Novel Applications of Surface Based Morphometry and Pattern Classification in Autism Spectrum Disorders

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Novel Applications of Surface Based Morphometry and Pattern Classification in Autism Spectrum Disorders

Thesis for the Degree of Doctor of Philosophy

by

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Abstract

Autism spectrum disorder (ASD) is a lifelong, behaviorally defined neurodevelopmental condition that is characterized by deficits in social communication, interaction, and repetitive behaviors. These behavioral symptoms are associated with atypical brain structure, function, and connectivity. The studies that comprise this thesis employed structural magnetic resonance imaging (MRI) to address aims in three areas of ASD research. First, we examined a novel neuroimaging feature based on signal intensity contrast between grey and white matter to quantify atypical microstructure at the grey-white matter boundary in ASD. We found reduced tissue contrast at the grey-white matter boundary among adults with ASD when compared typically developing (TD) controls. This result indicates that measures of tissue contrast may serve as an in vivo proxy measure of atypical cortical microstructure that has previously been reported in histological studies. Second, we trained multivariate pattern recognition models to identify individuals with ASD based on measures of cortical morphometry, and examined the predictive value of these models in a representative clinical sample. We demonstrated that these models have modest ability to distinguish cases from controls in the research setting. Only one model that was based on measures of grey-white matter tissue contrast identified individuals with and without ASD diagnoses at high overall accuracy (81%) in the clinical setting. However, this model did not provide significant accuracies above chance in the research setting, and therefore these results should be considered as preliminary and suggestive only. Third, we established normative models of phenotypic diversity in brain structure associated with biological sex in a sample of TD males and females which was subsequently applied to males and females with ASD. Across different morphometric features, females with ASD displayed a significant shift towards a more male-typical presentation of the brain. Sample probabilities for ASD also increased with predicted probabilities for male-typical brain phenotypes across both sexes. These studies highlight advances in the field of structural neuroimaging research in areas of feature development, clinical translation, and efforts to understand the modulating role of biological sex on the prevalence of ASD. Taken together, the work presented within this thesis thus constitutes an important step toward establishing translational imaging tools for ASD that may one day be applied in the clinical setting.
# Table of Contents

Abstract .................................................................................................................. 2
Table of Contents .................................................................................................... 3
Table of Figures ......................................................................................................... 7
Table of Tables ......................................................................................................... 9
Acknowledgments ..................................................................................................... 10
Declaration of Authorship ......................................................................................... 11
Abbreviations ........................................................................................................... 12

## Chapter 1: Introduction ......................................................................................... 13
1.1 Autism Spectrum Disorder ................................................................................. 13
1.2 Sex Differences in Autism Spectrum Disorder .................................................. 16
1.3 The Etiology of Autism Spectrum Disorder ....................................................... 18
  1.3.1 Environmental Factors .............................................................................. 18
  1.3.2 Genetic Factors .......................................................................................... 19
  1.3.3 Competing Aetiological Theories ............................................................... 20
1.4 The Structure of the Brain in Autism Spectrum Disorder .................................. 22
  1.4.1 Volumetric Differences in the Autistic Brain .............................................. 24
  1.4.2 Cortical Morphometry of the Autistic Brain .............................................. 25
1.5 Beyond Group Differences: Pattern Recognition in Autism Spectrum Disorder ................................................................................................................................. 31
  1.5.1 Multivariate Pattern Classification in Autism Spectrum Disorder ............. 32
  1.5.2 Towards Translational Tools ..................................................................... 37
1.6 The Current Thesis: Aims and Hypotheses ....................................................... 37
  1.6.1 Study 1: In Vivo Evidence of Reduced Integrity of the Grey-White Matter Boundary in Autism Spectrum Disorder ................................................................. 37
  1.6.2 Study 2: Clinical Validation of Pattern Classification Models for Autism Spectrum Disorder ........................................................................................................ 38
  1.6.3 Study 3 and 4: Association Between the Probability of Autism Spectrum Disorder and Normative Sex-Related Phenotypic Diversity in Brain Structure ................................................................................................................................. 38
1.7 References ......................................................................................................... 40

## Chapter 2: Magnetic Resonance Imaging and Analysis Methods .......................... 56
2.1 Principles of Nuclear Magnetic Resonance ....................................................... 56
2.2 Magnetic Characteristics of Tissue ..................................................................... 58
2.3 Magnetic Resonance Images ............................................................................. 62
  2.3.1 Basic Magnetic Resonance Imaging Pulse Sequences ................................ 62
  2.3.2 Spatial Encoding and Image Reconstruction of Magnetic Resonance Signal .............................................................................................................................. 65
2.4 Driven Equilibrium Single Pulse Observation of $T_1$ and $T_2$ Mapping .......... 66
2.5 Cortical Surface Reconstruction using Freesurfer Software ............................. 67
2.6 Mass Univariate Magnetic Resonance Imaging Analyses ................................ 70
  2.6.1 The General Linear Model ....................................................................... 70
  2.6.2 Correcting for Multiple Comparisons ........................................................ 73
2.7 Multivariate Pattern Classification ................................................................... 74
  2.7.1 The Support Vector Machine ..................................................................... 76
  2.7.2 Gaussian Process Classifiers ...................................................................... 79
  2.7.3 Model Performance and Validation for Multivariate Pattern Classification .......................................................................................................................... 85
  2.7.4 Feature Extraction and Selection ................................................................ 86
  2.7.5 Model Visualization .................................................................................... 88
2.8 References ......................................................................................................... 90
### Table of Figures

<table>
<thead>
<tr>
<th>Figure</th>
<th>Description</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.1</td>
<td>The Autism Spectrum</td>
<td>14</td>
</tr>
<tr>
<td>1.2</td>
<td>Multifactorial Liability Model of Autism Spectrum Disorder</td>
<td>17</td>
</tr>
<tr>
<td>1.3</td>
<td>Brain Regions Indicated in Autism Spectrum Disorder</td>
<td>23</td>
</tr>
<tr>
<td>1.4</td>
<td>Development of the Human Cerebral Cortex</td>
<td>26</td>
</tr>
<tr>
<td>2.1</td>
<td>Interactions between RF Energy and Magnetic Nuclei in a Magnetic Field</td>
<td>59</td>
</tr>
<tr>
<td>2.2</td>
<td>Longitudinal Magnetization</td>
<td>60</td>
</tr>
<tr>
<td>2.3</td>
<td>Transverse Magnetization</td>
<td>61</td>
</tr>
<tr>
<td>2.4</td>
<td>Development of $T_1$ Contrast between Two Tissues</td>
<td>62</td>
</tr>
<tr>
<td>2.5</td>
<td>Development of Image Contrast in a Spin-Echo Sequence</td>
<td>64</td>
</tr>
<tr>
<td>2.6</td>
<td>Development of $T_1$ Contrast in an Inversion Recovery Sequence</td>
<td>65</td>
</tr>
<tr>
<td>2.7</td>
<td>The Support Vector Machine</td>
<td>77</td>
</tr>
<tr>
<td>2.8</td>
<td>The Gaussian Process in a One Dimensional Regression Problem</td>
<td>82</td>
</tr>
<tr>
<td>2.9</td>
<td>The Gaussian Process Classifier</td>
<td>83</td>
</tr>
<tr>
<td>2.10</td>
<td>Nested Parameter Tuning via Cross Validation</td>
<td>89</td>
</tr>
<tr>
<td>3.1</td>
<td>Grey and White Matter Signal Intensity Sampling Procedure</td>
<td>100</td>
</tr>
<tr>
<td>3.2</td>
<td>Regions of Decreased Grey-to-White Matter Signal Intensity Percent Contrast in ASD</td>
<td>101</td>
</tr>
<tr>
<td>3.3</td>
<td>Regional Differences in Grey and White Matter Signal Intensities in ASD</td>
<td>103</td>
</tr>
<tr>
<td>4.1</td>
<td>Support Vector Machine Predictions Within the Training Set</td>
<td>117</td>
</tr>
<tr>
<td>4.2</td>
<td>Gaussian Process Classification Predictions within the Training Set</td>
<td>118</td>
</tr>
<tr>
<td>4.3</td>
<td>Support Vector Machine Predictions within the Training Set using GLM based Feature Selection</td>
<td>118</td>
</tr>
<tr>
<td>4.4</td>
<td>Gaussian Process Classification Predictions within the Training Set using GLM based Feature Selection</td>
<td>119</td>
</tr>
<tr>
<td>4.5</td>
<td>Support Vector Machine Predictions within the Testing Set</td>
<td>120</td>
</tr>
<tr>
<td>4.6</td>
<td>Gaussian Process Classification Predictions within the Testing Set</td>
<td>121</td>
</tr>
<tr>
<td>4.7</td>
<td>Gaussian Process Classification Predictions within the Testing Set using measures of Tissue Contrast and GLM based Feature Selection</td>
<td>122</td>
</tr>
<tr>
<td>4.8</td>
<td>Accuracy as a Function of Number of Included Patients in the Classification of Psychiatric and Neurodevelopmental Conditions</td>
<td>123</td>
</tr>
<tr>
<td>5.1</td>
<td>Gaussian Process Classification of Biological Sex</td>
<td>136</td>
</tr>
<tr>
<td>5.2</td>
<td>Regions of Decreased Grey-to-White Matter Signal Intensity Percent Contrast in ASD</td>
<td>137</td>
</tr>
<tr>
<td>5.3</td>
<td>Regional Differences in Grey and White Matter Signal Intensities in ASD</td>
<td>138</td>
</tr>
<tr>
<td>6.1</td>
<td>Gaussian Process Classification of Biological Sex</td>
<td>155</td>
</tr>
<tr>
<td>6.2</td>
<td>Probability for ASD as a Function of Normative Sex-Related Phenotypic Variance in Brain Morphology</td>
<td>157</td>
</tr>
<tr>
<td>6.3</td>
<td>Prediction Maps for High and Low ASD Probability Males and Females</td>
<td>158</td>
</tr>
</tbody>
</table>
Figure 6.4, Group by Sex Interaction Effects in Cortical Morphometry. ............... 160
Table of Tables

Table 1.1, Multivariate Pattern Classification Studies of Autism Spectrum Disorder ........................................................................................................................................34

Table 3.1, Participant Demographics .........................................................................................................................................................................................99
Table 3.2, Clusters of Significant Reductions in Grey White Matter Percent Contrast and Increases in Grey Matter Intensity in ASD ...................................... 102

Table 4.1, Participant Demographics (Research Sample) .................................................................................................................................................111
Table 4.2, Participant Demographics (Clinical Sample) .........................................................................................................................................................112
Table 4.3, Classification Accuracies within Subgroups of the Clinical Sample ..................................................................................................................................122

Table 5.1, Participant Demographics ..........................................................................................................................................................................................134

Table 6.1, Participant Demographics and Global Brain Measures ..................................................................................................................................149
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Declaration of Authorship

I, Derek Andrews, declare that this thesis and the work presented in it are my own and have been generated by me as the result of my own original research.

I confirm that:

1. This work was done wholly or mainly while in candidature for a research degree at King’s College London;
2. Where I have consulted the published work of others, this is always clearly attributed;
3. Where I have quoted from the work of others, the source is always given;
4. I have acknowledged all main sources of help;
5. Parts of this work have been published or are currently in submission as: [see list references below]. In cases where I was not lead author my contribution to the work has been detailed.

