Atypical brain asymmetry in autism – a candidate for clinically

meaningful stratification

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Abstract

Background. Autism Spectrum Disorder (henceforth 'autism') is a highly heterogeneous neurodevelopmental condition with few effective treatments for core and associated features. To make progress we need to both identify and validate neural markers that help to parse heterogeneity to tailor therapies to specific neurobiological profiles. Atypical hemispheric lateralization is a stable feature across studies in autism, however its potential of lateralization as a neural stratification marker has not been widely examined. Methods. In order to dissect heterogeneity in lateralization in autism, we used the large EU-AIMS Longitudinal European Autism Project dataset comprising 352 individuals with autism and 233 neurotypical (NT) controls as well as a replication dataset from ABIDE (513 autism, 691 NT) using a promising approach that moves beyond mean-group comparisons. We derived grey matter voxelwise laterality values for each subject and modelled individual deviations from the normative pattern of brain laterality across age using normative modeling. **Results**. Results showed that individuals with autism had highly individualized patterns of both extreme right- and leftward deviations, particularly in language-, motor- and visuospatial regions, associated with symptom severity. Language delay (LD) explained most variance in extreme rightward patterns, whereas core autism symptom severity explained most variance in extreme leftward patterns. Follow-up analyses showed that a stepwise pattern emerged with individuals with autism with LD showing more pronounced rightward deviations than autism individuals without LD. **Conclusion**. Our analyses corroborate the need for novel (dimensional) approaches to delineate the heterogeneous

neurophenotype for clinically meaningful stratification in autism.

Introduction

Autism spectrum disorder (autism) is a neurodevelopmental condition characterized by social-communicative deficits, restricted, repetitive behaviors and sensory abnormalities (1). One of the key characteristics of autism is its great phenotypic and biological heterogeneity (2,3). Particularly in the neuroimaging literature, results are mixed and inconsistent and have been attributed to this heterogeneity in the autism population. To provide a more coherent picture of the complex neuropathology of autism, it is critical to tackle its heterogeneity and identify the neural markers that are consistent across studies. Such markers can provide clinically relevant stratification of individuals with autism.

When it comes to identifying a consistently implicated neural feature in autism, a large body of literature converges towards a disruption of hemispheric specialization - one of the most fundamental biological properties of our brains (4,5). This basic organizational principle describes that the two hemispheres differ in their functional specialization and exhibit pronounced structural asymmetries (6,7). Functional specialization involves leftward lateralization of language and motor skills and rightward lateralization of spatial perceptual abilities (8). It is also evident in grey matter (GM) asymmetries with frontal opercular and temporal perisylvian regions, and hippocampus exhibiting

leftward asymmetries, whereas thalamus and posterior parietal cortex showing rightward asymmetries (9,10).

Individuals with autism exhibit impairments in left hemisphere skills such as social-communication, language and motor-related symptoms, whilst appearing relatively intact in right hemisphere functions such as visuospatial skills (11). This lateralized pattern of deficits and strengths in autism has given rise to theories trying to reconcile its complex clinical profile with atypical structural hemispheric specialization (11). In the largest autism cohort to date comprising over 3000 subjects individuals with autism presented with widespread leftward cortical reductions (12). Smaller studies are in line with this showing either a reduction or even reversal of typical leftward asymmetries in language- and motor-related regions (13-20). Thus, atypical lateralization in autism is among the most replicated findings with moderate to high effect sizes (21) in the otherwise heterogeneous neuroimaging literature. However, inconsistencies in results remain regarding regional specificity and direction of observed patterns. Such inconsistencies are usually attributed to sample heterogeneity and delineating heterogeneity still remains one of the central tasks in autism research.

The relationship between heterogeneity and brain asymmetry in autism may be further co-dependent on age, sex, handedness, different symptom profiles and comorbidities such as attention-deficit hyper-activity disorder (ADHD). *Age*: Lateralization in neurotypicals becomes more pronounced through agerelated maturational processes (22). This typical trajectory is disrupted in

individuals with autism showing increasing reversed rightward lateralization (23). *Sex*: NT males are usually more strongly lateralized, while NT females have a more symmetric distribution pattern (24). How sex affects atypical asymmetry in autism is unknown. *Handedness*: Left-handed individuals have a higher chance of a different organization in the brain than right-handed individuals. Individuals with autism exhibit elevated rates of non-right-handedness, which has been attributed to atypical specialization in the brain (25). *Language delay (LD):* Individuals with autism and early LD show more pronounced deviations from typical asymmetry than those without LD (15,26). *ADHD*: Some conditions that are also a common comorbidity in autism, such as ADHD, are also associated with atypical lateralization (27).

Despite increased recognition of heterogeneity in autism (28), there is little effort to address such challenges methodologically. One first example towards quantifying biological variation at the individual level in autism was recently demonstrated (29) using a novel normative modeling method (30,31). Similar to the use of growth charts in paediatric medicine, normative modelling aims to place each individual with respect to centiles of variation in the population and thereby facilitates a move away from classical case-control analyses that ignore individual differences. Applying normative modelling to cortical thickness estimates, Zabihi et al., 2018 (29) showed that individuals with autism exhibit highly individualized atypicalities of cortical development.

This study was designed to address heterogeneity in autism with regard to age, sex and core and co-occurring symptoms in the context of brain

lateralization using novel individualized analyses: 1) we transcend classical case-control analysis and address inter-individual variation by applying normative modeling (30,32). 2) We aim to identify laterality-related subtypes by considering co-occurring clinical symptoms in autism such as language development. Through capturing variation at the individual level in combination with addressing different sources of heterogeneity and using a consistent imaging feature in autism, our work provides a step towards precision neuroscience in autism.

Methods and Materials

Participants

Participants were part of the EU-AIMS and AIMS-2-TRIALS Longitudinal European Autism Project (LEAP) (33,34) cohort – the largest European multicenter initiative aimed at identifying biomarkers in autism. Participants underwent comprehensive clinical, cognitive and MRI assessment at one of six collaborating sites (Figure S1). All participants with ASD had an existing clinical diagnosis of autism. For details on participants, study design and exclusion criteria, see Supplemental Information (SI) and (34). The final sample comprised 352 individuals with autism (259 males and 93 females), and 233 NT controls (154 males and 79 females) between 6 and 30 years. For details on demographic information, see Table 1.

Clinical and cognitive measures

IQ was assessed using the Wechsler Abbreviated Scales of Intelligence. The Autism Diagnostic Observation Schedule (ADOS-G (35)) measured clinical core symptoms of autism. The Autism Diagnostic Interview-Revised (ADI-R) (36) was used to measure parent-rated autistic symptoms and LD, which was defined as having onset of first words later than 24 months and/or first phrases later than 33 months. ADHD symptoms were assessed with the DSM-5 ADHD rating scale. Handedness was assessed with the short version of the Edinburgh Handedness Inventory (37). For further details on all cognitive measures, see SI and (33).

