJ, Gu H, Gerig G, Elison JT, Styner M, Gouttard S, Botteron KN, Dager SR, Dawson G, Estes AM, Evans AC, Hazlett HC, Kostopoulos P, McKinstry RC, Paterson SJ, Schultz RT, Zwaigenbaum L, Piven J. Differences in white matter fiber tract development present from 6 to 24 months in infants with autism. Am J Psychiatry. 2012;169:589–600.
- Shen MD, Nordahl CW, Young GS, Wootton-Gorges SL, Lee A, Liston SE, Harrington KR, Ozonoff S, Amaral DG. Early brain enlargement and elevated extra-axial fluid in infants who develop autism spectrum disorder. Brain. 2013;136(Pt 9):2825–35.
- Alexander GE, DeLong MR, Strick PL. Parallel organization of functionally segregated circuits linking basal ganglia and cortex. Annu Rev Neurosci. 1986;9:357–81.
- Rothwell PE, Fuccillo MV, Maxeiner S, Hayton SJ, Gokce O, Lim BK, Fowler SC, Malenka RC, Südhof TC. Autism-associated neuroligin-3 mutations commonly impair striatal circuits to boost repetitive behaviors. Cell. 2014;158:198–212.
- Ahmari SE, Spellman T, Douglass NL, Kheirbek MA, Simpson HB, Deisseroth K, Gordon JA, Hen R. Repeated cortico-striatal stimulation generates persistent OCD-like behavior. Science. 2013;340:1234–9.
- Canales JJ, Graybiel AM. A measure of striatal function predicts motor stereotypy. Nat Neurosci. 2000;3:377–83.
- Estes A, Shaw DWW, Sparks BF, Friedman S, Giedd JN, Dawson G, Bryan M, Dager SR. Basal ganglia morphometry and repetitive behavior in young children with autism spectrum disorder. Autism Res. 2011;4:212–20.
- Wolff JJ, Hazlett HC, Lightbody AA, Reiss AL, Piven J. Repetitive and selfinjurious behaviors: associations with caudate volume in autism and fragile X syndrome. J Neurodev Disord. 2013;5:12.
- Langen M, Bos D, Noordermeer SDS, Nederveen H, van Engeland H, Durston S. Changes in the development of striatum are involved in repetitive behavior in autism. Biol Psychiatry. 2014;76:405–11.
- Di Martino A, Kelly C, Grzadzinski R, Zuo X-N, Mennes M, Mairena MA, Lord C, Castellanos FX, Milham MP. Aberrant striatal functional connectivity in children with autism. Biol Psychiatry. 2011;69:847–56.
- Sears LL, Vest C, Mohamed S, Bailey J, Ranson BJ, Piven J. An MRI study of the basal ganglia in autism. Prog Neuropsychopharmacol Biol Psychiatry. 1999;23:613–24.
- Hollander E, Anagnostou E, Chaplin W, Esposito K, Haznedar MM, Licalzi E, Wasserman S, Soorya L, Buchsbaum M. Striatal volume on magnetic resonance imaging and repetitive behaviors in autism. Biol Psychiatry. 2005; 58:226–32.
- Rojas DC, Peterson E, Winterrowd E, Reite ML, Rogers SJ, Tregellas JR. Regional gray matter volumetric changes in autism associated with social and repetitive behavior symptoms. BMC Psychiatry. 2006;6:56.
- Langen M, Schnack HG, Nederveen H, Bos D, Lahuis BE, de Jonge MV, van Engeland H, Durston S. Changes in the developmental trajectories of striatum in autism. Biol Psychiatry. 2009;66:327–33.
- Tanimura Y, King MA, Williams DK, Lewis MH. Development of repetitive behavior in a mouse model: roles of indirect and striosomal basal ganglia pathways. Int J Dev Neurosci. 2011;29:461–7.
- Koziol LF, Budding D, Andreasen N, D'Arrigo S, Bulgheroni S, Imamizu H, Ito M, Manto M, Marvel C, Parker K, Pezzulo G, Ramnani N, Riva D, Schmahmann J, Vandervert L, Yamazaki T. Consensus paper: the cerebellum's role in movement and cognition. Cerebellum. 2014;13:151–77.
- 22. Bostan AC, Dum RP, Strick PL. The basal ganglia communicate with the cerebellum. Proc Natl Acad Sci U S A. 2010;107:8452–6.