Signed:

Date: June 21st, 2017


Abbreviations

ADD: Attention-Deficit Disorder
ADI: Autism Diagnostic Interview
ADOS: Autism Diagnostic Observation Schedule
ADHD: Attention-deficit hyperactivity disorder
ASD: Autism Spectrum Disorder
CT: Cortical Thickness
CURV: Mean radian curvature
CV: Cross Validation
DSM: Diagnostic and Statistical Manual for Mental Disorders
DTI: Diffusion Tensor Imaging
FA: Fractional anisotropy
fMRI: Functional Magnetic Resonance Imaging
FWHM: Full width at half maximum
GLM: General linear model
GM: Grey Matter
GP: Gaussian Process
GPC: Gaussian Process Classification
GWR: Grey:White Matter signal contrast ratio
GW: Grey:White Matter percent contrast
ICD: International Statistical Classification of Diseases
iPSCs: induced pluripotent stem cells
JACOB: jacobian metric distortion
lGI: Local gyrification index
LOO-CV: Leave one out cross validation
MD: Mean Diffusivity
MMR: Measles Mumps & Rubella
MRI: Magnetic Resonance Imaging
MRS: Magnetic Resonance Spectroscopy
MVPC: Multivariate pattern classification
OCD: Obsessive-compulsive disorder
ODD: Oppositional defiant disorder
RFT: Random Field Theory
ROI: Region of interest
rsMRI: Resting state Magnetic resonance imaging
SA: Surface Area
SBM: Surface Based Morphometry
SE: Sensitivity
SP: Specificity
sMRI: Structural Magnetic Resonance Imaging
SULC: Sulcal depth
SVM: Support Vector machine
TD: Typically Developing
VBM: Voxel Based Morphometry
WM: White Matter
Chapter 1: Introduction

Autism spectrum disorder (ASD) is a complex neurodevelopmental condition. While progress continues to be made in understanding the neurological origins of ASD the underlying neurodevelopmental mechanisms and biological markers of the condition remain unclear. The broad aim of this thesis is to advance our knowledge of ASD in a way that offers real world translational benefits to patients through applications of structural brain imaging. More specifically the studies that compose this work focus on (i) novel ways to measure atypical brain structure in ASD, (ii) testing the efficacy of measures of brain structure to identify ASD within the clinic, and (iii) examining how biological sex/gender may modulate risk for the condition.

I begin this introductory chapter by introducing the behavioral ASD phenotype and associated diagnostic criteria. The role of biological sex/gender in ASD will then be discussed followed by current etiological theories for the condition. I will then discuss evidence for atypical brain structure in ASD with a focus on cortical magnetic resonance imaging (MRI) findings and review studies that apply particular types of ‘machine learning’ methods to classify ASD using MRI scans. Lastly, I will outline the aims of the three studies, which compose the body of this thesis.

1.1 Autism Spectrum Disorder

Autism spectrum disorder (ASD) is a lifelong behaviorally defined neurodevelopmental condition characterized by deficits in social interaction and communication in addition to the presentation of repetitive and/or restricted behaviors (World Health Organization, 2004). The earliest reports of the ASD phenotype are attributed to Kanner (1943) and Asperger (1944). In these foundational works Kanner and Asperger both detail case studies of children in which typical autistic behaviors are observed. By definition ASD is phenotypically heterogeneous, encapsulating a spectrum of individuals whose behavioral deficits range from severe impairment (e.g. learning disability, non-verbal) to high functioning (e.g. average or above average IQ, relatively minor impairments to social communication) (Wing, 1997). In addition to the core features, individuals with ASD experience high rates of comorbidity with other psychiatric and medical conditions including; social anxiety disorder, attention-deficit disorder (ADD), attention-deficit hyperactivity disorder (ADHD), oppositional defiant disorder (ODD), obsessive-compulsive disorder (OCD), specific phobia,
gastrointestinal problems, and epilepsy (Simonoff et al., 2008, Leyfer et al., 2006, Amiet et al., 2008) (Figure 1.1).

Figure 1.1, The Autism Spectrum (autismspeaks): Autism Spectrum Disorder (ASD) is defined by core symptoms (red circles). However, the condition is often accompanied by several associated neurological (dark grey circles) and systemic (gold circles) issues. Individuals with ASD also experience increased rates of related comorbid disorders (light grey circles) and intellectual disability (dotted line).

Large population studies carried out around the globe report ASD prevalence to range from 0.5% to 2% in the general population. Over the past few decades it has been widely speculated that ASD prevalence is increasing. However, empirical evidence suggests that these increases can largely be attributed to growing awareness of the condition and greater accessibility of diagnostic services (Mazumdar et al., 2013, Hansen et al., 2015).

As a behaviorally defined disorder the diagnosis of ASD is reliant on parental interviews and/or assessments and observations of patient’s behaviors. A clinical ASD diagnosis can be granted by meeting the diagnostic criteria outlined in the
The clinical diagnostic criterion outlined by the ICD-10 encompasses significant phenotypic heterogeneity. While there is extensive evidence to suggest that ASD is associated with several genetic and neurobiological markers (Ruggeri et al., 2013, Ecker and Murphy, 2014) currently there are no reliable diagnostic biomarkers validated for clinical use in the condition. Therefore, two ‘gold-standard’ assessment tools may be used as additional confirmation of a diagnosis of ASD, namely the *Autism Diagnostic Interview-Revised* (ADI-R) (Lord et al., 1994) and *Autism Diagnostic Observation Schedule* (ADOS) (Lord et al., 2000). Both these measures require administration by an experienced clinical interviewer.

The ADI-R is a clinical structured caregiver interview appropriate for children and adults with a mental age of 18 or more months. The ADI-R provides a diagnostic algorithm for ASD as described by the ICD-10 and focuses on probing behaviors in the three main symptom areas of social interaction, communication, and repetitive/restricted behaviors. The 93-item interview covers the patient’s full developmental history, and usual lasts 1-2 hours. Questions focus on current behavior as well as a period between the ages of 4 and 5 years when several behaviors associated with ASD are likely to be most evident. Exceptions to this are questions...
regarding group play (4-10 years), reciprocal friendships (5+ years), and circumscribed interests (3+ years) that are appropriate only for particular ages. As ASD is a lifelong neurodevelopmental condition, meeting ADI-R criteria for a diagnosis is key for establishing a childhood origin of ASD, particularly in diagnoses of adults on the spectrum.

In contrast to the ADI-R, the ADOS involves a series of semi-structured assessments of communication, social interaction, play and/or imaginative thinking scenarios conducted with the patient. Four modules of the ADOS are available to allow for administration across the spectrum of impairment observed in ASD. The choice of module depends on the individual’s competency in using phrase speech (i.e. non-echoed, meaningful word combinations that are spontaneous, contain at least three words, and sometimes a verb), verbal fluency, and age. As the ADOS is used to assess current symptom presentation it is considered a gold standard metric not only for diagnosis but also for quantification of current ASD symptoms.

1.2 Sex Differences in Autism Spectrum Disorder

Of significant note is the preponderance of males with ASD. Based on large-scale population studies conducted around the globe currently it is estimated that ASD is 2-5 times more prevalent in males than females (Lai et al., 2017). This male bias in ASD diagnoses may partly be related to differences in the ASD phenotype in females who have been shown to display less severe impairments in the repetitive behaviors domain and improved skills in coping with social deficits (Szatmari et al., 2012, Mandy et al., 2012). Such phenotypic differences between males and females with ASD have likely contributed to the historical view of ASD as a ‘male condition’ and subsequent under diagnosis of females on the spectrum.

However, as awareness of females with ASD increases population studies continue to show a male bias in prevalence, albeit in a lower range of 1.7-2.5 males for every 1 female with the condition (Kim et al., 2011, Mattila et al., 2011, Idring et al., 2012). This indicates that in addition to male biased diagnostic criteria and tools, sex differentiated developmental processes may have a modulating effect on the risk for ASD. For example, multifactorial liability models propose that liability for ASD is
distributed throughout the population with males and females experiencing different liability thresholds for the development of ASD (Werling and Geschwind, 2013) (Figure 1.2).

Figure 1.2 Multifactorial Liability Model of Autism Spectrum Disorder (Werling and Geschwind, 2013, Reich et al., 1975): Within this model liability for ASD is distributed throughout the population. However female specific ‘protective factors’ shift liability away from and male-specific ‘risk factors’ towards the minimum total liability needed for an individual to develop ASD.

Multifactorial liability models of ASD propose so-called ‘male risk’ and/or ‘female protective’ factors inherent to the sexes that shift liability toward or away the minimum liability threshold required for an ASD phenotype to manifest. Genetic studies have provided some evidence for a ‘female protective effect’ reporting increased genetic load for ASD risk genes among siblings of females with the condition than siblings of males with ASD (reviewed in (Werling, 2016)). Beyond well-established findings of increased brain size in males (Voigt & Pakkenburg, 1983), there is growing evidence that sex differences in brain structure exist (Jahanshad & Thompson, 2017). Several of these differences have been identified in brain regions indicated in ASD (for review see (Ruigrok et al., 2014)) suggesting diversity in brain structure associated with sex may modulate ASD risk. However, to date the role of sex differences in ASD liability has been largely understudied. Thus,
while there is some evidence that suggests differences between the sexes in both ASD clinical and biological phenotypes (Lai et al., 2017, Werling 2016), much work remains in understanding the role of sex related risk factors in ASD.

### 1.3 The Etiology of Autism Spectrum Disorder

Postmortem evidence indicates that ASD pathology is likely to first manifest during the early prenatal stages of development (Bauman and Kemper, 2005). For example, reduced numbers of cerebellar Purkinje cells have been observed in both childhood and adult ASD cases (Bailey et al., 1998, Duong et al., 1986, Arin et al., 1991). During fetal development axons climbing from the inferior olive region of the brainstem synapse with Purkinje cell dendrites within a transitory zone beneath the Purkinje cells known as the lamina dessicans (Rakic, 1971). Within human fetuses the lamina dessicans is no longer present after 28-30 weeks gestation (Rakic and Sidman, 1970). After this time, due to the strong connective relationship between Purkinje cells and olivary neurons, loss or damage to Purkinje cells causes retrograde losses of olivary neurons (Norman, 1940, Holmes and Stewart, 1908, Greenfield and Aring, 1954). Thus, findings of reduced Purkinje cells with persevered olivary neurons within ASD cases indicate that the development of ASD pathology occurs at some time prior to the resolution of the lamina dessicans at 28-30 weeks gestation.

Consensus thinking recognizes the role of complex interactions between several possible genetic, epigenetic, and environmental risk factors in the development and manifestation of an ASD phenotype (Sandin et al., 2014). It would be beyond the scope of this thesis to discuss all of these factors in detail. Here I will briefly highlight a selection of some of the factors known to be associated with increased risk for the development of ASD followed by a discussion of current etiological theory.

#### 1.3.1 Environmental Factors

Various environmental factors have been implicated in ASD. Of these the most widely discussed is the idea that vaccinations (particularly the measles, mumps, and rubella [MMR] vaccination) cause ASD. Despite the large amount of attention paid to this potential relationship there is no empirical evidence that strongly supports this claim. In fact a large body of evidence shows no relationship between MMR

Some environmental factors that have been associated with increased risk for ASD include; parental age (Gardener et al., 2009), low birth weight (Schendel and Bhasin, 2008), multiple births (Croen et al., 2002), maternal infections and illness during pregnancy (Sandin et al., 2014, Lichtenstein et al., 2010, Hallmayer et al., 2011a, Atladóttir et al., 2010, Lee et al., 2015) and exposure to toxins (Adams et al., 2009). However, each of these factors taken in isolation accounts for only a small minority of total ASD cases.