Image Preprocessing

For MRI data acquisition parameters, see SI and Table S1. Structural T1weighted images were preprocessed according to a validated laterality pipeline (15,38,39) using the CAT12 toolbox (http://www.neuro.unijena.de/cat) (see Figure S2). All original images were segmented and affine registered to a symmetric tissue probability map before being reflected across the cerebral midline ($x \Box = \Box 0$). All segmented reflected and original (non-reflected) grey matter (GM) maps were then used to generate a symmetrical study-specific template via a flexible, high-dimensional nonlinear diffeomorphic registration algorithm (DARTEL) (40). They were next registered to the symmetrical study-specific template as per standard DARTEL procedures. An intensity modulation step was included to retain voxel-wise information on local volume (39). The final resulting images were modulated, warped, reflected (I_{ref}) and non-reflected (I_{nref}) GM intensity images. A laterality index (LI) was calculated at each voxel using the following formula: LI = 2($I_{nref}-I_{ref}$)/($(I_{nref} + I_{ref})$. Positive values in the right hemisphere of the LI image indicate rightward asymmetry, and negative values in the right hemisphere of the LI image indicate leftward asymmetry. Values of LI in the left hemisphere have identical magnitude, but opposite sign and were therefore excluded from further analyses. LI images were smoothed with a 4-mm FWHM isotropic Gaussian kernel.

Normative Modelling

The normative modeling method has been described in detail previously (29,30,41–43). In summary, we estimated a normative brain aging model at each GM laterality voxel by using Gaussian process regression (GPR) (44), a Bayesian non-parametric interpolation method that yields coherent measures of predictive confidence in addition to point estimates (for details see SI). With this method, we could predict both the expected regional GM asymmetry changes and the associated predictive uncertainty for each individual allowing us to quantify the voxelwise deviation of GM asymmetry from the NT range across the entire brain.

First, we trained a GPR model at each voxel on the NT cohort using age, sex and site as covariates to predict GM asymmetry resulting in a developmental model of GM asymmetry in NT individuals. To avoid over-fitting, assess generalizability and determine whether NT individuals fall within the normative range, we used 10-fold cross-validation in NT individuals before retraining the model in the entire sample to make predictions in individuals with autism (see SI). We generated normative probability maps (NPM), which quantify the deviation of each participant from the normative model for GM asymmetry at each voxel. These subject-specific Z-score images provide a statistical estimate of how much each individual's true laterality value differs from the predicted laterality value with reference to the NT pattern at each voxel given the participants age, sex and site. NPMs were thresholded at an absolute value of Z>|2.6| (30,42). Based on this fixed threshold, we defined extreme rightwardlateralized (positive) and extreme leftward-lateralized (negative) deviations for each participant.

All extreme deviations per subject were summarized into (log-transformed) scores representing the percentage of extreme rightward and extreme leftward deviations per subject in relation to the total number of intracerebral voxels. These percentage scores were compared calculating a GLM including diagnosis and sex as the regressors of interest, and age as covariate. To compare our normative modeling against a conventional group-mean difference analysis we ran Permutation Analysis of Linear Models (PALM) on the LI images examining the sex-by-diagnosis(-by-age) interaction with site as covariate.

Spatial characterization of deviations

We applied two strategies to spatially characterize the extreme rightward and leftward deviations: 1) We generated spatial overlap maps for individuals in the same diagnostic and gender group by summing up the number of extreme deviations in each voxel for each subject. These were then divided by the total

number of subjects (multiplied by 100) to represent the percentage of extreme right- and leftward deviations per brain voxel. 2) We extracted extreme rightward and leftward deviations within structurally (Harvard-Oxford atlas (HOA (45))), and functionally (neurosynth (http://neurosynth.org, accessed June 2019) (46)) defined regions of interest (ROIs) (see SI). For latter, we used the search terms 'language', 'motor' (left-lateralized), 'visuospatial', 'attention' (right-lateralized) and 'monitoring', 'mentalizing' (no lateral bias). Results were FDR-corrected. Effect sizes were computed using Cohen's *d*. For robustness and replicability analyses, see SI.

Relative contributions of different cognitive and behavioral measures

We ran relative importance analyses within individuals with autism including variables related to lateralization such as LD, ADHD, handedness, sex and symptom severity (CSS). This was done in a sample with reduced sample size (N=305) without missing values on these variables. We used averaging over orderings (47) to rank the relative contribution of highly correlated regressors to the linear regression model. The variable showing strongest contribution to explained variance was followed-up with additional analyses.

Symptom Associations

An individual-level atypicality score was estimated for each individual through extreme value statistics by computing the 10% trimmed mean of the 1% top deviations (29). We then computed one-tailed, FDR-corrected Pearson's correlations between these individual atypicality scores and the ADI-R and ADOS-2 symptom severity scores.

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Robustness and Replicability

To assess robustness, we estimated a separate normative model without including site as a covariate, but removing site effects using ComBat (48). For the sake of comparability, we also applied FDR-correction (49) to the NPMs. We also included full-scale intelligence (FIQ) and handedness as nuisance covariates. To test whether the effects were robust against the influence of intellectual disability (ID), we reran analyses excluding individuals with FIQ <70. Finally, as individuals with autism with and without LD were not matched on certain demographic variables, we additionally created a sub-sample matched for age and symptom severity and re-ran second-level statistical analyses to assess robustness of results. To assess replicability, we selected a sample from the publically available Autism Brain Imaging Data Exchange (ABIDE) I and II (50,51) to examine extreme right- and leftward deviations in an independent datasets. For details, see the SI and Table S2.

Results

The classic PALM analysis to assess mean group differences did not yield any significant results. Figure 1 depicts the spatial representation of the voxelwise normative model in NT males in the largest site KCL (N=42). Results were comparable for NT females (Figure S3) and when using ComBat to address site effects (Figures S4a and S4b). The forward model for all other sites by sex is shown in Figure S5. Both in NT males and females leftward shifts in GM asymmetry were mainly evident in Heschl's gyrus, planum temporale, parietal and central operculum, angular gyrus (AG), supramarginal gyrus (SMG), pars triangularis, postcentral gyrus, precentral gyrus, superior

and middle frontal gyrus, lateral occipital cortex (LOC) and cerebellum. Rightward shifts in GM asymmetry mainly occurred in middle and inferior temporal gyrus, posterior SMG, AG, superior parietal lobule, precuneus, anterior cingulate cortex, supracalcarine cortex, lingual gyrus, occipital pole, and caudate.

The accuracy of the normative model was evaluated using the correlation between the true and the predicted voxel values (Rho) generated under 10fold cross-validation. NT Rho maps were restricted to positive values and further analyses were conducted on normative model maps thresholded with these. For unthresholded Rho maps along with root mean square error maps, see Figures S6 and S7.

