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HYPERPERFUSION OF FRONTAL WHITE AND SUBCORTICAL GRAY MATTER IN AUTISM SPECTRUM DISORDER

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Abstract

Background: Our aim was to assess resting cerebral blood flow (rCBF) in children and adults with Autism Spectrum Disorder (ASD).

Methods: We acquired pulsed arterial spin labeling MRI data in 44 generally high functioning ASD simplex participants and 66 typically developing (TD) controls with comparable mean fullscale IQ's. We compared rCBF values voxel-wise across diagnostic groups and assessed correlations with symptom scores. We also assessed the moderating influences of participant age, sex, and IQ on our findings and the correlations of rCBF with NAA (N-acetylaspartate) metabolite levels.

Results: We detected significantly higher rCBF values throughout frontal white matter and subcortical gray in ASD participants. rCBF correlated positively with socialization deficits in regions where ASD hyperperfusion was greatest. rCBF declined with increasing IQ in the TD group, a correlation that was absent in ASD participants because their rCBF values were elevated across all IQ levels. rCBF in the ASD group correlated inversely with NAA metabolite levels throughout frontal white matter, with greater rCBF accompanying lower and increasingly abnormal NAA levels relative to TD controls.

Conclusions: These findings taken together suggest the presence of altered metabolism, likely of mitochondrial origin, and dysfunctional maintenance processes that support axonal functioning in ASD. These disturbances in turn likely reduce neural efficiency for cognitive and social

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functioning, and trigger compensatory responses from supporting glial cells, which subsequently increase rCBF to affected white matter. These findings, if confirmed, suggest cellular and molecular targets for novel therapeutics that address axonal pathology and bolster glial compensatory responses in ASD.

Keywords

Autism; Magnetic Resonance Imaging; Arterial Spin Labeling; white matter; glia; cerebral blood flow

INTRODUCTION

Regional cerebral blood flow (rCBF) is a surrogate measure of brain metabolism, because blood flow and metabolism are usually tightly coupled(1–5). Arterial Spin Labeling (ASL) provides absolute and reproducible measures of rCBF(6, 7) by magnetically labeling water in arterial blood as a diffusible tracer, analogous to the use of ^{15}O water in PET scanning(8). Unlike PET, ASL does not require the use of a radioactive tracer and therefore can be used in children, and it is a less costly and procedurally less complicated set of procedures. Moreover, because perfusion signals are obtained by pair-wise subtraction of adjacently acquired tagged and control images, ASL is less prone to motion artifact, low frequency physiological noise, and baseline drift than more conventional BOLD-based functional imaging(9). Finally, perfusion maps quantify rCBF at every voxel in the brain, whereas taskbased BOLD imaging quantifies neural activity only in locations that the task activates, often confounded by systematic biases in task performance across participants(10).

We are aware of only one previous preliminary study employing ASL in persons with Autism Spectrum Disorder (ASD) and typically developing (TD) controls, which had a limited sample size, included only children, and reported abnormalities (increased rCBF) only in cortical gray matter of the ASD group(11). PET metabolism studies and SPECT rCBF studies in ASD have had small numbers of participants, and some have either employed sedation or compared against atypically developing patient controls; most have interrogated only gray matter, and their findings have been rather inconsistent. Nevertheless, FDG PET studies generally have reported reduced metabolism in cortical gray matter (anterior cingulate, parietal(12, 13), and temporal(14–16) cortex) and increased metabolism in occipital(17, 18) and other cortical regions(19). One preliminary PET study reported hyperperfusion of basal ganglia and posterior cingulate cortex in ASD(20), whereas larger ¹⁸FDG PET studies have found reduced basal ganglia and thalamus metabolism $(21, 22)$. One FDG PET study assessed white matter metabolism explicitly, reporting higher metabolic rates that were greatest in prefrontal regions and internal capsule in 25 ASD compared with 55 TD adults during a verbal learning task(23). SPECT blood flow studies have generally reported reduced rCBF in temporal(24–31), parietal(25, 30, 31), and frontal $(26, 27, 29-33)$ cortices and basal ganglia $(28, 31, 33)$. Prior resting state functional MRI (fMRI) studies have generally reported long-range underconnectivity and local overconnectivity(34–45), suggesting the presence of axonal pathology in ASD.