- Mosconi MW, Mohanty S, Greene RK, Cook EH, Vaillancourt DE, Sweeney JA. Feedforward and feedback motor control abnormalities implicate cerebellar dysfunctions in autism spectrum disorder. J Neurosci. 2015;35:2015–25.
- Ritvo ER, Freeman BJ, Scheibel AB, Duong T, Robinson H, Guthrie D, Ritvo A. Lower Purkinje cell counts in the cerebella of four autistic subjects: initial findings of the UCLA-NSAC Autopsy Research Report. Am J Psychiatry. 1986:143:862–6.
- Bailey A, Luthert P, Dean A, Harding B, Janota I, Montgomery M, Rutter M, Lantos P. A clinicopathological study of autism. Brain. 1998;121(Pt 5):889–905.
- Fatemi SH, Aldinger KA, Ashwood P, Bauman ML, Blaha CD, Blatt GJ, Chauhan A, Chauhan V, Dager SR, Dickson PE, Estes AM, Goldowitz D, Heck DH, Kemper TL, King BH, Martin LA, Millen KJ, Mittleman G, Mosconi MW, Persico AM, Sweeney JA, Webb SJ, Welsh JP. Consensus paper: pathological role of the cerebellum in autism. Cerebellum. 2012;11:777–807.

- D'Mello AM, Crocetti D, Mostofsky SH, Stoodley CJ. Cerebellar gray matter and lobular volumes correlate with core autism symptoms. NeuroImage Clin. 2015;7:631–9.
- 28. Pierce K, Courchesne E. Evidence for a cerebellar role in reduced exploration and stereotyped behavior in autism. Biol Psychiatry. 2001;49:655–64.
- Tsai PT, Hull C, Chu Y, Greene-Colozzi E, Sadowski AR, Leech JM, Steinberg J, Crawley JN, Regehr WG, Sahin M. Autistic-like behaviour and cerebellar dysfunction in Purkinje cell Tsc1 mutant mice. Nature. 2012; 489-647. E1
- Boyd BA, Baranek GT, Sideris J, Poe MD, Watson LR, Patten E, Miller H. Sensory features and repetitive behaviors in children with autism and developmental delays. Autism Res. 2010;3:78–87.
- Wigham S, Rodgers J, South M, McConachie H, Freeston M. The interplay between sensory processing abnormalities, intolerance of uncertainty, anxiety and restricted and repetitive behaviours in autism spectrum disorder. J Autism Dev Disord. 2015;45:943–52.
- Chen Y-H, Rodgers J, McConachie H. Restricted and repetitive behaviours, sensory processing and cognitive style in children with autism spectrum disorders. J Autism Dev Disord. 2009;39:635–42.
- Rogers SJ, Ozonoff S. Annotation: What do we know about sensory dysfunction in autism? A critical review of the empirical evidence. J Child Psychol Psychiatry. 2005;46:1255–68.
- Germani T, Zwaigenbaum L, Bryson S, Brian J, Smith I, Roberts W, Szatmari P, Roncadin C, Sacrey LAR, Garon N, Vaillancourt T. Brief report: assessment of early sensory processing in infants at high-risk of autism spectrum disorder. J Autism Dev Disord. 2014;44:3264–70.
- Ben-Sasson A, Hen L, Fluss R, Cermak SA, Engel-Yeger B, Gal E. A metaanalysis of sensory modulation symptoms in individuals with autism spectrum disorders. J Autism Dev Disord. 2009;39:1–11.
- Green SA, Hernandez L, Tottenham N, Krasileva K, Bookheimer SY, Dapretto M. Neurobiology of sensory overresponsivity in youth with autism spectrum disorders. JAMA Psychiatry. 2015;72:778–86.
- 37. Green SA, Rudie JD, Colich NL, Wood JJ, Shirinyan D, Hernandez L, Tottenham N, Dapretto M, Bookheimer SY. Overreactive brain responses to sensory stimuli in youth with autism spectrum disorders. J Am Acad Child Adolesc Psychiatry. 2013;52:1158–72.
- 38. Kern JK. The possible role of the cerebellum in autism/PDD: disruption of a multisensory feedback loop. Med Hypotheses. 2002;59:255–60.
- Mosconi MW, Wang Z, Schmitt LM, Tsai P, Sweeney JA. The role of cerebellar circuitry alterations in the pathophysiology of autism spectrum disorders. Front Neurosci. 2015;9:296.