Characterization of extreme deviations

Overall, males and females with autism showed both significantly more extreme rightward ($F_{(1)}$ =12.5, p<0.001, d=0.31) and leftward deviations ($F_{(1)}$ =12.0, p<0.001, d=0.29) compared to NT males and females (Figure 2). There were no sex differences (right: $F_{(1)}$ =0.2, p=0.67, d=0.06; left: $F_{(1)}$ =0.2, p=0.67, d=0.06) nor sex-by-diagnosis interactions (right: $F_{(1)}$ =2.2, p=0.14, d=0.26; left: $F_{(1)}$ =0.5, p=0.47, d=0.11). Controlling the FDR at the individual level of each NPM led to identical conclusions (see SI and Figure S8). Overall individuals with autism showed a trend towards more extreme right- than leftward deviations (χ^2 =3.59, p=0.05). Further, there was a significant positive correlation between extreme right- and leftward deviations (r=0.41, p<0.001). For rightward deviations, females with autism showed the highest overlap in parahippocampal gyrus, putamen and amygdala, while males with autism mostly in the middle temporal gyrus and hippocampus (Figure 2a). For leftward deviations, females with autism showed highest overlap in orbitofrontal cortex, frontal pole and postcentral gyrus, while males with autism mostly in LOC, temporal pole and thalamus (Figure 2b). On average females with autism had higher overlapping deviating regions than males with autism (right: χ^2 =667, *p*<0.001; left: χ^2 =277, *p*<0.001).

When considering deviations by structural ROIs, we found a significant main effect of diagnosis in the frontal ($F_{(1)}$ =15.7, p<0.001, d=0.35) and central operculum ($F_{(1)}$ =14.9, p<0.001, d=0.34) with individuals with autism showing more extreme rightward deviations, and in the superior lateral occipital cortex ($F_{(1)}$ =14.4, p<0.001, d=0.32) with individuals with autism showing more extreme leftward deviations. Further, there was a significant sex-by-diagnosis interaction in the temporal occipital fusiform cortex (TOFC) ($F_{(1)}$ =19.2, p<0.001, d=0.79) with females with autism having more extreme rightward deviations by functional ROIs, individuals with autism showed extreme rightward deviations in the motor network ($F_{(1)}$ =9.8, p=0.001, d=0.27), whereas more extreme leftward deviations in the visuospatial network ($F_{(1)}$ =11.6, p<0.001, d=0.3). Results were robust across sensitivity analyse and replicated in the large ABIDE sample (see SI).

Relative contributions

Results revealed that LD had relatively the largest importance for explaining extreme rightward deviations (LD: 62.1% > handedness: 17.3% > sex: 10.9% >

ADHD: 9.9% > CSS: 0.5%), while symptom severity for extreme leftward deviations (CSS: 40.4% > LD: 34.1% > sex: 23.6% > ADHD: 12.3% > handedness: 2.3%) (Figure 3). LD was further followed up showing a significant main effect for rightward deviations ($F_{(2)}$ =10.5, p<0.001). Specifically, individuals with autism and LD were different from both individuals with autism without LD ($t_{(213)}$ =2.5, p=0.01, d=0.32) and NT individuals ($t_{(152)}$ =4.6, p<0.001, d=0.58), while individuals with autism without LD were not different from NT individuals ($t_{(256)}$ =1.9, p=0.06, d=0.21). This stepwise pattern was overall more pronounced in males than in females with autism (Figure 4 and Figure S9).

Results by language-related ROIs showed a main effect of LD for extreme rightward deviations in the frontal operculum ($F_{(2)}$ =10.6, p<0.001), the central operculum ($F_{(2)}$ =14.9, p<0.001) and the motor network ($F_{(2)}$ =6.9, p=0.001). Rightward deviations in the language network were significant ($F_{(2)}$ =4.1, p=0.01), but did not survive FDR-correction. For all, there was a stepwise pattern, with individuals with autism and LD showing more pronounced deviations. For details, see SI.

Symptom Associations

Across the whole brain, there were significant associations between extreme rightward deviations and ADI-R communication scores (r=0.14, p=0.004). No correlations survived FDR-correction in individuals with autism with and without LD. In males with autism, there were significant positive correlations between rightward deviations and core autism symptoms (ADI-R social:

r=0.18, p=0.002; ADI-R communication: r=0.19, p=0.001). In females with autism, as well as in males and females with autism with and without LD, no results survived FDR-correction.

Replicability

In line with the results of the EU-AIMS LEAP sample, males and females with autism of the combined ABIDE dataset showed both significantly more extreme rightward ($F_{(1)}$ =7.7, p=0.006, d=0.14) and leftward deviations ($F_{(1)}$ =17.1, p<0.001, d=0.24) compared to NT individuals (see Figures S10a-b). As in EU-AIMS LEAP, there were no sex differences (right: $F_{(1)}$ =1.1, p=0.3, d=0.14; left: $F_{(1)}$ =2.9, p=0.13, d=0.19) and no sex-by-diagnosis interactions (right: $F_{(1)}$ =3.6, p=0.06, d=0.26; left: $F_{(1)}$ =3.4, p=0.06, d=0.26). When considering deviations by functional ROIs significant in the primary analysis, individuals with autism also showed extreme rightward deviations in the motor network ($F_{(1)}$ =5.2, p=0.02, d=0.21), and extreme leftward deviations in the visuospatial network ($F_{(1)}$ =4.0, p=0.05, d=0.12). There were no significant group differences in the frontal operculum ($F_{(1)}$ =0.2, p=0.6, d=0.02). For further details, see the SI and Figures 10a-b.

Discussion

In this study, we mapped extreme deviations in structural asymmetry at the individual level in comparison to a normative model of laterality development. A further aim was to explore laterality as a stratification marker in autism in a large and deeply phenotyped cohort and an equally large replication cohort. We found highly individualized patterns of both right- and leftward asymmetry

deviations in males and females with autism. In contrast, when using classical case-control analyses, we did not detect any significant group differences, emphasizing the need to move beyond group averages to capture an accurate representation of the phenotype at the individual level. Similarly, a recent study addressing heterogeneity dimensionally points out that traditional case-control analyses yield smaller effects and miss atypicalities detected with a dimensional subtyping approach (52). By deriving statistical inferences at the individual level, we demonstrated that individuals with autism show a different, individualized pattern that would otherwise be missed when focusing on a common neurobiological signature. In fact, spatial overlap of extreme deviations across subjects was minimal in both males and females with autism (Figure 2 and Figure S10) corroborating the high degree of heterogeneity across subjects.

Moreover, extreme right- and leftward deviations were highly correlated with each other, which suggests that biological factors acting on atypical laterality shifts might not be biased towards one side in the same individual in autism. It thus also appears that a disruption in establishing the typical laterality pattern is not confined to one hemisphere, but there are inter-related alterations in both. In line with several previous reports (13,15,53), our results also show that individuals with autism have slightly more extreme rightward than leftward deviations. The biological underpinning are unknown, but likely include both genetic (54), prenatal endocrine (55) and environmental factors. For instance, Geschwind et al., (2002) (56) reported that left frontal and extreme rightward deviations regions are under less genetic control as compared to the right hemisphere and therefore are more susceptible to environmental influences during neurodevelopment. Whether prenatal androgens and related early immune activation (55,57) contribute to atypical hemispheric development in autism, remains to be established.