We aimed to exploit some of the advantages of ASL to compare rCBF across a large sample of children and adults with ASD and age-, sex-, and IQ-matched TD controls. We also assessed the voxel-wise associations of ASL-based measures of rCBF and brain metabolism with MR spectroscopy-based measures of NAA (N-acetylaspartate)(46, 47) to understand how rCBF associates with this index of mitochondrial functioning(48–50), cellular metabolism(51), and neuronal abundance and viability(49, 50). Based on the prior metabolism and blood flow findings in ASD, we hypothesized a priori that we would detect reduced cortical and increased white matter rCBF. We also assessed the moderating effects of age, sex, and IQ on rCBF values in an effort to use rCBF values to parse the phenotypic heterogeneity of ASD. Finally, we assessed the voxel-wise correlation of rCBF with measures of NAA in ASD participants.

METHODS

Sample Ascertainment & Characterization

This was a subset of 110 participants(44 ASD,66 TD) drawn from a larger MRI study of 200 ASD (simplex cases) and TD controls(36, 46); the ASL pulse sequence was not available for the earliest recruited participants in the larger study. ASD participants were recruited through the Developmental Neuropsychiatry Clinic at Columbia University and community awareness events. Because screening of participants occurred prior to the publication of DSM 5 diagnostic criteria for ASD, screening was based on DSM-IV-TR criteria for PDD. Potential participants were required to have a previous clinical DSM-IV-TR(52) diagnosis of Pervasive Developmental Disorder (PDD) and a score >15 on either the Social Communication Questionnaire Lifetime Version(53) or Autism-spectrum Quotient(54). Exclusion criteria specific to ASD were known medical conditions associated with autism (e.g., Fragile-X syndrome, tuberous sclerosis). The final consensus diagnoses were established by a physician or clinical psychologist with established reliability by administering Module $3(n=16)$ or Module $4(n=25)$ of the Autism Diagnostic Observation Schedule(ADOS-G)(55). The revised algorithm was used for all modules(56, 57). Diagnoses for younger participants were also aided by administering the parent Autism Diagnostic Interview-Revised(ADI-R)(58)(n=17). In our final sample, 12 (27%) met DSM-IV-TR criteria for Autistic Disorder, 23 (52%) for Asperger's Disorder, and 8 (18%) for PDD-NOS. Because substantial evidence suggests that most individuals with DSM-IV-based PDD diagnoses also meet DSM 5 criteria for a diagnosis of ASD(59), for ease of reference we herein used the DSM 5 designation.

TD participants were recruited through flyers, online advertisements, and random sampling from a telemarketing list of local households eligible for participation, group-matched to ASD participants on age, sex, and ethnicity. TD controls participated after a clinical interview including the Kiddie Schedule for Affective Disorders and Schizophrenia Present and Lifetime Version for those under age 18(60) or the Structured Clinical Interview for DSM-IV Axis I Disorders for those age 18(61). Additional exclusion criteria for all participants included contraindications to MRI, a history of premature birth(<36 weeks), low birth weight(<2000g), birth complications or injury, prior head trauma with loss of consciousness, seizures, or chronic medical illness. The Institutional Review Board of the

New York State Psychiatric Institute approved the study procedures. An independent clinical monitor assessed the capacity for all adult ASD participants to provide informed consent. One participant was deemed to lack that capacity and therefore designated a surrogate to provide it. All children provided written assent.

Intelligence quotient(IQ) for 1 ASD participant was assessed using the Wechsler Intelligence Scale for Children(62) and 1 using the Wechsler Adult Intelligence Scale(63); all other ASD and TD participants were assessed using the Wechsler Abbreviated Scale of Intelligence(64). As all ASD participants had fluent speech, full-scale IQ was available for all. The Edinburgh Inventory assessed laterality of handedness(65), and Hollingshead's four-factor index estimated socioeconomic status(66).