1.3.2 Genetic Factors

The role of genetics in ASD susceptibility has been highlighted by studies showing an increased recurrence risk of up to 20% in families with a child with ASD (Ozonoff et al., 2011). Early twin studies showed ASD to be a highly heritable condition with concordance rates of 82-92% amongst monozygotic twins compared to 1-10% in dizygotic twins (Bailey et al., 1995, Folstein and Rutter, 1977), although a more recent study revealed higher concordance rates (>20%) among dizygotic twins (Hallmayer et al., 2011b).

While there is a consensus that genetic factors contribute to ASD, extensive heterogeneity exists in the specific genetic pathways leading to the development of the condition. For example, in ~10% of ASD cases specific genetic conditions (e.g.
fragile X syndrome, Rett syndrome, 22q-11 deletion disorder etc.) can be identified (Tammimies et al., 2015, Carter and Scherer, 2013). These cases where ASD is a secondary diagnosis to a known genetic condition are referred to as ‘syndromic’ autism (as opposed to ‘idiopathic’ autism). Within ~10-20% of the ASD population chromosomal rearrangements as well as rare and de novo copy-number variants can be observed compared to 1-2% in the general population (Jacquemont et al., 2006, Sebat et al., 2007). Additionally significant increases in rare and de novo coding sequence mutations affecting neuronal genes have been observed in ASD and could account for approximately 5-10% of cases (Huguet et al., 2013). Several risk genes for ASD have only recently been identified (C Yuen et al., 2017) and add to the over 850 genes currently implicated in ASD (sfari.org/resources/sfari-gene). These genes largely focus around processes that effect neurogenesis and neuronal and synaptic homeostasis such as chromatin remodeling, metabolism, mRNA translation, and synaptic functioning (Huguet et al., 2013, Packer, 2016). However, much like known environment risk factors, each of these specific risk genes in isolation is more than likely to confer a small amount of risk for ASD and therefore account for a small proportion of cases within the total population.

1.3.3 Competing Aetiological Theories

1.3.3.1 The “Autisms”

The complex heterogeneity of ASD phenotypes and genotypes has led to competing aetiological theories for the condition. On one hand, it has been proposed that ASD is actually comprised of several disorders that are individually rare each having its own aetiology. This concept of ASD as the “Autisms” helps explain the heterogeneity of the condition in terms of both behavioral manifestation and apparent underlying casual mechanisms (Geschwind and Levitt, 2007).

1.3.3.2 Unitary Theory

Another contrasting view is that ASD is a unitary disorder resulting from a common cause. Support for this theory is hindered due to a failure to identify a common causal mechanism for ASD, the observed heterogeneity of the condition, and the ever-increasing list of environmental and genetic risk factors (Boucher, 2011). However, recently it has been proposed that ASD may arise from the single pathophysiological
mechanism, namely heterochronic division of germinal cells during early
development leading to subsequent neural migration abnormalities (Casanova, 2014,
Packer, 2016). Under this unitary theory differences in individual ASD phenotypes can be explained by the severity and nature of the initial triggering exogenous factor, differences in the neurodevelopmental period which the insult was experienced, and the genetic susceptibility of the patient (Casanova, 2007).

1.3.3.3 Atypical Brain Connectivity

A widely accepted hypothesis that accommodates both the “Autisms” and unitary aetiological theories attributes ASD behaviors to aberrant brain connectivity (Belmonte et al., 2004). Under this view atypical neural connectivity is described in two key facets, namely local over connectivity (i.e. within brain regions) and long-range under connectivity (i.e. between brain regions). Behaviorally, aspects of the ASD phenotype such as hyper-arousal, reduced selectivity, and increased computational noise have been linked to possible local over connectivity (Belmonte and Yurgelun-Todd, 2003). Respectively, under connectivity between distant cortical regions may explain deficits in information integration observed in the condition (i.e. weak “central coherence”) (Geschwind and Levitt, 2007, Frith, 2004, Just et al., 2012).

Evidence for atypical connectivity in ASD has been provided by both postmortem and in vivo studies. For example, reports of increased neuronal dendritic spine counts in ASD (Tang et al., 2014, Hutsler and Zhang, 2010) are suggestive of excessive local connectivity in the condition. Structural imaging results also show reduced intrinsic wiring costs within cortical grey matter in ASD (Ecker et al., 2013b) that may represent a bias towards local over global information processing (Happé and Frith, 2006).

Poor connectivity between brain regions in ASD is supported by functional magnetic resonance imaging (fMRI) findings of lower degrees of synchronization across cortical networks responsible for language processing and executive functioning (Just et al., 2007, Just et al., 2004). Other studies of under connectivity have focused on white matter deficits, particularly in the corpus callosum due to its role in providing
long range connections between the brain’s two hemispheres and potential mediation of inter-regional cortical synchronization (Uhlhaas et al., 2009). Within ASD, diffusion-imaging measures of lower fractional anisotropy and increased radial diffusivity in the corpus callosum have been reported (Alexander et al., 2007, Barnea-Goraly et al., 2004, Shukla et al., 2010). Studies using magnetic transfer imaging have also reported atypical myelin development in the structure (Gozzi et al., 2012). These results likely reflect underlying changes in myelination, density, and size of axons within the tract (Hong et al., 2011) which are likely to adversely affect long range connectivity in the ASD brain. Additional ways that atypical cortical structure observed in ASD might affect brain connectivity are discussed in section 1.4.2.

1.4 The Structure of the Brain in Autism Spectrum Disorder

Indications of abnormalities in the neuroanatomical structure of the autistic brain arose from some of the first documented observations of the condition, where Kanner (1943) noted ASD patients to have significantly larger head size than typically developing (TD) individuals. Since these early observations several different research techniques have been applied in order to better understand atypical neuroanatomical structure in ASD (Ecker and Murphy, 2014). These studies have led to the identification of several brain regions indicated in ASD behaviors (Figure 1.3).
This thesis focuses on applications of structural magnetic resonance imaging (MRI) methods, namely surface based morphometry (SBM) in order to (i) identify novel ways to measure atypical brain structure in ASD, (ii) test the efficacy of measures of brain structure to identify ASD within clinical samples, and (iii) examining how normative variance in SBM measures associated with biological sex are associated with risk for the condition. Accordingly, the focus of this section will be a review of in vivo structural neuroimaging studies of the brain in ASD with a focus on SBM studies cortical structure.
1.4.1 Volumetric Differences in the Autistic Brain

Consistent with early observations (Kanner, 1943), findings of increased global brain size in ASD, particularly among younger patients, remains one of the most widely reported and cited neuroanatomical findings in the condition. Some of the first MRI studies in ASD showed patients to have significantly larger mid-sagittal brain areas apposed to healthy controls (Piven et al., 1992, Piven et al., 1996). Furthermore, it was revealed that these increases were not gross in nature, but rather differed between the frontal, temporal, parietal, and occipital lobes (Piven et al., 1996, Piven et al., 1995). In line with these MRI studies, findings using measures of head circumference have shown ASD patients to have higher rates of macrocephaly than the general population (Davidovitch et al., 1996, Woodhouse et al., 1996, Fidler et al., 2000, Miles et al., 2000, Dementieva et al., 2005).

Current evidence shows the ASD brain to have a nonlinear growth trajectory that when compared to TD controls experiences periods of precocious growth during the first few years of life, which are then followed by a deceleration leading to harder to distinguish differences in adolescence and later adulthood (Courchesne et al., 2011a). These early increases in brain volumes among young children with ASD appear to have an anterior to posterior growth gradient (Courchesne et al., 2004) showing the greatest GM differences in the frontal and temporal lobes (Carper et al., 2002). There is also some preliminary evidence to suggest that the observed enlargements in brain volumes in young children with ASD are driven by disproportional increases in WM compared to GM (Courchesne et al., 2001, Hazlett et al., 2005, Ben Bashat et al., 2007), yet it remains unclear if these WM increases persist into later life (Palmen et al., 2005, Hazlett et al., 2006, Barnea-Goraly et al., 2004). Furthermore, in line with the heterogeneous nature of the condition, recent evidence suggests that brain overgrowth in ASD may be specific to a subset of the ASD population (Libero et al., 2016).

Several MRI studies have used voxel based morphometry (VBM) methods (Ashburner and Friston, 2000) to explore regional differences in brain volumes (Nickl-Jockschat et al., 2012). While these studies are largely consistent in reporting distributed patterns of volumetric differences in ASD (Ecker et al., 2012, Hyde et al.,
they often differ on specific brain regions identified as well as the directionality of their findings. Some limitations common to all imaging studies of ASD contribute to the lack of replicability of VBM findings. These include i) heterogeneity of ASD phenotypes and genotypes ii) non linear changes in brain structure across developmental stages (Ecker et al., 2014, Courchesne et al., 2011a), and iii) small samples often drawn from homogenous subgroups of the autism spectrum (e.g. high functioning adult males, ect.). Despite inconsistencies, VBM studies have been valuable in linking regional atypical brain volumes to ASD symptomatology (Ecker et al., 2012) and highlighting anatomical heterogeneity across the spectrum (McAlonan et al., 2009, McAlonan et al., 2008).

1.4.2 Cortical Morphometry of the Autistic Brain

The pathological processes underlying volumetric abnormalities in ASD remain unclear. More recent studies of cortical structure have moved beyond measures of volume to investigations of the dimensions and shape (i.e. morphometry) of the cortex. Surface based morphometry (SBM) comprises a series of structural MRI methods that use tessellated reconstructions of the cortical surface to quantify a variety of morphological features relating to brain structure. These studies are particularly valuable in ASD for their potential to indicate developmental pathways in the condition that are associated with specific features of neural architecture, particularly in cortical grey matter (i.e. the cerebral cortex).

The cerebral cortex is organized according to different cellular layers and follows a columnar architecture. At their core, cortical minicolumns are comprised of radially oriented arrays of pyramidal projection neurons. Inputs and outputs from these pyramidal neurons are modulated by GABAergic interneurons that lie within the core and periphery of mini-columns allowing for area and task specific information processing (Kawaguchi and Kubota, 1997, Sanders et al., 2012, DeFelipe, 1997). Several types of progenitor cells and related molecular and genetic signaling pathways regulate the development of this cortical organization (Figure 1.4).
Figure 1.4, Development of the Human Cerebral Cortex (Fernández et al., 2016): This model depicts the main types of progenitor cells involved in the development of the cerebral cortex. Arrows indicate lineage relationships between cell types. During cortical expansion most neuroepithelial cells divide symmetrically to amplify their numbers and generate apical radial glial cells (aRGC). During neurogenesis these aRGCs divide into neuron cells directly or via intermediate progenitor cells (IPC). IPCs divide solely into neurons, allowing for rapid proliferation of cortical neurons. Neurons migrate along scaffolding provided by aRGC fibers to form cortical units known as minicolumns. Basal radial glial cells (bRGC) provide additional scaffolding for migrating neurons across convoluted cortical surfaces and are thought to be important to the development of cortical gyration. Boxes highlight some of the many molecular and genetic pathways that regulate these processes. Marginal zone (MZ), cortical plate (CP), intermediate zone (IZ), subventricular zone (VZ), ventricular zone (VZ).