Extreme rightward deviations were most pronounced in the motor network and frontal operculum and extreme leftward deviations in the visuospatial network. Accordingly, rightward shifts in asymmetry in language related regions, particularly in Broca's area, are frequently reported in autism (16,17,58). However, atypical lateralization of motor and visuospatial performance is underexplored in autism. Accumulating evidence suggests the important role of motor-related asymmetries in the neurobiology of autism (5,13,59,60). Despite mostly intact visuospatial performance in autism, atypical activation patterns have also been reported in individuals with autism while performing visuospatial tasks (61). More specific cognitive measures are needed to establish the functional relevance of these alterations.

Males and females with autism showed a similar degree of extreme deviations (Figure 2). Specifically, in the TOFC, females with autism showed stronger rightward deviations, while males with autism and NT females both show fewer rightward deviations than NT males. This 'occipital face area' is strongly right-lateralized in NT individuals (62). Individuals with autism exhibit atypical face processing strategies and these are more pronounced in females with autism (63). Being one of the most reported impairments in social cognition in

autism, face processing and related atypical, sex-differential lateralization awaits further exploration.

When considering the spatial extent of deviations, females with autism show on average greater overlap across differently implicated regions than males with autism (Figure 2 and Figure S10). This suggests that females with autism constitute a less heterogeneous group than males with autism who show less pronounced overlap of focal atypicalities in laterality. Also, when considering males and females separately, atypical asymmetry was only associated with more social- communicative symptoms in males with autism, but not females with autism, suggesting that phenotypically similar manifestations of atypical lateralization appear to have differential cognitive implications for males and females with autism.

We found that both extreme left- and rightward deviations were associated with LD, but only rightward deviations differentiated individuals with autism with and without LD from each other. This is in line with prior reports (15,26). Two structural studies further showed differential morphological alterations in individuals with autism with different developmental language profiles (64) and a loss of leftward asymmetry in the arcuate fasciculus in non-verbal children with autism (65). The degree of atypical rightward lateralization may thus constitute a biological marker of different etiological subgroups with different language profiles in autism. Evidence suggests onset of language before the age of two years (66), and the level of language at the ages five and six years (67), predict functional outcome later in life in autism. Developing stratification

markers based on LD will thus have important clinical implications for early prognosis, individualized diagnoses and early language-based interventions in individuals at risk.

Strengths and Limitations

We present analyses in a large-scale, deeply phenotyped and prospectively harmonized dataset. Results present robust across several analyses addressing potential confounds and are overall also observable in the largescale, yet not harmonized ABIDE. Replicability and robustness of findings are important prerequisites as a first step towards establishing neural biomarkers.

Nevertheless, our findings should be considered in light of some limitations. 1) Despite the large dataset, when dividing individuals by diagnostic group, sex, LD and site, sub-samples decrease substantially in size. While our Bayesian GPR model adapts to the availability of data (i.e. gives more conservative estimates when data density decreases (31)), the model would yield more precise estimates with larger sample sizes. 2) Large datasets come at the expense of additional confounds associated with site. Here, we addressed this both by including site in the model and by running an alternative method for controlling batch effects. Our results are robust, however, residual site effects cannot fully be excluded. 3) The question arises whether there are also experience-dependent influences on cortical asymmetry. Longitudinal analyses are needed to pinpoint the onset and trajectory of atypical development in subgroups on the autism spectrum. 4) The normative modeling approach is suitable for detecting extreme deviations from a

normative pattern. However, an alternative hypothesis that individuals with autism might lack specialization of either hemisphere (59) which would be expressed in reduced laterality might be less detected with the current approach.

Conclusion

We estimated a normative model of GM voxelwise asymmetry based on a large NT cohort and applied this to a large, deeply phenotyped and heterogeneous autism sample. Our results confirm that atypical asymmetry is a core feature of the autism neurophenotype, shows highly individualized patterns across individuals and is differentially related to different symptom profiles such as language delay. Further exploration of such associations has the potential to yield clinically relevant stratification markers needed for precision medicine.

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Disclosures

JKB has been a consultant to, advisory board member of, and a speaker for Takeda/Shire, Medice, Roche, and Servier. He is not an employee of any of these companies and not a stock shareholder of any of these companies. He has no other financial or material support, including expert testimony, patents, or royalties. CFB is director and shareholder in SBGneuro Ltd. TC has received consultancy from Roche and received book royalties from Guildford Press and Sage. DM has been a consultant to, and advisory board member, for Roche and Servier. He is not an employee of any of these companies, and not a stock shareholder of any of these companies. TB served in an advisory or consultancy role for Lundbeck, Medice, Neurim Pharmaceuticals, Oberberg GmbH, Shire, and Infectopharm. He received conference support or speaker's fee by Lilly, Medice, and Shire. He received royalities from Hogrefe, Kohlhammer, CIP Medien, Oxford University Press; the present work is unrelated to these relationships. The other authors report no biomedical financial interests or potential conflicts of interest.

References

1. American Psychiatric Association (2013): Diagnostic and Statistical Manual

of Mental Disorders 1. American Psychiatric Association. Arlington.

https://doi.org/10.1176/appi.books.9780890425596.893619

- Lombardo M V., Lai MC, Baron-Cohen S (2019): Big data approaches to decomposing heterogeneity across the autism spectrum. *Mol Psychiatry*. https://doi.org/10.1038/s41380-018-0321-0
- Lenroot RK, Yeung PK (2013): Heterogeneity within autism spectrum disorders: What have we learned from neuroimaging studies? *Frontiers in Human Neuroscience*. https://doi.org/10.3389/fnhum.2013.00733
- Lindell AK, Hudry K (2013): Atypicalities in cortical structure, handedness, and functional lateralization for language in autism spectrum disorders. *Neuropsychol Rev* 23: 257–70.
- Floris DL, Howells H (2018): Atypical structural and functional motor networks in autism. *Progress in Brain Research*. https://doi.org/10.1016/bs.pbr.2018.06.010
- Gazzaniga MS (1995): Principles of human brain organization derived from split-brain studies. *Neuron* 14: 217–28.
- Rogers LJ, Zucca P, Vallortigara G (2004): Advantages of having a lateralized brain. *Proc Biol Sci* 271 Suppl: S420–S422.
- Springer SP, Deutsch G (1998): Left brain, right brain: Perspectives from cognitive neuroscience, 5th ed. Left Brain, Right Brain: Perspectives from Cognitive Neuroscience, 5th Ed.
- Good CD, Johnsrude I, Ashburner J, Henson RN, Friston KJ, Frackowiak RS (2001): Cerebral asymmetry and the effects of sex and handedness on brain structure: a voxel-based morphometric analysis of 465 normal adult human brains. *Neuroimage* 14: 685–700.