MRI Scanning & Processing

All participants were unsedated for the scan. Details for scanning procedures, MRI pulse sequences, scan times, and image processing methods are provided in Supplementary Materials.

Statistical Analyses

All group-level statistical analyses employed a general linear model(GLM) at each voxel of the spatially normalized, individual rCBF maps, while covarying for age and sex, applying a False Discovery Rate (FDR) procedure at FDR=0.05 to control for false positives and colorcoding voxels on the T1 template where P-values survived the FDR procedure. All statistical models were hierarchically well-formulated, with any interaction terms including the component main effects. Our *a priori* hypothesis assessed the main effect of diagnosis on rCBF in all participants with no other covariates or interactions in the model. We also assessed the influences of FSIQ and medication use on our findings in post hoc analyses by covarying separately for FSIQ and psychotropic medication use and by applying again our model for a priori hypothesis testing while excluding any ASD participants who were taking psychotropic medications at the time of scan.

Secondary analyses assessed the moderating effects of age, sex, and FSIQ on group differences in separate models that included the respective interactions with their component main effects. These interactions assessed whether the slopes of the correlations differed across the ASD and TD control groups. In addition, within the ASD group we assessed the correlations of rCBF with Total and Domain scores on the developmentally appropriate ADOS module and with standardized Total and Subscale scores on SRS, while covarying for age and sex. We also assessed the voxel-wise correlations of rCBF with MRS-based measures of NAA metabolite levels as a continuous variable, and we compared rCBF values in the 6 ASD participants in whom we previously detected the presence of lactate with 30 ASD participants in whom we did not detect the presence of lactate.

RESULTS

Sample Characteristics

The ASD and TD samples did not differ significantly in age, sex, ethnicity, FSIQ, handedness, or socioeconomic status(Table 1). By design, all SRS domain scores were higher in the ASD sample (all $P<10^{-11}$), reflecting more severe social symptoms(Table 1). Fourteen of the 44 ASD participants (and no TD participants) were taking psychotropic medication at the time of scan: stimulants(n=5), antidepressants(n=7; includes SSRIs and SNRIs), lithium(n=2), anticonvulsants/mood stabilizers(n=1), antipsychotics(n=4), benzodiazepine(n=2), MAOI(n=1), and non-stimulant ADHD medication(n=2) (these total >n=14 because some were taking multiple medications).

Group Comparisons of rCBF

ASD compared with TD participants had higher rCBF values throughout large expanses of white matter, primarily in frontal regions comprising the centrum semiovale(CS), corona radiata(CR), and internal capsule(IC) bilaterally. They also had greater rCBF bilaterally in gray matter of the caudate (Cd) and lenticular nucleus (L) more dorsally, the ventral striatum(VS), orbitofrontal cortex(OFC), and amygdala(Am).

Correlations of rCBF with ASD Symptom Measures

Within the ASD group, rCBF correlated positively with the Total ADOS score in much of the same frontal white matter regions where a main effect of diagnosis was located(CS,CR), as well as within the hypothalamus, hippocampus, and middle temporal gyrus(Fig. 1). The social affect domain score of the ADOS almost exclusively contributed to this correlation, with little or no contribution from the repetitive behaviors domain score(Supplemental Figs.S22-S23). Of the various SRS subscale scores, social awareness correlated most robustly with rCBF measures, both within more dorsal white matter of the frontal lobe(CS), as well as in gray matter of the hypothalamus, hippocampus, middle temporal gyrus, basal ganglia(Cd,L,VS), amygdala, hippocampus, and brainstem(Supplemental Figs.S24-S28).

Moderators of Diagnosis Effect

Age significantly moderated the effects of diagnosis in the subgenual, pregenual, and dorsal ACC, inferior frontal gyrus, DLPFC, occipital gyrus, middle and inferior temporal gyrus, and amygdala(Fig. 2), with the effects deriving in each instance from a steeper inverse correlation of rCBF with age in the ASD group than in TD controls(Fig. 3).