According to the radial unit hypothesis (Rakic, 1995) the quantity of neuron cells within each individual minicolumn is a main determinant of cortical thickness (CT) while cortical surface area (SA) is closely related to the number of minicolumns (i.e. radial units). It is now widely believed that intermediate progenitor cells (IPC) are a principle modulator of CT (Pontious et al., 2007). These cells divide solely into neurons (Miyata et al., 2004, Noctor et al., 2004) and are responsible for the vast proliferation of neuronal cells within the cortical sheet. These neuronal cells migrate from the ventricular zone along scaffolding fibers provided by apical radial glial cells (aRGC) leading to the formation of cortical minicolumns. Increases in SA have thus been linked to the length of symmetrical division cycles for neuroepithelial progenitor cells responsible for the expansion of aRGC (Pontious et al., 2007).
There is a large body of postmortem evidence that suggests disruptions to the above developmental processes in ASD. For example, individuals with ASD have been shown to have increased minicolumn density in frontal and temporal regions (Casanova et al., 2006, Casanova et al., 2002). Shorter distances between these cellular units could introduce hyper connectivity within local brain regions resulting in signal delays and metabolic inefficiencies in connecting disparate brain regions, potentially leading to ASD behaviors (Belmonte et al., 2004, Horwitz et al., 1988, Just et al., 2004, Casanova et al., 2003). As could be predicted by this increased column density, greater numbers of prefrontal neuron cells in young children with ASD have also been reported (Courchesne et al., 2011b). The presence of supernumerary neurons within white matter below the cortical sheet and poor definition of the grey-white matter boundary in ASD (Avino and Hutsler, 2010, Bailey et al., 1998) also point towards disruptions to neuronal migratory processes in the condition. Together these findings provide evidence for atypical cell division of neuronal progenitors and the subsequent migration of neuron cells to their target locations within the developing cortex in ASD (Casanova, 2014). Furthermore, it can be predicted that disruptions to these foundational developmental processes have lasting impact on the structural morphology of the cortex.

Several researchers have applied SBM methods to quantify differences in cortical features associated with diagnoses of ASD. To date CT is the most reported in the condition, although results have been mixed. Some studies have reported decreased CT in ASD (Hadjikhani et al., 2006, Scheel et al., 2011, Ekman et al., 1975), others no differences (Hazlett et al., 2011, Hutsler et al., 2007), and others still CT increases in the condition (Hardan et al., 2006). These inconsistent findings can likely be largely attributed to nonlinear changes in CT across developmental stages (Ecker et al., 2014). For example, within a large multi-centre study including individuals from 6-35 years of age, Khundrakpam et al. (2017) reported CT increases in young children with ASD were greatest around the age of ten. Due to accelerated cortical thinning these differences decreased into adolescence and adulthood such that little to no difference was observed at 35 years (Khundrakpam, 2017). These findings are supported by both histological (Hutsler et al., 2007) and longitudinal MRI findings (Hardan et al., 2009) reporting accelerated reductions in CT with age in ASD.
compared to TD controls, which is consistent with the observed developmental trajectory of volume measures within the condition (Courchesne et al., 2011a).

However, imaging studies of both CT and SA have found closer links between SA expansion and observed early life volumetric increases in ASD (Hazlett et al., 2011, Hazlett et al., 2017). One recent study reported that accelerated SA expansion between 6 and 12 months predicted increased brain volumes at 24 months (Hazlett et al., 2017). Furthermore, Ecker et al. (2013a) found that 56% of a distributed pattern of volumetric differences in adults with ASD was driven by underlying changes in SA measures compared to 8% for CT, with only 5% shared between the two. The small degree of overlap between CT and SA findings highlights the need for studies that differentiate volumetric measures into these component measures. Furthermore, findings by Ecker et al. (2013a) and Hazlett et al. (2017) of a greater contribution of SA expansion to volumetric differences advance theories that processes governing SA expansion are a possible aetiological agent in ASD. However, it should be noted that Ecker et al. (2013a) was only able to highlight these underlying relationships through liberal statistical thresholding (uncorrected \( p < 0.05 \)). When corrected for multiple comparisons no significant differences in SA measures were observed, which is consistent with a few other reports (Wallace et al., 2013, Raznahan et al., 2009). Furthermore, another study has reported reduced cortical SA within the left fusiform and bilateral middle temporal cortices in ASD (Libero et al., 2014), which were also reported by Ecker et al. (2013a).

In addition to SA and CT, individuals with ASD have been shown to have atypical morphology associated with cortical folding. For example, an early MRI study conducted by Piven et al. (1990) found significantly more instances of cortical malformations relating to folding among ASD cases including; i) polymicrogyria (excessive small folding), ii) schizencephaly (clefts lined within the cortical grey matter) and iii) macrogyria (increased gyri size) (Piven et al., 1990). Other early MRI studies have reported atypical placement of major sulci within the frontal and temporal lobes (Levitt et al., 2003) and increased gyrification in frontal brain regions (Hardan et al., 2004) in children and adolescences with ASD.
More recent studies using MRI based gyrification index (GI) measures (Zilles et al., 1988, Su et al., 2013) to quantify cortical folding have reported decreased GI in ASD (Bos et al., 2015). Recently SBM studies using a vertex wise implementation of the GI, namely the local gyrification index (lGI) (Schaer et al., 2008), have reported decreased gyrification in inferior fronto-parietal and posterior medial cortical regions (Schaer et al., 2013) as well as the supramarginal gyrus (Libero et al., 2014). Others have reported regional patterns of increased lGI in ASD (Wallace et al., 2013, Ecker et al., 2016). Some studies have linked differences in gyrification in ASD to underlying changes in WM diffusion measures (Ecker et al., 2016, Bos et al., 2015, Schaer et al., 2013) which provides preliminary evidence for what are likely to be complex interrelations between white matter and grey matter development which may contribute to atypical intra- and inter-cortical connectivity in the condition.

In addition to gyrification other studies have sought to quantify various aspects of sulcal morphometry and the relationship between these features and ASD symptomology. For example, increased sulcal depths within ASD participants have been reported and found to correlate with ADOS social interaction scores (Auzias et al., 2014, Nordahl et al., 2007). Furthermore, abnormalities in sulcal length and SA have been noted in ASD (Shokouhi et al., 2012, Nordahl et al., 2007). Some of these abnormal findings appear to be larger within younger children on the spectrum suggesting a “normalization” of these measures into adolescence and adulthood similar to the trajectory observed in volumetric measures (Nordahl et al., 2007).

The mechanisms underlying and relating to cortical folding are numerous and complex (Fernández et al., 2016). Folding has been related to differential growth rates between outer and inner cortical layers (Richman et al., 1975, Kriegstein et al., 2006), proliferation of bRGCs (Nonaka - Kinoshita et al., 2013, Reillo et al., 2011), axonal tension (Van Essen, 1997), and the neural complexity of a particular brain region (i.e. numbers of neurons, synaptic density, and/or dendritic arborization) (Welker, 1990). Studies of cortical folding are further complicated by large variances in the degree of gyrification across individuals in the normative population (Bartley et al., 1997). This is supported by findings of discordant folding patterns between monozygotic twins pairs where only one twin had a diagnosis of ASD (Kates et al., 2009). This is in
contrast to volumetric measures of the brain, which have been found to be highly concordant between twins (White et al., 2002). These twin studies and the lack of correlation between cortical folding and measures that are largely genetically determined such as total brain volume and body weight and length (Zilles et al., 1988, Rogers et al., 2010) indicate that measures of gyrification may be more sensitive to environmental factors than measures that are largely genetically determined (Zilles et al., 2013).

The identification of group differences in SBM measures such as CT, SA, sulcal depth and GI are valuable for their ability to highlight different developmental pathways that may contribute to ASD pathology. In doing so these studies can narrow focus on or introduce novel etiological targets for the condition. For example, IPCs and the proliferation of neuron cells (Pontious et al., 2007), dendritic arborization (Huttenlocher, 1990), and myelination (Sowell et al., 2004) have all been attributed to changes in CT while the length of symmetrical division cycles for aRGC progenitor cells appear to correspond with proliferation of cortical minicolumns and SA (Pontious et al., 2007). Parsing out the deterministic mechanisms responsible for cortical folding abnormalities in ASD may be more difficult due to the potentially large contribution of environmental factors in the development of folding patterns (Zilles et al., 2013). However, current evidence suggests that cortical folding relates to bRGCs and differential expansion rates of the outer and inner cortical surface (Richman et al., 1975, Kriegstein et al., 2006, Nonaka - Kinoshita et al., 2013, Reillo et al., 2011) thus, cortical folding may be related to processes governing the expansion of cortical SA.

To date, results reported by SBM studies in ASD have been somewhat inconsistent. In many cases this can be attributed to the complex developmental trajectories that morphometric cortical features undergo (Ecker et al., 2014). This is further complicated by relatively small samples sizes and the large degree of phenotypic heterogeneity among the ASD population. Several of these studies are likely underpowered due to such small samples as one power study estimated that 50 subjects per group are needed to detect a 0.25mm differences in CT between two groups (Pardoe et al., 2013). To date, most SBM studies fall below this number of
participants. Thus, work remains to be done in confirming differences in morphological phenotypes across the spectrum as well as untangling the numerous genetic and molecular pathways that regulate their development. Additionally, some atypical structural features that have been identified in postmortem studies of ASD brains, such as integrity of the grey-white matter boundary, have yet to be investigated *in vivo*.

### 1.5 Beyond Group Differences: Pattern Recognition in Autism Spectrum Disorder

Until recently, the majority of neuroimaging research in psychiatric conditions has employed mass-univariate statistical methods based the general linear model (GLM) (Friston et al., 1994). The GLM is a valuable tool for identifying mean differences between groups. However, these models are not suited to make predictions for individual cases based on multivariate patterns within the data (i.e. predictions regarding group membership or clinical outcome), and thus offer limited translational value to clinical applications. Alternatively, a class of ‘machine learning’ methods known as multivariate pattern classification (MVPC) can be applied to generate image-based predictive models that have the potential to be utilized as diagnostic tools and intervention outcome measures.

Within ‘machine learning’ an algorithm is said to learn from experience $E$ with respect to some class of tasks $T$ and performance measure $P$. Learning is said to occur when performance in task $T$ improves with experience $E$ as measured by performance $P$ (Mitchell, 1997). Within this context classification poses a supervised learning problem in which examples from a training set of empirical data are mapped to a known output such as a class label. In brief MVPC algorithms differentiate cases between two or more classes in the following way. During a ‘training phase’ the model is initially trained using well-defined samples or data subsets (e.g. ASD and controls) to establish a mathematical criterion or decision function that best distinguishes between the groups. During the ‘testing phase’ the decision function is applied to predict the label of an unseen data point. Often MVPC model performance is validated within the same sample using cross-validation methods. However, gold
standard assessment of a predictive model’s accuracy in distinguishing between classes is reliant on validation using completely independent testing data.

A variety of MVPC algorithms including the Support Vector Machine (SVM) (Vapnik, 1995) and Gaussian Process Classification (GPC) (Rasmussen and Williams, 2006) as well as numerous others (e.g. linear discriminant classifiers, logistic regression, decision trees, random forest, etc.) have been successfully trained using various structural and functional imaging data modalities to distinguish individuals diagnosed with various neuropsychiatric disorders from healthy controls and/or to predict treatment outcome or subjective states (Wolfers et al., 2015, Arbabshirani et al., 2016). These findings provide proof of concept for the development of safe, in vivo diagnostic and prognostic tools and biomarkers for a variety of psychiatric conditions. However, until these models undergo stringent validation on independent data sets that accurately reflect the clinical environment their potential translational value remains to be realized.