- Jäncke L, Schlaug G, Huang Y, Steinmetz H (1994): Asymmetry of the planum parietale. *Neuroreport*. https://doi.org/10.1097/00001756-199405000-00035
- Prior MR, Bradshaw JL (1979): Hemisphere Functioning in Autistic Children. *Cortex.* https://doi.org/10.1016/S0010-9452(79)80008-8
- Postema MC, Rooij D van, Anagnostou E, Arango C, Auzias G, Behrmann M, *et al.* (2019): Altered structural brain asymmetry in autism spectrum disorder: large-scale analysis via the ENIGMA Consortium. *bioRxiv*. https://doi.org/10.1101/570655
- Floris DL, Barber AD, Nebel MB, Martinelli M, Lai MC, Crocetti D, et al. (2016): Atypical lateralization of motor circuit functional connectivity in children with autism is associated with motor deficits. *Mol Autism*. https://doi.org/10.1186/s13229-016-0096-6
- Floris DL, Chura LR, Holt RJ, Suckling J, Bullmore ET, Baron-Cohen S, Spencer MD (2013): Psychological correlates of handedness and corpus callosum asymmetry in autism: the left hemisphere dysfunction theory revisited. *J Autism Dev Disord* 43: 1758–72.
- Floris DL, Lai M-C, Auer T, Lombardo M V., Ecker C, Chakrabarti B, *et al.* (2016): Atypically rightward cerebral asymmetry in male adults with autism stratifies individuals with and without language delay. *Hum Brain Mapp.* https://doi.org/10.1002/hbm.23023
- Herbert MR, Harris GJ, Adrien KT, Ziegler D a, Makris N, Kennedy DN, *et al.* (2002): Abnormal asymmetry in language association cortex in autism.
 Ann Neurol 52: 588–96.
- 17. De Fossé L, Hodge SM, Makris N, Kennedy DN, Caviness VS, McGrath L,

et al. (2004): Language-association cortex asymmetry in autism and specific language impairment. *Ann Neurol* 56: 757–66.

- Rojas DC, Camou SL, Reite ML, Rogers SJ (2005): Planum temporale volume in children and adolescents with autism. *J Autism Dev Disord* 35: 479–86.
- Rojas DC, Bawn SD, Benkers TL, Reite ML, Rogers SJ (2002): Smaller left hemisphere planum temporale in adults with autistic disorder. *Neurosci Lett* 328: 237–40.
- 20. Lange N, Dubray MB, Lee JE, Froimowitz MP, Froehlich A, Adluru N, *et al.*(2010): Atypical diffusion tensor hemispheric asymmetry in autism. *Autism Res.* https://doi.org/10.1002/aur.162
- 21. Preslar J, Kushner HI, Marino L, Pearce B (2014): Autism, lateralisation, and handedness: A review of the literature and meta-analysis. *Laterality*. https://doi.org/10.1080/1357650X.2013.772621
- 22. Brown TT, Lugar HM, Coalson RS, Miezin FM, Petersen SE, Schlaggar BL (2005): Developmental changes in human cerebral functional organization for word generation. *Cereb Cortex*. https://doi.org/10.1093/cercor/bhh129
- 23. Flagg EJ, Cardy JEO, Roberts W, Roberts TPL (2005): Language lateralization development in children with autism: Insights from the late field magnetoencephalogram. *Neurosci Lett*. https://doi.org/10.1016/j.neulet.2005.05.037
- Kansaku K, Kitazawa S (2001): Imaging studies on sex differences in the lateralization of language. *Neurosci Res.* https://doi.org/10.1016/S0168-0102(01)00292-9

- Dane S, Balci N (2007): Handedness, eyedness and nasal cycle in children with autism. *Int J Dev Neurosci*. https://doi.org/10.1016/j.ijdevneu.2007.03.005
- Escalante-Mead PR, Minshew NJ, Sweeney JA (2003): Abnormal brain lateralization in high-functioning autism. *J Autism Dev Disord* 33: 539– 543.
- 27. Silk TJ, Vilgis V, Adamson C, Chen J, Smit L, Vance A, Bellgrove MA (2016): Abnormal asymmetry in frontostriatal white matter in children with attention deficit hyperactivity disorder. *Brain Imaging Behav*. https://doi.org/10.1007/s11682-015-9470-9
- Wolfers T, Floris DL, Dinga R, van Rooij D, Isakoglou C, Kia SM, *et al.* (2019): From pattern classification to stratification: towards conceptualizing the heterogeneity of Autism Spectrum Disorder. *Neurosci Biobehav Rev.* https://doi.org/10.1016/j.neubiorev.2019.07.010
- Zabihi M, Oldehinkel M, Wolfers T, Frouin V, Goyard D, Loth E, *et al.* (2019): Dissecting the Heterogeneous Cortical Anatomy of Autism Spectrum Disorder Using Normative Models. *Biol Psychiatry Cogn Neurosci Neuroimaging.* https://doi.org/10.1016/j.bpsc.2018.11.013
- Wolfers T, Doan NT, Kaufmann T, Alnæs D, Moberget T, Agartz I, *et al.* (2018): Mapping the Heterogeneous Phenotype of Schizophrenia and Bipolar Disorder Using Normative Models. *JAMA Psychiatry*. https://doi.org/10.1001/jamapsychiatry.2018.2467
- Marquand AF, Kia SM, Zabihi M, Wolfers T, Buitelaar JK, Beckmann CF (2019): Conceptualizing mental disorders as deviations from normative functioning. *Molecular Psychiatry*. https://doi.org/10.1038/s41380-019-

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- 32. Marquand AF, Wolfers T, Mennes M, Buitelaar J, Beckmann CF (2016): Beyond Lumping and Splitting: A Review of Computational Approaches for Stratifying Psychiatric Disorders. *Biol Psychiatry Cogn Neurosci Neuroimaging* 1: 433–447.
- 33. Charman T, Loth E, Tillmann J, Crawley D, Wooldridge C, Goyard D, et al. (2017): The EU-AIMS Longitudinal European Autism Project (LEAP): Clinical characterisation. *Mol Autism*. https://doi.org/10.1186/s13229-017-0145-9
- 34. Loth E, Charman T, Mason L, Tillmann J, Jones EJH, Wooldridge C, *et al.* (2017): The EU-AIMS Longitudinal European Autism Project (LEAP): Design and methodologies to identify and validate stratification biomarkers for autism spectrum disorders. *Mol Autism*. https://doi.org/10.1186/s13229-017-0146-8
- 35. Lord C, Risi S, Lambrecht L, Cook EH, Leventhal BL, DiLavore PC, et al. (2000): The autism diagnostic observation schedule-generic: a standard measure of social and communication deficits associated with the spectrum of autism. J Autism Dev Disord 30: 205–223.
- 36. Lord C, Rutter M, Le Couteur A (1994): Autism Diagnostic Interview-Revised: a revised version of a diagnostic interview for caregivers of individuals with possible pervasive developmental disorders. J Autism Dev Disord 24: 659–685.
- 37. Oldfield RC (1971): The assessment and analysis of handedness: the Edinburgh inventory. *Neuropsychologia* 9: 97–113.
- 38. Kurth F, Mackenzie-Graham A, Toga AW, Luders E (2014): Shifting brain

asymmetry: the link between meditation and structural lateralization. *Soc Cogn Affect Neurosci.* https://doi.org/10.1093/scan/nsu029