Sex significantly moderated the effect of diagnosis, though localized almost exclusively to the subgenual and pregenual ACC and small portions of the caudate, ventral striatum, and amygdala(Fig. 2), with the effects deriving in each case from a reversal in the ASD group of the female>male rCBF differences present in TD controls(Fig. 3).

FSIQ also significantly moderated the effect of diagnosis within frontal white matter(CR,CS,IC), hypothalamus, and insular cortex(Fig. 2), with the effect in each case deriving from a steeper decline in FSIQ with increasing rCBF, and more broadly throughout

frontal white matter and subcortical gray matter, in the TD controls than in the ASD participants(Fig. 3).

Correlations of rCBF with MRS Metabolites

In the ASD and TD groups separately and combined, rCBF correlated inversely with NAA metabolite levels throughout frontal white matter, as well as in white matter of the internal and external capsules(Fig. 4). The 6 ASD participants who had detectable lactate in their brains, compared with the 30 who did not, had significantly lower rCBF in the white matter of the CC genu and posterior CS, and in gray matter of the DPFC and pregenual ACC. They also had higher rCBF values in the midcingulate cortex(Fig. 4).

Post Hoc Analyses

We conducted post hoc analyses to assess the effects of potential confounds on our findings. Covarying for FSIQ in our models had negligible effects on our findings(Supplemental Figs. S29-S31). Likewise, covarying for the use of psychotropic medications at the time of MRI scan had negligible effects, as did excluding from analyses those ASD participants taking psychotropic medications, though maps were overall somewhat less statistically significant because of reduced statistical power associated with fewer participants(Supplemental Figs.S32-S37). Covarying for total cortical gray matter or total white matter volumes also had no discernible effects on our findings(Supplemental Figs.S38-S39). Finally, we also conducted our primary analyses (shown in Fig. 1) separately in youth(≤ 21 years; N=20) and adults(>21 years; N=24); despite reduced statistical power, hyperperfusion was present in both groups, with the frontal locations most prominent in youth and dorsal parietal location more prominent in the adults(Supplemental Fig.S40). rCBF correlations with ADOS total scores were stronger in ASD adults, whereas SRS Awareness correlations were stronger in ASD youth(Supplemental Figs.S41-S42).

DISCUSSION

To the best of our knowledge, this is the first report of widespread hyperperfusion in ASD, and the first to report voxelwise correlations of rCBF with NAA metabolite concentrations throughout the brain.

Hyperperfusion in ASD

Hyperperfusion was present throughout frontal white matter (WM) and subcortical gray in ASD participants of all ages and in both sexes. rCBF correlated positively with measures of socialization deficits within the ASD group, including social affect domain scores on the ADOS and the social awareness subscale of the SRS, in regions where ASD hyperperfusion in group comparisons of rCBF were greatest.

Our findings of WM hyperperfusion are consistent with one prior FDG PET brain metabolism study reporting higher metabolic rates in the WM of 25 ASD compared with 55 TD adults performing a verbal learning task, with the greatest increases in prefrontal WM and anterior limb of the internal capsule(23). They are also consistent with findings from numerous MRI(67, 68) and DTI(69, 70) studies reporting volumetric and organizational

WM abnormalities in ASD, and with postmortem findings of fewer long axons, more short local axons, and thinner myelin(71, 72), which together suggest that compromised integrity of axons or oligodendrocytes may be the anatomical basis for reported long-range underconnectivity and local overconnectivity in functional MRI studies(34–45).

CNS WM consumes approximately one-third the energy of gray matter $(1-5)$ due to the greater energy efficiency of axon action potentials and especially WM synapses(73), with most WM energy devoted to maintaining resting potentials and various housekeeping tasks(73, 74). Hyperperfusion in ASD could derive from pathology in any one or a combination of these maintenance processes in neuronal axons, the most abundant cellular tissue in WM and its largest component by volume.