1.5.1 Multivariate Pattern Classification in Autism Spectrum Disorder

In a behaviorally defined disorder such as ASD the development of biologically based diagnostic and prognostic markers would represent an invaluable tool to clinicians. Furthermore, identification of a biomarker for ASD could provide pathways towards better understanding the casual mechanisms for the condition. While several genetic and neurobiological markers have been proposed, there are currently no reliable biomarkers that have been validated for clinical use in ASD (Ruggeri et al., 2013, Ecker and Murphy, 2014). Classification of ASD using MRI features is one pathway towards the development of a biomarker for the condition. To date several MVPC methods have been applied to a variety of neuroimaging data and psychiatric conditions (Wolfers et al., 2015, Arbabshirani et al., 2016). The results of MVPC studies in ASD are reviewed bellow and summarized in Table 1.1.
<table>
<thead>
<tr>
<th>Modality</th>
<th>Features</th>
<th>Classifier</th>
<th>Number of Subjects</th>
<th>Biological Gender</th>
<th>ASD age</th>
<th>Overall Accuracy</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>sMRI</td>
<td>Cortical Thickness and Volumetric ROIs</td>
<td>SVM</td>
<td>TDC=325, ASD=325</td>
<td>Male and Female</td>
<td>17.8±7.4</td>
<td>59%</td>
<td>Sabuncu et al. (2014)</td>
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<tr>
<td>sMRI</td>
<td>Cortical Thickness and Volumetric ROIs</td>
<td>Multi-kernel SVM</td>
<td>TDC=59, ASD=58</td>
<td>Male and Female</td>
<td>10.8±4.0</td>
<td>96.3%</td>
<td>Wee et al. (2014)</td>
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<tr>
<td>sMRI</td>
<td>VBM GM and WM volume maps</td>
<td>SVM</td>
<td>TDC=24, ASD=24</td>
<td>Male and Female</td>
<td>13.2±0.6</td>
<td>92%</td>
<td>Uddín et al. (2011)</td>
</tr>
<tr>
<td>sMRI</td>
<td>Cortical Thickness</td>
<td>Logistic model trees</td>
<td>TDC=16, ASD=22</td>
<td>Male and Female</td>
<td>9.2±2.1</td>
<td>87%</td>
<td>Jiao et al (2010)</td>
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<tr>
<td>sMRI</td>
<td>VBM and SBM based ROI</td>
<td>SVM</td>
<td>TDC=20, ASD=21</td>
<td>Male and Female</td>
<td>4.1±0.8</td>
<td>74%</td>
<td>Gori et al. (2015)</td>
</tr>
<tr>
<td>sMRI</td>
<td>GM volume map</td>
<td>SVM</td>
<td>TDC=40, ASD=52</td>
<td>Male and Female</td>
<td>14.4±1.7</td>
<td>80-85%</td>
<td>Segovia et al. (2014)</td>
</tr>
<tr>
<td>sMRI</td>
<td>VBM GM and WM volume maps</td>
<td>SVM</td>
<td>TDC=22, ASD=22</td>
<td>Male</td>
<td>27.0±7.0</td>
<td>77%</td>
<td>Ecker et al.(2010)</td>
</tr>
<tr>
<td>sMRI</td>
<td>Cortical Thickness, Surface Area, Sulcal depth, Curvature, Metric Distortion</td>
<td>SVM</td>
<td>TDC=20, ASD=20</td>
<td>Male</td>
<td>33.0±11.0</td>
<td>85%</td>
<td>Ecker et al.(2010)</td>
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<tr>
<td>sMRI</td>
<td>VBM GM volume maps</td>
<td>SVM</td>
<td>TDC=38, ASD=30</td>
<td>Female</td>
<td>4.4±1.5</td>
<td>80%</td>
<td>Calderoni et al. (2012)</td>
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<td>sMRI</td>
<td>Volumetric measures</td>
<td>Discriminant function analysis</td>
<td>TDC=15, ASD=52</td>
<td>Male</td>
<td>3.4±0.7</td>
<td>95.8%</td>
<td>Akshoomoff et al. (2004)</td>
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<td>sMRI</td>
<td>Age, Sex, Volume, Surface Area, and Cortical Thickness ROIs</td>
<td>Three-Stage Deep Learning Network</td>
<td>HRN = 145, ASD=34</td>
<td>Male and Female</td>
<td>6 and 12 months</td>
<td>94%</td>
<td>Hazlett et al. (2017)</td>
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<td>sMRI</td>
<td>Extra-axial Cerebrospinal Fluid</td>
<td>Decision Tree</td>
<td>HRN = 174, ASD=47</td>
<td>Male and Female</td>
<td>6 months</td>
<td>66-72%</td>
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<td>dMRI</td>
<td>FA and MD based ROI</td>
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<td>TDC=30, ASD=45</td>
<td>Male and Female</td>
<td>10.5±2.5</td>
<td>80%</td>
<td>Ingalhalikar er al. (2011)</td>
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<td>fMRI</td>
<td>Activation of selected voxels during social interaction task</td>
<td>Gaussian naïve Bayes</td>
<td>TDC=17, ASD=17</td>
<td>Male and Female</td>
<td>25.6±6.7</td>
<td>97%</td>
<td>Just et al. (2014)</td>
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<td>fMRI</td>
<td>Language and theory of mind task based functional connectivity</td>
<td>Logistic regression</td>
<td>TDC=14, ASD=13</td>
<td>Male</td>
<td>21.4±3.9</td>
<td>96%</td>
<td>Murdaugh et al. (2012)</td>
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<tr>
<td>fMRI</td>
<td>Activation of selected voxels during social interaction task</td>
<td>SVM</td>
<td>TDC=14, ASD=15</td>
<td>Male and Female</td>
<td>28.6±1.8</td>
<td>92.3%</td>
<td>Chanel et al. (2016)</td>
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<td>rsfMRI</td>
<td>rsfMRI ICA components</td>
<td>Logistic regression</td>
<td>TDC=20, ASD=20</td>
<td>Male and Female</td>
<td>9.9±1.5</td>
<td>78%</td>
<td>Uddín et al (2013)</td>
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Table 1: Multivariate Pattern Classification Studies of Autism Spectrum Disorder (continued)

<table>
<thead>
<tr>
<th>Modality</th>
<th>Features</th>
<th>Classifier</th>
<th>Number of Subjects</th>
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<th>ASD age ± standard deviation</th>
<th>Overall Accuracy</th>
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<td>ROI based functional connectivity</td>
<td>Random forest</td>
<td>TDC=126, ASD=126</td>
<td>Male and Female</td>
<td>14.8±1.6</td>
<td>79%</td>
<td>Chen et al. (2015)</td>
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<tr>
<td>rsfMRI</td>
<td>ROI based functional connectivity</td>
<td>Logistic regression</td>
<td>TDC=59, ASD=59</td>
<td>Male and Female</td>
<td>17.66±2.7</td>
<td>77%</td>
<td>Plitt et al (2015)</td>
</tr>
<tr>
<td>rsfMRI</td>
<td>ROI based functional connectivity</td>
<td>Probabilistic neural network</td>
<td>TDC=328, ASD=312</td>
<td>Male and Female</td>
<td>13.2±3.1</td>
<td>90%</td>
<td>Iidaka (2015)</td>
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<tr>
<td>rsfMRI</td>
<td>ROI based functional connectivity</td>
<td>Thresholding</td>
<td>TDC=40, ASD=40</td>
<td>Male</td>
<td>22.7±7.4</td>
<td>79%</td>
<td>Anderson et al. (2011)</td>
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<td>rsfMRI and fMRI</td>
<td>Functional connectivity and FA values</td>
<td>SVM</td>
<td>TDC=15, ASD=15</td>
<td>not reported</td>
<td>21.1±0.9</td>
<td>95.9%</td>
<td>Deshpande et al. (2013)</td>
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<td>rsfMRI and dMRI</td>
<td>Cortical Thickness, FA and neurochemical concentration</td>
<td>Decision tree</td>
<td>TDC=18, ASD=19</td>
<td>Male and Female</td>
<td>27.1±1.3</td>
<td>91.9%</td>
<td>Libero et al. (2015)</td>
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<tr>
<td>rsfMRI and sMRI</td>
<td>rsfMRI, Cortical Thickness and Volume based graph theory metrics</td>
<td>Random forest</td>
<td>TDC=153, ASD=127</td>
<td>Male and Female</td>
<td>13.5±6.0</td>
<td>70%</td>
<td>Zhou et al. (2014)</td>
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Table 1, Multivariate Pattern Classification Studies of Autism Spectrum Disorder: Twenty-two MVPC studies that use magnetic resonance imaging (MRI) data to classify ASD are summarized. Ages are given in mean years ± standard deviation. sMRI = structural MRI, VBM = voxel based morphometry, SBM = surface based morphometry dMRI = diffusion MRI, FA = fractional anisotropy, MD = mean diffusivity, fMRI = functional MRI, rsfMRI = resting state functional MRI, MRS = magnetic resonance spectroscopy, SVM = support vector machine, TDC = typically developing controls.
The use of MVPC models based on brain imaging data in ASD was first reported by Akshoomoff et al. (2004). Using a discriminate function and pre-selected volumetric regions of interest (ROIs) from young children and TD controls a high classification accuracy (95%) was achieved. However, this rate was not based on a thorough cross validation but on reclassification of the entire sample and is therefore inherently circular. More recent studies employing SVM algorithms to classify ASD cases using volumetric GM and WM ROIs have reported overall classification accuracies ranging from 65-90% for different brain regions (Uddin et al., 2011, Segovia et al., 2014). In addition to these volumetric ROI studies whole brain VBM data has been shown to classify adult male ASD cases with a cross-validated accuracy of 81% (Ecker et al., 2010b).

When comparing structural MRI measures used to predict ASD diagnoses, models using SBM features such as CT have been generally shown to outperform those using volumetric features (Jiao et al., 2010). A study by Ecker et al. (2010a) investigating the predictive value of five SBM measures (CT, SA, sulcal depth, mean radial curvature, and metric distortion) used a SVM to classify adult males with ASD and matched controls. These studies revealed that CT measures in the left hemisphere produced the highest cross-validated prediction accuracy (90%). Other SVM studies using a variety of regional cortical volume and thickness features have also successfully classified ASD cases from controls albeit at lower discriminative accuracies (59-74%) (Gori et al., 2015, Sabuncu and Konukoglu, 2014). In addition to being able to classify cases from controls, measures of interregional CT covariance have also been shown to be predictive of current symptom measures among ASD patients (Sato et al., 2013).

In addition to volumetric and morphometric features both diffusion and functional MRI (fMRI) features have been used to classify ASD. To date, two studies investigating the predictive power of mean diffusivity and fractional anisotropy diffusion measures in ASD using a SVM have reported classification accuracies in line with the best performing SBM based models (~80%-89%) (Ingahalikar et al., 2010, Ingahalikar et al., 2011). Activation patterns during theory-of-mind and language fMRI tasks as well as resting state brain activity (rsMRI) have been used to classify individuals with ASD from controls with varying reported accuracies.
Recent studies have begun to implement methodologies that combine multiple neuroimaging modalities into single predictive models. Decision tree algorithms have been used to combine selections of diffusion, morphometric, and MR spectroscopy features as inputs for ASD classification. Under this framework one model was able to achieve a 91.9% classification accuracy between ASD cases and controls. These findings also highlighted the potential utility for MVPC methods to aid in understanding ASD risk factors and etiology. For example, if certain favorable values among features within the decision tree were present an individual could be protected from an ASD classification, even in the presence of other unfavorable metrics (Libero et al., 2015). Other studies have combined diffusion MRI with auditory task based magnetoencephalography (MEG) data, resulting in the successful classification of ASD cases from TD controls at a CV accuracy of 83.3% (Ingalhalikar et al., 2014). Another recent study employed multi-kernel learning SVM methods to investigate the predictive capacity of combined vertex based surface measures (i.e. CT, grey and white matter surface volumes), subcortical volumetric ROI data, and interregional correlation measures. Using this methodology the researchers reported a CV accuracy over 96% (Wee et al., 2014).