- Kurth F, Gaser C, Luders E (2015): A 12-step user guide for analyzing voxel-wise gray matter asymmetries in statistical parametric mapping (SPM). *Nat Protoc.* https://doi.org/10.1038/nprot.2015.014
- 40. Ashburner J (2007): A fast diffeomorphic image registration algorithm. *Neuroimage* 38: 95–113.
- Marquand AF, Rezek I, Buitelaar J, Beckmann CF (2016): Understanding Heterogeneity in Clinical Cohorts Using Normative Models: Beyond Case-Control Studies. *Biol Psychiatry*.

https://doi.org/10.1016/j.biopsych.2015.12.023

- 42. Wolfers T, Beckmann CF, Hoogman M, Buitelaar JK, Franke B, Marquand AF (2019): Individual differences v. The average patient: Mapping the heterogeneity in ADHD using normative models. *Psychol Med.* https://doi.org/10.1017/S0033291719000084
- 43. Rezek I, Beckmann CF (2012): Models of Disease Spectra. arXiV.
- 44. Rasmussen CE, Williams CKI (2018): Model Selection and Adaptation of Hyperparameters. *Gaussian Processes for Machine Learning*. https://doi.org/10.7551/mitpress/3206.003.0008
- 45. Smith SM, Jenkinson M, Woolrich MW, Beckmann CF, Behrens TEJ, Johansen-Berg H, et al. (2004): Advances in functional and structural MR image analysis and implementation as FSL. *NeuroImage*. https://doi.org/10.1016/j.neuroimage.2004.07.051
- 46. Yarkoni T, Poldrack RA, Nichols TE, Van Essen DC, Wager TD (2011): Large-scale automated synthesis of human functional neuroimaging data.

Nat Methods. https://doi.org/10.1038/nmeth.1635

47. Sen PK, Lindeman RH, Merenda PF, Gold RZ (1981): Introduction to

Bivariate and Multivariate Analysis. J Am Stat Assoc.

https://doi.org/10.2307/2287559

- 48. Nielson DM, Pereira F, Zheng CY, Migineishvili N, Lee JA, Thomas AG, Bandettini PA (2018): Detecting and harmonizing scanner differences in the ABCD study - annual release 1.0. *bioRxiv*. https://doi.org/10.1101/309260
- Benjamini Y, Hochberg Y (2000): On the adaptive control of the false discovery rate in multiple testing with independent statistics. *J Educ Behav Stat.* https://doi.org/10.3102/10769986025001060
- 50. Di Martino A, Yan CG, Li Q, Denio E, Castellanos FX, Alaerts K, et al. (2014): The autism brain imaging data exchange: Towards a large-scale evaluation of the intrinsic brain architecture in autism. *Mol Psychiatry*. https://doi.org/10.1038/mp.2013.78
- 51. Di Martino A, O'Connor D, Chen B, Alaerts K, Anderson JS, Assaf M, *et al.* (2017): Enhancing studies of the connectome in autism using the autism brain imaging data exchange II. *Sci Data* 4: 170010.
- 52. Tang S, Sun N, Floris DL, Zhang X, Di Martino A, Yeo BTT (2019): Reconciling Dimensional and Categorical Models of Autism Heterogeneity: a Brain Connectomics & Behavioral Study. *Biol Psychiatry*. https://doi.org/10.1016/j.biopsych.2019.11.009
- 53. Cardinale RC, Shih P, Fishman I, Ford LM, Müller RA (2013): Pervasive rightward asymmetry shifts of functional networks in autism spectrum disorder. *JAMA Psychiatry*.

https://doi.org/10.1001/jamapsychiatry.2013.382

- 54. Sun T, Patoine C, Abu-Khalil A, Visvader J, Sum E, Cherry TJ, *et al.*(2005): Early asymmetry of gene transcription in embryonic human left and right cerebral cortex. *Science* 308: 1794–8.
- Baron-Cohen S, Tsompanidis A, Auyeung B, Nørgaard-Pedersen B, Hougaard DM, Abdallah M, *et al.* (2019): Foetal oestrogens and autism. *Mol Psychiatry*. https://doi.org/10.1038/s41380-019-0454-9
- 56. Geschwind DH, Miller BL, DeCarli C, Carmelli D (2002): Heritability of lobar brain volumes in twins supports genetic models of cerebral laterality and handedness. *Proc Natl Acad Sci U S A* 99: 3176–81.
- 57. Geschwind N, Behan P (1982): Left-handedness: Association with immune disease, migraine, and developmental learning disorder. *Proc Natl Acad Sci U S A*. https://doi.org/10.1073/pnas.79.16.5097
- Herbert MR, Ziegler D a, Deutsch CK, O'Brien LM, Kennedy DN, Filipek P a, *et al.* (2005): Brain asymmetries in autism and developmental language disorder: a nested whole-brain analysis. *Brain* 128: 213–26.
- Nebel MB, Joel SE, Muschelli J, Barber AD, Caffo BS, Pekar JJ, Mostofsky SH (2014): Disruption of functional organization within the primary motor cortex in children with autism. *Hum Brain Mapp*. https://doi.org/10.1002/hbm.22188
- 60. Thompson A, Murphy D, Dell'Acqua F, Ecker C, McAlonan G, Howells H, et al. (2017): Impaired Communication Between the Motor and Somatosensory Homunculus Is Associated With Poor Manual Dexterity in Autism Spectrum Disorder. *Biol Psychiatry*. https://doi.org/10.1016/j.biopsych.2016.06.020

- Kana RK, Liu Y, Williams DL, Keller T a, Schipul SE, Minshew NJ, Just MA (2013): The local, global, and neural aspects of visuospatial processing in autism spectrum disorders. *Neuropsychologia* 51: 2995– 3003.
- 62. Gauthier I, Tarr MJ, Moylan J, Skudlarski P, Gore JC, Anderson AW
 (2000): The fusiform "face area" is part of a network that processes faces at the individual level. *J Cogn Neurosci*. https://doi.org/10.1162/089892900562165
- 63. Coffman MC, Anderson LC, Naples AJ, McPartland JC (2013): Sex Differences in Social Perception in Children with ASD. J Autism Dev Disord. https://doi.org/10.1007/s10803-013-2006-5
- 64. Lai M-C, Lombardo M V, Ecker C, Chakrabarti B, Suckling J, Bullmore ET, et al. (2014): Neuroanatomy of Individual Differences in Language in Adult Males with Autism. *Cereb Cortex*. https://doi.org/10.1093/cercor/bhu211
- 65. Wan CY, Marchina S, Norton A, Schlaug G (2012): Atypical hemispheric asymmetry in the arcuate fasciculus of completely nonverbal children with autism. *Ann N Y Acad Sci.* https://doi.org/10.1111/j.1749-6632.2012.06446.x
- 66. Mayo J, Chlebowski C, Fein D a, Eigsti I-M (2013): Age of first words predicts cognitive ability and adaptive skills in children with ASD. J Autism Dev Disord 43: 253–64.
- 67. Howlin P (2003): Outcome in high-functioning adults with autism with and without early language delays: implications for the differentiation between autism and Asperger syndrome. *J Autism Dev Disord* 33: 3–13.