Oligodendrocytes, which comprise approximately 5% of all cells and 10–15% of all human glial cells in the brain(75, 76), ensheath WM axons with myelin to speed saltatory conduction(77). Their pathological functioning could alter myelination and, secondarily, WM metabolism and perfusion. Myelination is particularly important for finely calibrating conduction speeds and coordinating signals over long axons, which fMRI studies have suggested is a prominent disturbance in ASD(34–45). Most new myelin in adulthood is formed through an activity-dependent, plastic remodeling of the number, length, and thickness of preexisting myelin sheaths(37, 78–85). Approximately 20% of new myelin in adulthood also derives from the neuronal activity-dependent differentiation of oligodendrocyte precursor cells (OPCs) into new oligodendrocytes(85–87). Thus primary abnormalities in oligodendrocytes could produce altered WM metabolism and blood flow or, alternatively, abnormalities secondary to disturbances in axonal activity, whether from molecular or genetic defects in ASD or altered experiences associated with the illness, that in turn could trigger ongoing, activity-dependent myelin remodeling and its attendant increased metabolic demands.

Astrocytes, the most abundant glial cell type in the CNS, constitute nearly half the volume of WM and contribute significantly to WM metabolism and blood flow(88). They play an essential supporting role in OPC proliferation and differentiation, production of myelin(89), and formation of the blood-brain barrier. They take up potassium and glutamate released into the interstitial fluid to sustain and modulate high-frequency neuronal firing. Though not electrically excitable, they form a large network of astrocyte signaling through gap junctions that propagates Ca^{+2} and ATP waves over long-distances(90, 91). These properties of astrocytes are increasingly believed to play an important role in learning, memory, and intelligence(92), and genetic mutations affecting astrocyte functioning even produce ASDlike phenotypes of maturational delay, intellectual disability, and seizures(89). Altered astrocyte metabolism could account for our perfusion findings.

Microglia, the least abundant glial cell type, are the primary responders in a CNS cellular immune response. Their relative scarcity makes their metabolism itself an unlikely cause of hyperperfusion in ASD, though the inflammatory responses that they generate could increase perfusion. Excessive microglial cell number and activation have been reported in previous PET and postmortem studies of ASD(93–95), and the reactive oxygen species and

proinflammatory cytokines they generate could produce the impaired mitochondrial functioning suggested in prior studies(96), including one study of our participants(46).

Hyperperfusion was also present in the basal ganglia and amygdala. Though predominantly gray matter structures, they do contain axons, myelin, and glia(97, 98), and therefore a disturbance affecting only one of these cell components could produce hyperperfusion in both white and gray matter.

Interactions with Age

We detected a significantly steeper inverse correlation of rCBF with age in the ASD group than in TD controls throughout much of cortical gray matter, but most significantly in the ACC, DLPFC, amygdala, and several other cortical gray matter regions, with rCBF generally higher in children and either similar or lower in adults with ASD compared with controls(Fig. 2b). Thus the hyperperfusion detected in ASD across all ages in WM and subcortical gray was also present in cortical gray matter and amygdala, but primarily in ASD children. Prior ¹⁵O PET studies have reported declining cerebral metabolic rates(1–5) and blood flow $(1-3, 5)$ with age in cortical gray matter in healthy individuals; age correlates generally have not been reported in ASD. We detected an age-related decline of rCBF in healthy individuals only in occipital cortex. The decline in other cortical regions was in ASD participants, and in the direction toward values in healthy controls with advancing age, suggesting either that rCBF normalizes with age in cortical gray or that values close to normal in those regions may be a marker for ASD illness that persists into adulthood. By extension, higher rCBF values in children could also represent some kind of adaptive response to the presence of illness. Myriad cellular aging processes could account for the age-related decline in cortical rCBF in ASD, including: fewer neurons(99), dendritic arbors, or spines; reduced activity of the synaptic sodium/potassium pump and resting membrane potential(100); or less oxygen consumption and energy production from neuronal mitochondria in older compared with younger ASD participants(101).

Interactions with Sex

We detected a significant sex-by-diagnosis effect on rCBF in limbic regions, including subgenual ACC, ventral striatum, and amygdala, as well as parietal WM(Fig. 2). In each instance, rCBF was greater in TD females than TD males, but this sex difference reversed in ASD, driven by particularly high values in ASD males, higher than in either TD males or females(Fig. 2b). These findings suggest that, for reasons unknown, the perfusion abnormalities and cellular processes underlying them in ASD are disproportionately greater in males in limbic gray matter regions, perhaps contributing to the well-known male predominance in prevalence of ASD and to the more severe problems with externalizing and prosocial behaviors commonly found in ASD males(102, 103).