- Lord C, Rutter M, Couteur A. Autism Diagnostic Interview-Revised: a revised version of a diagnostic interview for caregivers of individuals with possible pervasive developmental disorders. J Autism Dev Disord. 1994;24:659–85.
- 41. Rutter M, Bailey A, Lord C, Berument S. Social Communication Questionnaire. Western Psychological Services: Los Angeles; 2003
- 42. Lord C, Risi S, Lambrecht L, Cook EH, Leventhal BL, DiLavore PC, Pickles A, Rutter M. The autism diagnostic observation schedule-generic: a standard measure of social and communication deficits associated with the spectrum of autism. J Autism Dev Disord. 2000;30:205–23.
- Mullen EM. Mullen Scales of Early Learning. Circle Pines: AGS Publishing; 1995.
- Sparrow S, Balla D, Cicchetti D. Vineland Adaptive Behavior Scales: Second Edition. Shoreview: AGS Publishing; 2005.
- Bodfish JW, Symons FJ, Parker DE, Lewis MH. Varieties of repetitive behavior in autism: comparisons to mental retardation. J Autism Dev Disord. 2000;30:237–43.
- Baranek GT, David FJ, Poe MD, Stone WL, Watson LR. Sensory Experiences Questionnaire: discriminating sensory features in young children with autism, developmental delays, and typical development. J Child Psychol Psychiatry. 2006;47:591–601.
- Turner M. Annotation: Repetitive behaviour in autism: a review of psychological research. J Child Psychol Psychiatry. 1999;40:839–49.
- Mirenda P, Smith IM, Vaillancourt T, Georgiades S, Duku E, Szatmari P, Bryson S, Fombonne E, Roberts W, Volden J, Waddell C, Zwaigenbaum L. Validating the Repetitive Behavior Scale-Revised in young children with autism spectrum disorder. J Autism Dev Disord. 2010;40:1521–30.
- Little LM, Freuler AC, Houser MB, Guckian L, Carbine K, David FJ, Baranek GT. Psychometric validation of the Sensory Experiences Questionnaire. Am J Occup Ther. 2011;65:207–10.

- Gouttard S, Styner M, Prastawa M, Piven J, Gerig G. Assessment of reliability of multi-site neuroimaging via traveling phantom study. Med Image Comput Comput Assist Interv. 2008;11(Pt 2):263–70.
- Liu Z, Wang Y, Gerig G, Gouttard S, Tao R, Fletcher T, Styner M. Quality Control of Diffusion Weighted Images. Proc SPIE Int Soc Opt Eng. 2010;7628: 7628011–9.
- Oguz I, Farzinfar M, Matsui J, Budin F, Liu Z, Gerig G, Johnson HJ, Styner M. DTIPrep: quality control of diffusion-weighted images. Front Neuroinform. 20148:4
- Verde AR, Budin F, Berger J-B, Gupta A, Farzinfar M, Kaiser A, Ahn M, Johnson H, Matsui J, Hazlett HC, Sharma A, Goodlett C, Shi Y, Gouttard S, Vachet C, Piven J, Zhu H, Gerig G, Styner M. UNC-Utah NA-MIC framework for DTI fiber tract analysis. Front Neuroinform. 2014;7:51.
- Geng X, Gouttard S, Sharma A, Gu H, Styner M, Lin W, Gerig G, Gilmore JH. Quantitative tract-based white matter development from birth to age 2 years. Neuroimage. 2012;61:542–57.
- Oishi K, Faria AV, Zijl PCM van, Mori S. MRI Atlas of Human White Matter. Volume 12. Cambridge: Academic Press; 2010.
- Gao W, Lin W, Chen Y, Gerig G, Smith JK, Jewells V, Gilmore JH. Temporal and spatial development of axonal maturation and myelination of white matter in the developing brain. AJNR Am J Neuroradiol. 2009;30:290–6.
- Sadeghi N, Prastawa M, Fletcher PT, Wolff J, Gilmore JH, Gerig G. Regional characterization of longitudinal DT-MRI to study white matter maturation of the early developing brain. Neuroimage. 2013;68:236–47.
- 58. Benjamini Y, Krieger AM, Yekutieli D. Adaptive linear step-up procedures that control the false discovery rate. Biometrika. 2006;93:491–507.
- Elison JT, Wolff JJ, Heimer DC, Paterson SJ, Gu H, Hazlett HC, Styner M, Gerig G, Piven J. Frontolimbic neural circuitry at 6 months predicts individual differences in joint attention at 9 months. Dev Sci. 2013;16:186–97.