Two recent studies have shown promise in predicting ASD diagnoses at 24 months based on MRI measures at 6 month, an age before diagnosis is possible using current gold standard criterion. The first of these studies used a three stage deep learning algorithm employing age and sex data along with volumetric, CT, and SA based ROIs. This study was able to classify high-risk infants who received an ASD diagnosis at 24 months from those who did not at accuracies of 95%. The second study utilized measures of extra-axial spinal fluid at 6 months to classify high-risk infants with ASD at 24 months (Shen et al., 2017). This study achieved a cross-validated classification accuracy of 69% within the training sample, as well as an accuracy of 72% when applied to an independent data set.
1.5.2 Towards Translational Tools

While the above studies provide proof of concept for imaging based diagnostic models in ASD, translating these tools from the research to ‘real-world’ clinical settings raises additional hurdles. To date no studies have tested classification models for ASD trained in the research environment on independent samples that are representative of the broader clinical population. This is important because several differences exist between research and clinical ASD cohorts. For example, individuals with ASD recruited within the research environment will generally be excluded if they have comorbid conditions, are taking certain medications, or fail to reach cut-offs on behavioral metrics such as the autism diagnostic interview (ADI) and/or autism diagnostic observation schedule (ADOS). As such, many individuals who would qualify for a clinical ASD diagnosis would not meet inclusion criteria for research. Thus, inherent differences in the clinical phenotype between individuals in research and clinical ASD samples are likely. Accordingly, if imaging-based prediction models are to be adopted in the clinical setting, a critical first step will be the testing of models trained in the research setting on separate (i.e. independent) samples that are representative of the clinical population.

1.6 The Current Thesis: Aims and Hypotheses

This chapter has outlined several advances that have been made in structural MRI studies of ASD. Broadly, this thesis aims to advance this literature in a way that offers the potential for real world benefits to patients and stakeholders. With regards to this aim this thesis includes fours studies that focus on i) the development of novel informative imaging features for ASD, ii) evaluating the efficacy of structural neuroimaging measures to predict ASD diagnosis within clinical samples, and iii) investigating the role of normative sex differentiation in brain structure on the probability for ASD.

1.6.1 Study 1: In Vivo Evidence of Reduced Integrity of the Grey-White Matter Boundary in Autism Spectrum Disorder

Reduced integrity of the grey-white matter boundary has previously been reported by postmortem studies in individuals with ASD (Avino and Hutsler, 2010). However, prior to this work no in vivo studies have examined this atypical structure in the condition. Here, we used structural MRI to examine differences in tissue contrast
between grey and white matter at the boundary between these tissue types as well as across different cortical layers. As individuals with ASD are thought to have a less well defined transition between grey and white matter we hypothesized that individuals with ASD would show significant reductions in measures of tissue contrast taken at and around the grey-white matter boundary.

1.6.2 Study 2: Clinical Validation of Pattern Classification Models for Autism Spectrum Disorder

Several proof of concept studies have shown promise in developing diagnostic imaging tools for ASD using multivariate pattern classification (MVPC) and SBM measures (see section 1.4.1 for review). However, to date these models have not been tested on samples that are representative of the clinical setting. Thus, this study first aimed to train SVM and GPC models to separate adult ASD patients from TD controls acquired in the research setting using a variety of morphological cortical features, including features that have not previously been reported (i.e. lGI and tissue contrast). The second and primary aim of this study was to then test the ability of classifiers trained within the research setting to make diagnostic predictions within an independent clinical sample consisting of individuals who had been referred for a specialist ASD diagnostic assessment.

1.6.3 Study 3 and 4: Association Between the Probability of Autism Spectrum Disorder and Normative Sex-Related Phenotypic Diversity in Brain Structure

Male preponderance is an understudied hallmark of ASD and other neurodevelopmental and psychiatric conditions. While some degree of male preponderance is likely attributable to sex differences in the ASD phenotype and male biased diagnostic tools, it has been purposed that variance in biology attributable to normative sex differences may modulate ASD risk (Werling, 2016). Thus, these studies sought to understand how biological sex/gender is associated with probability for ASD. The aims of these studies were three-fold; i) provide a phenotypic characterization of normative sex related diversity in brain structure, ii) determine how diversity in sex based neurophenotypes is associated with probability for ASD,
and iii) estimate how much ASD prevalence in the general population might vary as a function of normative sex-related phenotypic diversity in brain structure
1.7 References


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Chapter 2: Magnetic Resonance Imaging and Analysis Methods

I will begin this chapter by discussing the basic physical principles underlying Magnetic Resonance Imaging (MRI) (Lauterbur, 1973, Mansfield, 1977), as reviewed by Sprawls (1987). Details of basic MRI scanning sequences and those used to acquire structural images of the brain within this thesis will then be covered. Last, I will discuss methods for the analysis of MRI images including surfaced based (SBM) morphometry, the general linear model (GLM), and multivariate pattern classification (MVPC) (i.e. “machine learning”) techniques.

2.1 Principles of Nuclear Magnetic Resonance

Magnetic resonance imaging (MRI) is made possible by interactions of atomic nuclei and radio waves that occur when objects are placed within a magnetic field. Under these conditions atoms take on a resonant characteristic whereby electromagnetic radiation is absorbed and rereleased at a particular frequency. This phenomena is generally known as Nuclear Magnetic Resonance (NMR) (Bloch, 1946, Purcell et al., 1946). The resonance frequency of materials (e.g. animal tissue) in a magnetic field typically occurs within the radio frequency (RF, ~3 kHz to 300GHz) of the electromagnetic spectrum. This allows for NMR RF signal to be captured using a radio-receiver coil and later displayed as images (i.e. MRI).

The magnetic properties and potential NMR RF signal of individual nuclei are reliant on their neutron-proton composition. Nuclei with an odd number of protons and neutrons act in themselves as small magnets with a magnetic dipole. Nuclei with this magnetic property have increased NMR susceptibility. Of all nucleotides, hydrogen-1 ($^1$H) is unique in that it occurs in relatively high concentrations in most human tissues and is highly sensitive to NMR. These properties make $^1$H key to the efficacy of MRI in humans.

In order to produce NMR a magnetic nucleus (e.g. $^1$H) must be placed within a static magnetic field denoted as $B_0$. MRI scanners use superconducting electrical coils to generate highly homogeneous magnetic fields in the range of 1.5 to 7 tesla (T), with 3T magnets being the standard at time of writing for clinical research. During
scanning, a patient lies within this electrical coil (i.e. bore) so that the magnetic field runs parallel to the individual’s body.

During MRI scanning protocols NMR RF signal is captured across segmented three-dimensional volumetric units known as voxels. The RF signal intensity at each individual voxel is determined by characteristics of the tissue, namely 1) the concentration of chemical elements, 2) quantity of magnetic isotopes, and 3) the NMR sensitivity of the specific nuclides within the tissue. Signal from a voxel is emitted as the result of stimulation by short RF energy pulses applied during the imaging cycle. The NMR process involves a series of interactions involving the magnetic nuclei, the magnetic field, and these RF pulses and signals.

When not undergoing strong magnetic forces magnetic nuclei (e.g. $^1$H) will be randomly oriented throughout the body. However, when placed within a magnetic field ($B_0$) these nuclei experience torque to bring their magnetic dipole in alignment with the direction of the field. Due to agitation caused by thermal energy only a small percentage of the total nuclei (on the order of a few out of a million) come into alignment during MRI scanning of the human body. The number of magnetic nuclei that align with the magnetic field is a function of the field strength of the MRI scanner.

Once aligned with $B_0$ nuclei are not static but precess (an oscillation similar to a spinning top wobbling) along the axis of the magnetic field (Figure 2.1 A). The precession of an atomic nucleus in a magnetic field occurs at a specific frequency making it susceptible to RF energy applied at the same frequency. This frequency is known as the resonant frequency, or Larmor frequency ($\omega$),

$$\omega = (\gamma_i)B_0$$

Where $B_0$ = the applied magnetic field strength, and $\gamma$ = the gyromagnetic ratio or resonance quality of the nuclei $i$. When RF energy is applied at the Larmor frequency the absorption of this energy by the nuclei will cause it to flip its alignment away from the direction of $B_0$, resulting in a state of increased energy or excitation (Figure
2.1 B). In the excited state the nuclei experiences increasing force from $B_0$ to align with the main direction of the magnetic field. A nuclei in an excited state returns to alignment with $B_0$ by transferring energy to surrounding material in a process known as relaxation (Figure 2.1 C).

![Figure 2.1](image)

**Figure 2.1**, Interactions between RF Energy and Magnetic Nuclei in a Magnetic Field (Sprawls, 1987): Parallel vertical arrows indicate the direction of the main magnetic field ($B_0$). A.) When placed inside $B_0$ a magnetic nuclei (red sphere) will align $B_0$ and precess (i.e. spin) around its magnetic dipole (small arrow). B.) When RF energy is applied at the Larmor frequency the nuclei enters a state of increased energy causing alignment away from $B_0$. C.) A nuclei in this excited state will then transfer energy to surrounding material, returning to alignment with $B_0$ through a process known as relaxation.

Relaxation rates of individual nuclei are influenced by the structure surrounding them and differ across various tissues. These differing relaxation rates are a key component in producing the contrasts seen in MR images of the brain.

### 2.2 Magnetic Characteristics of Tissue

When tissue is placed within a magnetic field the nuclei of that tissue will align in parallel to the main direction of the field. Although only a small proportion of nuclei will come into alignment a net magnetization vector ($M_0$) is formed within the tissue voxel. An equilibrium state known as longitudinal magnetization occurs when $M_0$ is aligned with $B_0$. If nuclei are perturbed out of this state via application of RF energy they will seek to regain longitudinal magnetization through relaxation. This relaxation
occurs at an exponential rate. As a convention, the time needed for recovery of 63% of the maximum longitudinal magnetization energy is known as $T_1$ (Figure 2.2). The angle at which a nuclei aligns itself in relation to $B_0$ as a result of an RF pulse is known as the flip angle (FA) or $\alpha$. Application of a 90 degree RF pulse (i.e. FA=90) results in longitudinal magnetization being reduced to zero. This unstable condition is known as saturation and will lead to longitudinal magnetization being recovered exponentially through relaxation.

When RF energy is applied to nuclei they will enter an excited state in which $M_0$ is not aligned with $B_0$, this is known as transverse magnetization. This excited state is unstable and quickly decays via a relaxation process known as free induction decay (FID). The time required for 63% of the initial transverse magnetization to dissipate after excitation is known as $T_2$ (Figure 2.3). Due to its instability, transverse magnetization decay occurs much quicker that $T_1$ relaxation. Therefore, $T_2$ values are considerably less than $T_1$ values for a majority of tissue types.
Differing $T_1$ and $T_2$ relaxation times across tissues are what lead to contrast within MRI images. In basic MRI sequences contrast is controlled via two factors Time of Repetition (TR) and Time to Echo (TE). TR is the length of one image acquisition cycle as well as the interval between the start of longitudinal relaxation and signal measurement. On the other hand, TE is the time interval between the beginning of transverse relaxation and signal measurement.

In addition to $T_1$ and $T_2$ the concentration of protons within a particular voxel (i.e. proton density) is a third factor that can influence MRI image contrast. Proton density contrast is relatively weak compared to $T_1$ contrast early in longitudinal relaxation. Therefore, proton density images are best captured with long TR times (e.g. three or more $T_1$ intervals) when longitudinal magnetization approaches its maximum value.