Interactions with IQ

We detected a significant IQ-by-diagnosis interaction effect on rCBF values in frontal WM and subcortical gray matter that derived from a significant decline in rCBF with increasing IQ in the TD group, a correlation that was absent in ASD participants(Fig. 2a&b). The hyperperfusion present in most members of the ASD group, though attenuated in younger

ASD participants, destroyed the normal association of IQ with rCBF in these regions(Fig. 3). The inverse correlation of resting rCBF with IQ in the TD controls is consistent with the neural efficiency hypothesis of intelligence, which posits that those with higher intelligence process information more efficiently, using less energy to accomplish any given task(104). Presumably this greater efficiency is present even in the "resting" brain that is not explicitly engaged in performing any task. The cellular and molecular determinants of greater efficiency are unknown, but could include improved synaptic or network connectivity(105), myelination(105), or maintenance of resting potentials and other housekeeping functions(106, 107). These need not be the same determinants as those that increase perfusion in ASD, but if they are the same, then disturbances in myelin remodeling or axon housekeeping functions are potential culprits in ASD, as both would require increased metabolism to support intellectual functioning at its highest potential.

Metabolite Correlations

rCBF throughout frontal WM in the ASD group correlated inversely with NAA metabolite levels. We reported elsewhere significantly lower NAA concentrations in the ASD group compared with controls(47). Thus, greater rCBF accompanied lower, and increasingly abnormal, NAA levels. As findings from prior studies indicate that reduced NAA concentrations represent a reduced density or functioning of axonal mitochondria(49, 50), then reduced frontal WM mitochondria could suggest that increased frontal WM rCBF and metabolism represents a compensatory process in ensheathing glia to maintain the energy state of WM axons.

The possibility of compensation is bolstered by evidence that glia support the formidable metabolic housekeeping demands of myelinated axons, which are challenged by sparse mitochondria and slow passive diffusion of nutrients across nodes of Ranvier(108). Myelinating oligodendrocytes couple with associated axons to support neuronal metabolism under stress(109), in part through release of exosomes containing trophic substances(73, 74, 109, 110), and in part through shuttling of lactate and pyruvate to axons to serve as substrates for aerobic ATP production(111). Axonal firing stimulates further oligodendrocyte production of lactate and pyruvate through release of glutamate, which activates oligodendrocyte NMDA receptors to stimulate uptake of glucose and glycolysis, thereby maintaining energy supplies to axons(89, 112, 113). Finally, the myelin sheath has been proposed as an ectopic site of glucose combustion and ATP generation for use by myelinated axons(74, 114–116), though direct experimental and mathematical modeling support for ATP generation in myelin sheaths is lacking(117).

The positive correlations of rCBF with measures of social impairment we observed suggest that this putative compensatory response is either unsuccessful or incomplete, because fully successful compensation would produce an inverse correlation. In the presence of unsuccessful or incomplete compensation, however, greater reductions of NAA and axonal metabolism would produce more severe symptoms and a proportionately greater compensatory response in glial cell metabolism – i.e., it would produce the observed positive associations of social impairment with rCBF. Alternatively, sustained increases in myelin synthesis by oligodendrocytes in ASD WM could deplete the associated axons of

NAA, because more myelin synthesis would require transportation of more NAA from neurons to oligodendrocytes for cleavage and use of its acetate moiety(118, 119). Increased myelin synthesis therefore could account for both reduced NAA in axons but increased rCBF in metabolic support of myelin synthesis. Human postmortem and imaging studies, however, generally have not provided evidence for excessive, ongoing myelination in ASD.

Limitations

Though our sample comprised ASD participants with a wide range of intellectual and social functioning, mean IQs were by design comparable to TD participants to limit the potential confounding effects of IQ differences on our findings. This sample selection, however, limits the generalizability of our findings to persons who are more severely affected with ASD, at least with respect to language and cognitive functioning. In addition, a larger sample size would improve statistical power and the stability of interaction effects. Longitudinal imaging studies are needed to support developmentally based interpretations of our findings(10).