- Solso S, Xu R, Proudfoot J, Hagler DJ, Campbell K, Venkatraman V, Carter Barnes C, Ahrens-Barbeau C, Pierce K, Dale A, Eyler L, Courchesne E. Diffusion tensor imaging provides evidence of possible overconnectivity in frontal lobes in autism spectrum disorder toddlers. Biol Psychiatry. 2016;79: 676–84.
- Ben Bashat D, Kronfeld-Duenias V, Zachor DA, Ekstein PM, Hendler T, Tarrasch R, Even A, Levy Y, Ben Sira L. Accelerated maturation of white matter in young children with autism: a high b value DWI study. Neuroimage. 2007;37:40–7.
- 62. Weinstein M, Ben-Sira L, Levy Y, Zachor DA, Ben Itzhak E, Artzi M, Tarrasch R, Eksteine PM, Hendler T, Ben Bashat D. Abnormal white matter integrity in young children with autism. Hum Brain Mapp. 2011;32:534–43.
- Hoeft F, Barnea-Goraly N, Haas BW, Golarai G, Ng D, Mills D, Korenberg J, Bellugi U, Galaburda A, Reiss AL. More is not always better: increased fractional anisotropy of superior longitudinal fasciculus associated with poor visuospatial abilities in Williams syndrome. J Neurosci. 2007;27:11960–5.
- Gao JH, Parsons LM, Bower JM, Xiong J, Li J, Fox PT. Cerebellum implicated in sensory acquisition and discrimination rather than motor control. Science. 1996;272:545–7.
- Proville RD, Spolidoro M, Guyon N, Dugué GP, Selimi F, Isope P, Popa D, Léna C. Cerebellum involvement in cortical sensorimotor circuits for the control of voluntary movements. Nat Neurosci. 2014;17:1233–9.
- Doyon J, Song AW, Karni A, Lalonde F, Adams MM, Ungerleider LG. Experience-dependent changes in cerebellar contributions to motor sequence learning. Proc Natl Acad Sci U S A. 2002;99:1017–22.
- 67. Chugani HT, Phelps ME, Mazziotta JC. Positron emission tomography study of human brain functional development. Ann Neurol. 1987;22:487–97.
- Ridler K, Veijola JM, Tanskanen P, Miettunen J, Chitnis X, Suckling J, Murray GK, Haapea M, Jones PB, Isohanni MK, Bullmore ET. Fronto-cerebellar systems are associated with infant motor and adult executive functions in healthy adults but not in schizophrenia. Proc Natl Acad Sci U S A. 2006;103:15651–6.
- Hoshi E, Tremblay L, Féger J, Carras PL, Strick PL. The cerebellum communicates with the basal ganglia. Nat Neurosci. 2005;8:1491–3.
- Middleton FA, Strick PL. Anatomical evidence for cerebellar and basal ganglia involvement in higher cognitive function. Science. 1994;266: 458–61.
- Bostan AC, Dum RP, Strick PL. Cerebellar networks with the cerebral cortex and basal ganglia. Trends Cogn Sci. 2013;17:241–54.
- Nakamae T, Narumoto J, Sakai Y, Nishida S, Yamada K, Nishimura T, Fukui K. Diffusion tensor imaging and tract-based spatial statistics in obsessivecompulsive disorder. J Psychiatr Res. 2011;45:687–90.

- Fitzgerald KD, Liu Y, Reamer EN, Taylor SF, Welsh RC. Atypical frontal-striatalthalamic circuit white matter development in pediatric obsessivecompulsive disorder. J Am Acad Child Adolesc Psychiatry. 2014;53:1225–33. 1233 e1–9.
- Ozonoff S, Young GS, Landa RJ, Brian J, Bryson S, Charman T, Chawarska K, Macari SL, Messinger D, Stone WL, Zwaigenbaum L, Iosif A-M. Diagnostic stability in young children at risk for autism spectrum disorder: a baby siblings research consortium study. J Child Psychol Psychiatry. 2015;56:988–98.

Submit your next manuscript to BioMed Central and we will help you at every step:

- We accept pre-submission inquiries
- Our selector tool helps you to find the most relevant journal
- We provide round the clock customer support
- Convenient online submission
- Thorough peer review
- Inclusion in PubMed and all major indexing services
- Maximum visibility for your research

Submit your manuscript at www.biomedcentral.com/submit