Images designed to maximize $T_1$ contrast are known as $T_1$ weighted, and are a commonly used form of structural MRI. $T_1$ weighted images can be formed by application of an RF pulse at the Larmor frequency resulting in excitation. From this state different tissues will recover longitudinal magnetization at different rates. The
physical composition of the voxel, the molecular size of the nuclei within that voxel, and the strength of the magnetic field combine to determine these $T_1$ values. For example, fluids (such as cerebral spinal fluid [CSF]) have relatively long $T_1$ values while the $T_1$ of lipids (i.e. fat) is relatively short.

![Diagram of Development of T1 Contrast between Two Tissues](image)

Figure 2.4, Development of $T_1$ Contrast between Two Tissues (Sprawls, 1987): After the application of a 90 degree RF pulse two tissues return to a state of equilibrium at different relaxation rates (250 and 500 mSec). TR is set to a time at which contrast between the two different tissues is maximized producing contrast in the pixel (i.e. voxel) intensities. Images that maximize $T_1$ contrast are often used in structural MRI research.

Setting a TR to capture RF signal at a point where $T_1$ contrast is significantly different across tissues, prior to a voxel fully regaining longitudinal magnetization, results in a $T_1$ weighted image (Figure 2.4). Within these images tissue with a short $T_1$ (e.g. white matter [WM]) will appear brighter than those with longer $T_1$ values (e.g. grey matter [GM]). The choice of flip angle ($\alpha$) is an important factor in producing $T_1$ weighted images. For each value of $T_1$ the optimum flip angle for acquiring signal is known as the Ernst angle and is calculated as,

$$\alpha_{Ernst} = \arccos \left( e^{-\frac{TR}{T1}} \right)$$

Additional MRI contrasts can be produced by utilizing differences in transverse magnetization relaxation rates across tissues. These images are known as $T_2$ weighted,
with contrast being dependent on the TE selected by the user. Furthermore, inherent inhomogeneity of $B_0$ within each individual voxel will cause dephasing of transverse magnetization at a rate much faster than would be indicated by the intrinsic $T_2$ characteristics of the tissue. The actual transverse relaxation time taking into account this dephasing effect is known as $T_{2^*}$, which is always less than $T_2$. Contrast resulting from $T_{2^*}$ is a key property in the imaging of changes in blood oxygenation, known as functional MRI (fMRI).

### 2.3 Magnetic Resonance Images

Magnetic resonance images are three-dimensional volumes comprised of individual units (i.e. voxels) that encode signal intensity resulting from emitted NMR RF energy. MRI protocols can be described in two main phases, namely signal acquisition and image reconstruction. During signal acquisition a particular sequence of RF pulses and magnetic field gradients (i.e. controlled inhomogeneities in $B_0$) are applied. Differences between MRI methods are determined by these pulse sequences. Here I will describe some characteristics of basic MRI pulse sequences followed by spatial encoding of MR signal and MR image reconstruction.

#### 2.3.1 Basic Magnetic Resonance Imaging Pulse Sequences

The process of MRI signal acquisition can be separated into longitudinal and transverse magnetization phases. Transition between these two phases is produced by the application of an RF pulse characterized by $\alpha$, causing excitation. The transverse magnetization phase concludes with the echo event, which is produced by either the application of RF pulses (e.g. spin-echo, inversion recovery methods) or field gradient (i.e. gradient-echo methods) and results in the emission of RF signal.

Basic spin-echo pulse sequences are characterized by the application of a 90-degree excitation pulse followed by a 180-degree pulse that produces the echo event and RF signal (Figure 2.5). Spin-echo imaging methods can be used to produce $T_1$, $T_2$, or proton density weighted images by manipulating TR and TE. For example, $T_1$ weighted images require a short TR and TE, proton density images a long TR and short TE, and $T_2$ weighted images a long TR and TE.
Figure 2.5, Development of Image Contrast in a Spin-Echo Sequence (Sprawls, 1987): Here the imaging cycle begins with the application of a 90 degree RF pulse. Tissue A and B then recover longitudinal magnetization at differing rates determined by their $T_1$, producing $T_1$ contrast. As longitudinal magnetization reaches its maximum, proton density becomes the key factor in contrast between tissues A and B. The end of this imaging cycle is indicated by the application of another 90-degree RF pulse, converting longitudinal magnetization to transverse magnetization. At the start of transverse relaxation tissue A and B have different magnetization as a result of differing $T_1$ and proton density characteristics. However, due to different $T_2$ properties between the tissues as transverse relaxation progresses $T_2$ contrast develops. At a time selected by the user an echo event is triggered by the application of a 180 degree RF pulse resulting in the emission of RF signal. Adjusting TR and TE times will result in different weighted images (i.e. $T_1$, $T_2$, proton density).

In addition to the 90-degree excitation pulse and 180-degree RF pulse used in basic spin-echo techniques, inversion recovery (IR) MRI protocols employ an additional 180-degree pulse at the beginning of the scanning sequence (Figure 2.6). The time between the initial 180-degree and 90-degree pulses is known as the inversion time (TI). There are two main benefits in the application of IR sequences that result from the longer longitudinal magnetization relaxation rates they provide. 1) IR sequences with a short TI can be used to nullify signal from tissues that have a relatively short $T_1$ (e.g. fat). 2) IR sequences with long TI give more time for $T_1$ contrast to develop. This makes IR MRI particularly well suited to produce $T_1$ weighted images commonly used in structural MRI analyses. The images used throughout this thesis are a form of synthetically calculated IR MRI (see section 2.4).
Inversion recovery (IR) sequences operate in a similar manner to spin-echo sequences but begin with the application of a 180 degree inversion recovery RF pulse. This causes increased longitudinal relaxation times as the process begins from a negative (inverted) value. Short inversion times (TI) (i.e. short time inversion recovery [STIR]) can be used to eliminate signal from tissues with short $T_1$ values (e.g. fat) while longer TI can be used to increase $T_1$ contrast.

Differing from spin-echo techniques, gradient-echo sequences are characterized by their application of a RF excitation pulse with a relatively small $\alpha$ (i.e. >90-degrees). Flip angles of less than 90-degrees cause less longitudinal magnetization to be converted into transverse magnetization. Therefore, over a short TR more longitudinal magnetization and less transverse magnetization signal intensity can be captured, with net signal intensity being the combination of the two. The relatively short TR needed to acquire these images gives the benefit of reduced total acquisition time.

Gradient-echo sequences employ a single RF energy pulse per imaging cycle in order to not affect the longitudinal magnetization. However, when TR is set to be only slightly longer than TE time it is possible for an echo event to be produced by each excitation pulse. Thus, transverse magnetization in a gradient-echo sequence has two components; free induction decay ($T_2^*$) and the spin-echo event resulting from the previous RF pulse ($T_2$). By not applying a second RF pulse during each imaging cycle gradient-echo differs from spin-echo techniques in their handling of transverse magnetization.
In order to produce $T_1$ weighted images with a gradient-echo sequence the spin-echo component resulting from carryover transverse magnetization across imaging cycles is spoiled (i.e. eliminated) by altering the phase relationship of the RF excitation pulses or through the application of dephasing gradient pulses. Popular gradient-echo sequences used for structural MRI include gradient recalled acquisition in steady state (GRASS) and spoiled gradient recalled-echo (SGPR).

2.3.2 Spatial Encoding and Image Reconstruction of Magnetic Resonance Signal

During the acquisition phase NMR RF signal from a particular voxel is encoded with distinct frequency and phase properties in order to identify the spatial origin of the signal during image reconstruction. Frequency and phase encoding are achieved through the application of magnetic field gradients (i.e. controlled changes in field strength) at specific times along three orthogonal directions (i.e. x, y, and z) within a MRI scanner.

A gradient applied along the direction of $B_0$ can be used to selectively image a particular slice of tissue. Tissue across the gradient will experience minor differences in field strength and therefore have slightly different resonant frequencies. Thus, over the course of a single TR an excitation pulse can be applied at different frequencies to selectively excite and capture signal from each slice of tissue. A phase encoding gradient can then be applied in order to alter the precession rate in voxels across this gradient. The phase encoding gradient is only applied for a short amount of time leaving each voxel in a different phase (i.e. position) of precession (i.e. phase encoded). In order to produce a distinct phase relationship for each voxel in an image phase encoding must be repeated $n$ times, where $n$ = the number of voxels. Last, the application of a frequency encoding gradient at the time of the echo event when signal is captured is used to isolate signal from a particular column of voxels within a slice. The frequency encoding gradient causes magnetic nuclei to precess at different rates and therefore produce different signal frequencies across the gradient.
Combined effects of slice, phase, and frequency encoding gradients cause individual voxels to have unique differences in phase in one direction and frequency in another. Thus, while signal from all voxels within an MRI image are captured as a single composite signal and stored in what is known as $K$ space a Fourier transformation can be used to isolate the spatial location of a particular voxel’s signal from the composite signal. The spatial sorting of the composite RF signal captured by the MRI scanner in $K$ space into an image via Fourier transformation comprises the most commonly applied form of MR image reconstruction.

Up to this point of the chapter I have described elements common to several standard structural MRI methods. I will now describe the quantitative relaxometry MRI technique used to produce synthetic $T_1$ weighted IR images described in this thesis, namely Driven Equilibrium Single Pulse Observation of $T_1$ and $T_2$ (DESPOT1 and DESPOT2) (Deoni et al., 2005, Deoni et al., 2003).

2.4 Driven Equilibrium Single Pulse Observation of $T_1$ and $T_2$ Mapping

Traditional structural MRI methods result in images that display qualitative differences in $T_1$ or $T_2$ contrast between tissue types. Alternatively intrinsic $T_1$ and $T_2$ relaxation times can be determined and mapped in a voxel-wise manner (Breger et al., 1986, Johnson et al., 1987, Kjær et al., 1987). MR sequences that quantify relaxation times are commonly referred to as relaxometry or quantitative mapping methods. These techniques offer several benefits over traditional structural MRI. For example, contrast within a synthetic $T_1$ weighted image produced using quantitative mapping is determined solely by differences in $T_1$ properties of the tissue being imaged with extraneous effects such as $T_2$ and proton density removed. Furthermore, relaxometry methods are less sensitive to hardware-related effects (i.e. RF coil sensitivity, electronic amplifier gains etc.) that may vary across scanners or even scanning sessions. This makes these methods ideal for comparison of images acquired across different scanning sites or for longitudinal acquisitions (Deoni et al., 2008).

The structural MRI data described within this thesis was acquired through a DESPOT1 sequence (Deoni et al., 2005, Deoni et al., 2003). In order to determine $T_1$ at each voxel, DESPOT1 acquires a series of SPGR images with a range of flip angles and constant TR. Signal from an SPGR can be calculated given,
\[
\frac{SI_{SPGR}}{\sin \alpha} = \frac{SI_{SPGR}}{\tan \alpha} \times E_{-1} + M_o(1 - E_{-1})
\]

Where \(SI_{SPGR}\) is the SPGR signal intensity given flip angle \(\alpha\), \(M_o\) is a factor proportional to the equilibrium longitudinal magnetization, and \(E_{-1} = \exp \left( -\frac{TR}{T_1} \right)\). By plotting \(\frac{SI_{SPGR}}{\sin \alpha}\) versus \(\frac{SI_{SPGR}}{\tan \alpha}\) \(T_1\) can be calculated from the slope of the resulting line,

\[
T_1 = -\frac{TR}{\ln(\text{slope})}
\]

The results of DESPOT1 mapping were then used to produce synthetic IR \(T_1\) weighted images using the following steps. 1.) Signal from non-brain tissue was removed using the brain extraction tool (BET) (Smith, 2002), 2.) intra-session images were non-linearly co-registered using an automated three-dimensional multi-scale approach based on normalized cross-correlation (Collins et al., 1994). 3.) \(T_1\) maps were then calculated in the ImageJ (http://rsb.info.nih.gov/ij/) analysis package using plug-ins freely available at www.fmrib.ox.ac.uk/~sdeoni/ (Deoni et al., 2008). This process resulted in the formation of high-resolution structural \(T_1\) weighted “synthetic” IR images, with 1x1x1mm resolution, a 256x256x176 matrix, TR=1800ms, TI=50ms, FA=20°, and FOV=5cm.