Conclusion

Extensive resting frontal WM hyperperfusion in ASD suggests increased metabolic demands of either axons or glia or both. Positive correlations of rCBF with measures of social impairment in the same WM regions suggest that either increased perfusion partially generates those symptoms or is a proportionate response to their presence. The altered associations of rCBF with IQ moreover suggests impaired neural efficiency in frontal WM. The combination of reduced NAA levels with proportionally greater rCBF values in frontal WM suggests that glial cells are attempting unsuccessfully or incompletely to compensate for reduced axonal metabolism in ASD. The most parsimonious explanation for all these findings is the presence of axonal pathology, including reduced NAA and altered mitochondrial metabolism, that disrupts one or more components of the housekeeping processes supporting axonal functioning, thereby reducing the efficiency of information processing. These disruptions trigger compensatory responses from supporting glial cells to support axon metabolism and proportionately increase rCBF to affected WM. The compensatory response is only partial, however, and in the ASD group disrupts the normal association of greater efficiency of neural processing with increasing IQ observed in the TD group. These findings, if confirmed, suggest cellular and molecular targets for studies of axonal pathology and development of novel therapeutics to bolster glial compensatory responses in ASD.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Group Comparisons and Symptom Correlations

The right sides of the images correspond to the right side of the brain. All analyses control for the effects of age and sex, and the significance levels in all maps are FDR-corrected for the number of statistical comparisons.First Row Shown is the T1-weighted anatomical template to which rCBF values for all participants were mapped at transaxial slice levels positioned parallel to the Anterior Commissure-Posterior Commissure (AC-PC) line, and corresponding to the positioning of the statistical maps shown in the other rows. The zvalues represent slice level (in millimeters) in the Talairach coordinate system. Second Row This shows the statistically significant differences in rCBF values between the ASD group and TD controls while covarying for age and sex, displayed at a threshold of P<0.05 after correction for multiple comparisons. Voxels in red indicate significantly increased rCBF, and blue voxels reduced rCBF, in ASD relative to controls. Perfusion values at rest in the ASD group were higher bilaterally throughout white matter of all the frontal lobe, internal capsule, and dorsal parietal lobe, and in gray matter of the basal ganglia, thalamus, and amygdala.

Third & Fourth Rows Red and blue voxels represent, respectively, significant positive or inverse correlations of ADOS total scores and SRS social affect scores with rCBF values in the ASD group, after FDR correction for multiple comparisons.

Abbreviations Am: amygdala; BS: brainstem; CC: corpus callosum; Cd: caudate; CR: corona radiata; CS: centrum semovale; H: hippocampus; IC: internal capsule; IF: inferior frontal gyrus; IT: inferior temporal gyrus; L: lenticular nucleus; MB: midbrain; MT: middle temporal gyrus; OFC: orbitofrontal cortex; OR optic radiations; PCC: posterior cingulate cortex; ST: superior temporal gyrus; Th: thalamus; VS: ventral striatum

Figure 2. Modifying Effects of Age, Sex, and FSIQ

the significance levels in all maps are FDR-corrected for the number of statistical comparisons Left Panel In the first two columns are maps showing the significance of age correlations with resting rCBF at each voxel in the ASD and TD groups separately, while covarying for sex. Red and blue voxels represent, respectively, significant positive or inverse correlations of age with rCBF values. The third column shows effects of the interaction of age with diagnosis on rCBF at each voxel, or the voxels where the correlations of age with rCBF differ significantly across the ASD and TD groups, while covarying for sex and including the main effects of age and diagnosis in a hierarchically well formulated statistical model.

Middle Panel In the first two columns are maps showing the significance of sex difference in rCBF at each voxel in the ASD and TD groups separately, while covarying for age. Red and blue voxels represent significantly higher rCBF values in males or females, respectively. The third column shows effects of the interaction of sex with diagnosis on rCBF at each voxel, or the voxels where sex differences in rCBF differ significantly across the ASD and TD groups, while covarying for age and including the main effects of sex and diagnosis in a hierarchically well formulated statistical model.