Figure Legends

Figure 1 - Normative developmental changes for GM laterality in neurotypical males

Figure 1 captures the spatial representation of the voxelwise normative model in neurotypical (NT) males from the KCL site. The upper panel shows the beta values of laterality change across 6 to 30 years of age. The lower panel shows the actual prediction of GM laterality at ages 6 and 30. Blue colors indicate a shift towards leftward asymmetry, whereas red colors indicate a shift towards rightward asymmetry. The regression line depicts the predicted laterality values extracted from the language network based on neurosynth between 6 and 30 years of age along with centiles of confidence. These are based on the normative model maps thresholded with positive Rho-maps. Blue dots are the true values for NT males in KCL.

Figure 2 – Characterization of extreme laterality deviations

Abbreviations: ASD=autism spectrum disorder, NT=neurotypicals. The upper panel characterizes extreme rightward deviations (a), the lower panel extreme leftward deviations (b). The left panels show the percentage of extreme rightand leftward deviations from the normative model at each brain locus for each diagnostic group and gender separately. We depict loci where at least 2% of the subjects show overlaps. The violin plots on the right show the extreme deviations for each individual within each diagnostic and gender group. On average, individuals with autism show more extreme deviations than NT controls for both right- and leftward deviations.

Figure 3 – Relative importance of extreme laterality deviations

Abbreviations: LD=language delay, ADHD=attention-deficit hyper-activity disorder, CSS=calibrated severity scores on the ADOS-2. Based on the R package 'relaimpo' (https://cran.r-

project.org/web/packages/relaimpo/relaimpo.pdf), LD has the largest contribution to R^2 for extreme rightward deviations, whereas symptom severity has the largest contribution to R^2 for extreme leftward deviations.

Figure 4 – Extreme laterality deviations as a function of language delay

Abbreviations: LD=individuals with autism with language delay, noLD=individuals with autism without language delay, NT=neurotypicals. The upper panel characterizes extreme rightward deviations (a), the lower panel extreme leftward deviations (b). The left panels show the percentage of extreme right-and leftward deviations from the normative model at each brain locus for each diagnostic group and gender separately. We depict loci where at least 2% of the subjects show overlaps. The violin and bar plots on the right show that individuals with autism with LD show more extreme rightward deviations than the other two groups.

Figure S1 – Distribution of subjects across acquisition sites

Abbreviations: ASD=autism spectrum disorder, NT=neurotypicals. The bar plot shows the distribution of subjects by diagnostic group and sex across the five acquisition sites. Cambridge: autism males=34, autism females=16, NT males=19, NT females=9; KCL= autism males=103, autism females=33, NT males=42, NT females=25; Mannheim: autism males=22, autism females=6, NT males=25, NT females=10; Nijmegen: autism males=72, autism females=27, NT males=40, NT females=23; Utrecht: autism males=28, autism females=11, NT males=28, NT females=12.

Figure S2 – Summary of GM VBM laterality pre-processing pipeline

Abbreviations: GM=grey matter, VBM=voxel-based morphometry, DARTEL= Diffeomorphic Anatomical Registration using Exponentiated Lie algebra (40), LI=laterality index, I_{nref} =non-reflected GM images, I_{ref}= reflected GM images. Figure depicts the pre-processing pipeline for structural T1-weighted images. Note that the standard VBM pipeline is adjusted to meet the needs for laterality analyses, i.e., using symmetric registration.

Figure S3 - Normative developmental changes for GM laterality in neurotypical females

Supplementary Figure 3 captures the spatial representation of the voxelwise normative model in neurotypical (NT) females from the KCL site. The upper panel shows the beta values of laterality change across 6 to 30 years of age. The lower panel shows the actual prediction of GM laterality at ages 6 and 30. Blue colors indicate a shift towards leftward asymmetry, whereas red colors indicate a shift towards rightward asymmetry. The regression line depicts the

predicted laterality values extracted from the language network based on neurosynth between 6 and 30 years of age along with centiles of confidence. These are based on the normative model maps thresholded with positive Rho-maps. Blue dots are the true values for NT females in KCL.

Figure S4a – Normative developmental changes for GM laterality in neurotypical males using ComBat

Supplementary Figure 4a captures the spatial representation of the voxelwise normative model in neurotypical (NT) males across all sites. The upper panel shows the beta values of laterality change across 6 to 30 years of age. The lower panel shows the actual prediction of GM laterality at ages 6 and 30. Blue colors indicate a shift towards leftward asymmetry, whereas red colors indicate a shift towards rightward asymmetry.

The regression line depicts the predicted laterality values extracted from the language network based on neurosynth between 6 and 30 years of age along with centiles of confidence. These are based on the normative model maps thresholded with positive Rho-maps. Blue dots are the true values for NT males across all sites.

Figure S4b – Normative developmental changes for GM

laterality in neurotypical females using ComBat

Supplementary Figure 4b captures the spatial representation of the voxelwise normative model in neurotypical (NT) females across all sites. The upper panel shows the beta values of laterality change across 6 to 30 years of age. The lower panel shows the actual prediction of GM laterality at ages 6 and 30.

Blue colors indicate a shift towards leftward asymmetry, whereas red colors indicate a shift towards rightward asymmetry. The regression line depicts the predicted laterality values extracted from the language network based on neurosynth between 6 and 30 years of age along with centiles of confidence. These are based on the normative model maps thresholded with positive Rho-maps. Blue dots are the true values for NT females across all sites.

Figure S5 – Normative model by sex and site in NT individuals for the language network

The regression lines depict the predicted laterality values extracted from the language network based on neurosynth between 6 and 30 years of age along with centiles of confidence. These are based on the normative model maps thresholded with positive Rho-maps. Blue dots are the true values for NT males on the left and NT females on the right categorized by sites.

Figure S6 – Mean model accuracy 1

The figure shows the correlation (Rho) between true and predicted grey matter laterality values in NT controls and autism individuals.

Figure S7 – Mean model accuracy 2

The figure shows the root mean square error of true and predicted mean of grey laterality values in NT controls and autism individuals.

Figure S8 – Characterization of extreme laterality deviations using FDRcorrection Abbreviations: ASD=autism spectrum disorder, NT=neurotypicals,

LD=individuals with autism with language delay, noLD=individuals with autism without language delay. The figure depicts the replication of results when using FDR-correction as an alternative thresholding method. The upper panel characterizes extreme rightward deviations (a), the lower panel extreme leftward deviations (b). The left panels show the percentage of extreme right-and leftward deviations from the normative model at each brain locus for each diagnostic group and gender separately. We depict loci where at least 2% of the subjects show overlaps. The violin plots in the middle show the extreme deviations for each individual within each diagnostic and gender group. On average, individuals with autism show more extreme deviations than NT controls for both right- and leftward deviations. The barplots on the right show that individuals with autism with LD have more extreme-rightward deviations.