Right Panel In the first two columns are maps showing the significance of correlations of FSIQ (full scale IQ) with resting rCBF at each voxel in the ASD and TD groups separately, while covarying for age and sex. Red and blue voxels represent, respectively, significant positive or inverse correlations of FSIQ with rCBF values. The third column shows effects of the interaction of FSIQ with diagnosis on rCBF at each voxel, or the voxels where the correlations of FSIQ with rCBF differ significantly across the ASD and TD groups, while covarying for age and sex and including the main effects of FSIQ and diagnosis in a hierarchically well formulated statistical model.

Abbreviations Am: amygdala; Cd: caudate; CR: corona radiata; CS: centrum semovale; dACC: dorsal anterior cingulate cortex; DLPFC: dorsolateral prefrontal cortex; Hy:

hypothalamus; IC: internal capsule; IF: inferior frontal gyrus; Ins: insula; IT: inferior temporal gyrus; LG: lingual gyrus; MO: middle occipital gyrus; OFC: orbitofrontal cortex; OR optic radiations; pgACC: pregenual anterior cingulate cortex; PCC: posterior cingulate cortex; sgACC: subgenual anterior cingulate cortex; ST: superior temporal gyrus; VS: ventral striatum

Figure 3. Data Displays for Age, Sex, and FSIQ Effects

These are shown for representative regions indicated from a slice taken from Figure 2, shown above each plot in the statistical map. The line leading from the regional label to the image indicates the voxels with the images that were sampled to generate the data shown in the scatterplots. The scatterplots illustrate the age-related decline in rCBF in most cortical regions for ASD and the inverse correlation of rCBF with full scale IQ (FSIQ) in the TD but not ASD group. Bar graphs for sex effects demonstrate in ASD participants the reversal of female>male sex differences present in the TD group. Error bars represent standard error.

Figure 4. Correlations of rCBF with NAA Metabolite Levels and Lactate

Top 3 Rows show the voxel-wise correlations of NAA level with rCBF in the ASD (N=36, 29 males, 7 females, mean age 25.6 years) and TD (N=63, 48 males, 15 females, mean age 22.3 years) groups combined (top row) and separately, while covarying for age and sex. Significance levels are FDR-corrected for the number of statistical comparisons. Primarily inverse correlations (blue voxels) are present in the ASD group in frontal white matter of the centrum semiovale (CS) and corona radiata (CR), internal capsule (IC), and external capsule (EC), and in the midcingulate cortex (MCC) and posterior cingulate cortex (PCC). In the TD group, inverse correlations are seen in similar locations, but are smaller in spatial extent and weaker in statistical significance; positive correlations (red voxels) in the TD group are scattered throughout cortical gray matter (insula: Ins; posterior cingulate cortex: PCC) and inverse correlations are present in frontal white matter (CS, CR). Bottom Row The 6 ASD participants with detectable lactate compared with the 30 without detectable lactate had significantly lower rCBF values (blue voxels) in the dorsal prefrontal cortex (DPFC), pregenual anterior cingulate cortex (pgACC), posterior cingulate cortex (PCC), genu of the corpus callosum (gCC), and posterior white matter of the centrum semiovale (CS) and parieto-occipital (PO) region.

Table 1.

Participant Characteristics

 a^a ADOS scores were available for 41 ASD participants

 b SRS scores were available for 38 ASD and 49 TD participants

ADOS=Autism Diagnostic Observation Schedule; Restrict & Repet Beh=Restricted and Repetitive Behaviors SRS=Social Responsiveness Scale; Awareness=Social Awareness; Cognition=Social Cognition; Communication=Social Communication; Motivation=Social Motivation; Mannerisms=Restricted Interests and Repetitive Behaviors SES=socioeconomic status measured with the Hollingshead; this was available in 31 ASD and 56 TD participants FSIQ was available in 43 ASD and 65 TD participants

RMS=Root Mean Square motion estimate during the ASL scan