Figure S9 – Extreme laterality deviations as a function of language delay in matched subsamples

Abbreviations: LD=individuals with autism with language delay, noLD=individuals with autism without language delay, NT=neurotypicals. Replicated results are shown in a sample where individuals with autism with and without LD are matched for age and symptom severity. The upper panel characterizes extreme rightward deviations (a), the lower panel extreme leftward deviations (b). The violin and bar plots on the right show that individuals with autism with LD show more extreme rightward deviations than the other two groups.

Figure S10 – Overlap between ABIDE and EU AIMS

Abbreviations: ASD=autism spectrum disorder, NT=neurotypicals, ABIDE=Autism Brain Imaging Data Exchange. The figure depicts the replication of results in an independent dataset. The upper panel characterizes extreme rightward deviations (a), the lower panel extreme leftward deviations (b). The left panels show the percentage of extreme rightand leftward deviations from the normative model at each brain locus for each diagnostic group and gender separately where ABIDE and EU-AIMS LEAP show overlap. The violin plots on the right show the extreme deviations for each individual within each diagnostic and gender group in the ABIDE dataset. On average, individuals with autism show more extreme deviations than NT controls for both right- and leftward deviations

Table 1 – Demographic and clinical characterization of the LEAP sample

Abbreviations: ASD=Autism Spectrum Disorder; NT=neurotypical, M=males, F=females, ADI-R= Autism Diagnostic Interview-Revised, ADOS=Autism Diagnostic Observation Schedule, RRB=restricted, repetitive behavior, CSS=calibrated severity score, ADHD= attention-deficit hyper-activity disorder, LD=language delay.

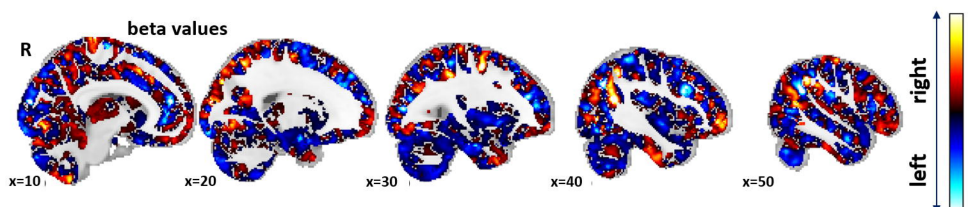
Table S1 – Summary of acquisition parameters across sites

 Table S2 - Demographic and clinical characterization of the ABIDE
 sample

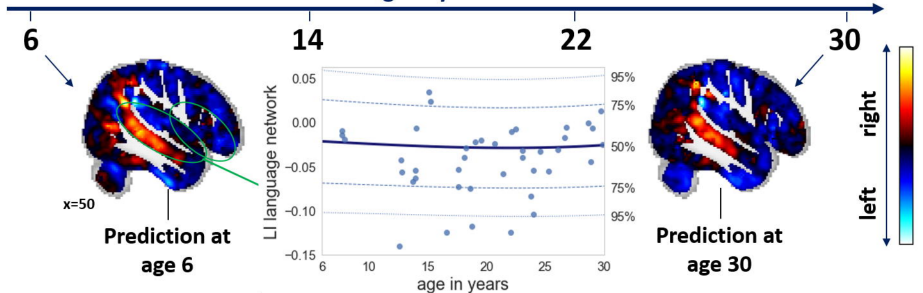
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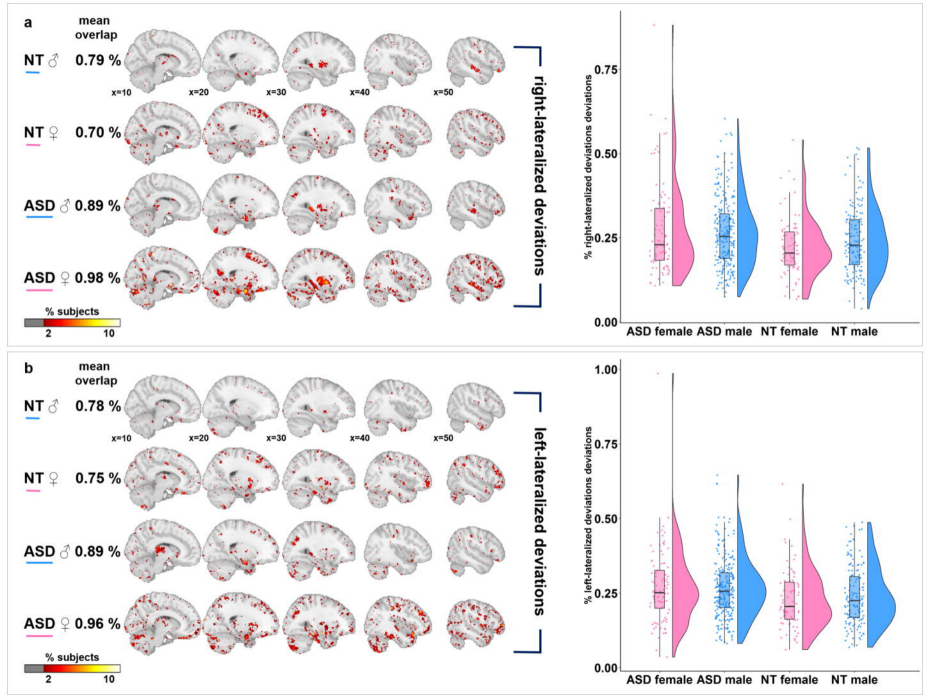
Abbreviations: ASD=Autism Spectrum Disorder; NT=neurotypical, M=males,

F=females, ADI-R= Autism Diagnostic Interview-Revised, ADOS=Autism Diagnostic Observation Schedule, RRB=restricted, repetitive behavior, CSS=calibrated severity score ^aFull-Scale IQ information was available for 384 ASD M, 87 ASD F, 435 NT M and 197 NT F. ^bVerbal IQ information was available for 330 ASD M, 76 ASD F, 382 NT M and 159 NT F. ^cPerformance IQ information was available for 335 ASD M, 78 ASD F, 398 NT M and 173 NT F. ^dADI-R Social Scores were available for 341 ASD M and 74 ASD F. ^eADI-R Communication and RRB Scores were available for 342 ASD M and 74 ASD F. ^fADOS-Gotham Social-Affect Scores were available for 132 ASD M and 28 ASD F. ^gADOS-Gotham RRB Scores were available for 134 ASD M and 29 ASD F. ^hADOS-Gotham Severity Scores were available for 134 ASD M and 30 ASD F.

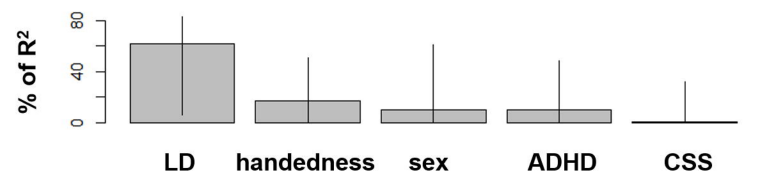


age in years





Rightward deviations



Leftward deviations