2.5 Cortical Surface Reconstruction using Freesurfer Software

In neuroimaging, morphometry methods comprise techniques that investigate the size and shape of the brain and its structures. Surface Based Morphometry (SBM) relies on reconstructions of the cortical surface from MRI images to quantify different structural aspects of brain anatomy. Here, I describe the pre-processing steps used to generate SBM models of the brain and morphometric features calculated using these models.

Volumetric segmentation and reconstruction of tessellated models of the cortical surface from the \(T_1\) weighted structural MRI images described above was performed using the Freesurfer image analysis suite, which is documented and freely available for download online (http://surfer.nmr.mgh.harvard.edu/). The technical details of the
Freesurfer segmentation and reconstruction procedures are described in prior publications (Dale et al., 1999, Dale and Sereno, 1993, Fischl and Dale, 2000, Fischl et al., 2001, Fischl et al., 2002, Fischl et al., 2004a, Fischl et al., 1999a, Fischl et al., 1999b, Fischl et al., 2004b, Han et al., 2006, Jovicich et al., 2006, Ségonne et al., 2004, Reuter et al., 2010, Reuter et al., 2012) and have been demonstrated to show good test-retest reliability across scanner manufacturers and field strengths (Reuter et al., 2012, Han et al., 2006).

Briefly, the Freesurfer processing pipeline includes motion correction and averaging (Reuter et al., 2010) of one or more volumetric T1 weighted MR images, removal of non-brain tissue using a hybrid watershed/surface deformation procedure (Ségonne et al., 2004), automated Talairach transformation, segmentation of the subcortical white matter and deep grey matter volumetric structures (including hippocampus, amygdala, caudate, putamen, ventricles) (Fischl et al., 2004a, Fischl et al., 2002), intensity normalization (Sled et al., 1998), tessellation of the grey-white matter boundary, automated topology correction (Ségonne et al., 2007, Fischl et al., 2001), and surface deformation following intensity gradients to optimally place the grey-white and grey-cerebrospinal fluid borders at the location where the greatest shift in intensity defines the transition to the other tissue class (Dale et al., 1999, Dale and Sereno, 1993, Fischl and Dale, 2000). Once the cortical models are complete, a number of deformable procedures can be performed for further data processing and analysis including surface inflation (Fischl et al., 1999a), registration to a spherical atlas which is based on individual cortical folding patterns to match cortical geometry across subjects (Fischl et al., 1999b), and parcellation of the cerebral cortex into units with respect to gyral and sulcal structure (Desikan et al., 2006, Fischl et al., 2004b).

Within this thesis tessellated models of the cortical surface were used to calculate a variety of morphometric measures of brain anatomy for later analysis. 1) Cortical thickness (CT) is calculated as the closest distance from the grey-white boundary to the grey-CSF boundary at each vertex on the tessellated surface (Fischl and Dale, 2000). CT maps are created using spatial intensity gradients across tissue classes and are therefore not simply reliant on absolute signal intensity. These maps are not restricted to the voxel resolution of the original data thus are capable of detecting submillimeter differences between groups. Freesurfer procedures for the measurement
of CT have been validated against histological analysis (Rosas et al., 2002) and manual measurements (Salat et al., 2004, Kuperberg et al., 2003). 2) Measures of cortical pial and white matter surface areas (SA) are derived using a pycnophylactic interpolation method described in detail by Winkler et al. (2012). 3) Measures of cortical volume are calculated as the product of pial SA and CT. This Freesurfer approach to calculating volume differs from traditional imaging measures of brain volumes such as VBM in which volume is calculated by counting voxels inclusive to segmentations of GM, WM, and CSF as determined by signal intensity.

Additional measures of the cortical surface may be used to describe the topography of the convolutions in the cortical sheet. These include 4) average convexity or concavity (SULC), which indicates the depth/height at each vertex above the average cortical surface, measuring sulcal depth or gyral height respectively. SULC captures large-scale cortical geometric features and is insensitive to noise in the form of small wrinkles in a surface and can be used to quantify the primary folding pattern of the cortical surface. 5) Mean radial curvature (CURV) quantifies folding of the small secondary and tertiary folds in the cortical surface that would not be captured by measures of large-scale geometric features (i.e. SULC measures). 6) Metric distortion (i.e. Jacobian) is calculated as the degree of displacement and convolution of the cortical surface relative to the average template. Unlike SULC and CURV, which measure specific aspects of cortical geometric features (e.g. folding patterns of sulci and gyri), metric distortion estimates the overall degree of folding including geometric distortions otherwise not specified. 7) Further estimates of the degree of cortical gyrification can be achieved through calculation of the local gyrification index (/GI). Methods used to calculate /GI have been previously validated by Schaer et al. (2008). In brief, /GI measures quantify of the proportion of pial surface invaginated within sulcal folds via a ratio of the outer to pial surface area within spherical regions of interest placed across each vertex on the cortical surface.

It is critical to note that all the above measures are susceptible artefacts caused by movement during image acquisition. The effects of these artefacts, such as motion induced blurring, are not trivial and are particularly relevant when studying conditions associated with atypical movement such as ASD. To date, outside of eliminating
motion during acquisition adequate solutions for dealing with these artefacts are limited (Reuter et al. 2015).

2.6 Mass Univariate Magnetic Resonance Imaging Analyses

Several different methodologies can be employed for analysis of MRI data. Often the goal of these methods is the identification and mapping of differences related to a variable(s) of interest across each individual voxel or vertex. These techniques are referred to as mass univariate methods as they involve repeating univariate statistical tests across each voxel or vertex. Applications of multiple regression, more specifically the general linear model (GLM) (Friston et al., 1994a) are among the most popular methods for this type of MRI investigation. Within this section I will introduce the GLM as well as discuss ways of dealing with statistical limitations that result from calculating significance in mass univariate analyses (i.e. the multiple comparison problem).

2.6.1 The General Linear Model

The GLM builds upon simple linear regression methods for identifying associations between predictor variables and responses. An example first order regression model with two predictor variables can be given as,

\[ Y_i = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2} + \epsilon_i \]

(2.5)

Where \( Y_i \) is the response given predictor variables \( X_{i1} \) and \( X_{i2} \) in the \( i \)th sample. The model parameters in this example are \( \beta_0, \beta_1, \) and \( \beta_2, \) with an error term \( \epsilon_i. \) The \( \beta \) parameters may be referred to as partial regression coefficients as they reflect the partial effect of the corresponding predictor variable when all other predictor variables within the model are held constant. Assuming \( E(\epsilon_i) = 0, \) the regression function (i.e. regression surface or response surface) for this model example is,

\[ E(Y) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 \]

(2.6)

Points on this regression surface correspond to the mean response \( Y \) given any combination of \( X_1 \) and \( X_2. \)
A GLM may be given using the same formula as the example above. However, the variables (e.g. \( X_1, X_2 \)) need not only represent qualitative predictor variables. For quantitative (i.e. categorical) variables binary indicator variables are used to identify different classes (e.g. male=0, female=1; patient=0, control=1, etc.). Furthermore, if the effects of predictor variables are not additive (i.e. the effect of one variable is dependent on the level of another) an interaction term can be included.

\[
Y_i = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2} + \beta_3 X_{i1} X_{i2} + \epsilon_i
\]

Here \( \beta_3 \) represents the interaction effect between variables \( X_1 \) and \( X_2 \) on \( Y \). The GLM may also take into account non-linear effects through the use of polynomials. For example, quadratic or cubic relationships can be investigated using \( X_i^2 \) or \( X_i^3 \) respectively,

\[
Y_i = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i1}^2 + \beta_3 X_{i2}
\]

The GLM encompasses most types of statistical inference including the \( t \)-test and analysis of variance (ANOVA). An ANOVA can be given in GLM terms when a predictor variable (i.e. independent variable) is categorical. Within an ANOVA this categorical variable is known as a factor and the categories as levels of the factor. Under GLM framework these levels are coded using dummy variables. For example, an ANOVA with a factor comprised of three levels can be given as,

\[
Y_i = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2}
\]

Here, \( X_i \) codes for the first level such that \( X_i=1 \) if the observation is within this level or \( X_i=0 \) if not. The same applies for the second level, which is coded for by \( X_2 \). The third level’s mean is predicted by \( \beta_0 \) and therefore is not coded.

Using the framework outlined above the GLM is able to take into account any number and combination of linear, non-linear, and/or interactions terms within a single model. Thus, the GLM offers flexibility to test for a variety of different effects. Some examples include; group comparisons, effects of age or gender, relationships between
age or gender and diagnosis, and correlations with specific behavioral measures or symptom metrics.

In order to perform a particular statistical test using a GLM, data included in the model is first defined as a design matrix where each column corresponds to a qualitative variable or indicator variable. For example, for the following GLM,

\[ Y_i = \beta_0 + \beta_1 Age_{i1} + \beta_2 Gender_{i2} + \beta_3 Age_{i1} Gender_{i2} + \varepsilon_i \]

a corresponding design matrix would be given as,

<table>
<thead>
<tr>
<th>( \beta_0 )</th>
<th>Age</th>
<th>Male</th>
<th>Female</th>
<th>Age * Male</th>
<th>Age * Female</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>23</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>23</td>
</tr>
<tr>
<td>1</td>
<td>24</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>24</td>
</tr>
<tr>
<td>1</td>
<td>29</td>
<td>1</td>
<td>0</td>
<td>29</td>
<td>0</td>
</tr>
<tr>
<td>1</td>
<td>30</td>
<td>1</td>
<td>0</td>
<td>30</td>
<td>0</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
</tbody>
</table>

Significance of an effect at each voxel or vertex \( i \) can be tested using a \( t \) test (Student, 1908) of linear contrasts between conditions. Using the above design matrix as an example, to test for a difference between genders we would use the contrast \( c = [0 \ 1 \ -1 \ 0 \ 0] \), for a main effect of age \( c = [1 \ 0 \ 0 \ 0 \ 0] \), or an interaction effect between age and gender \( c = [0 \ 0 \ 0 \ 1 \ -1] \). The significance of each of these effects at data point \( i \) is then tested with,

\[ t_i = c \cdot b_i / \varepsilon_i \]  

Where \( b_i \) is the corresponding parameter estimates at data point \( i \). This \( t \) statistic and the number of degrees of freedom can then be used to calculate the probability of significance (i.e. \( p \) value) for an effect at a particular vertex or voxel. Spatial mapping of \( t_i \) is referred to as statistical parametric mapping (SPM) and reflects data points that are statistically related to the user defined contrast (Friston et al., 1994a, Worsley et al., 1996).
Within this thesis, GLM analyses of morphometric brain features were conducted using the SurfStat toolbox (http://www.math.mcgill.ca/keith/surfstat/) implemented in MATLAB (R2014a; MathWorks).

2.6.2 Correcting for Multiple Comparisons

Mass univariate analyses of brain imaging data apply a large number of independent statistical tests. Therefore, probability theory dictates that a large degree of false positives will be detected in these analyses (type I errors). For example, a \( p \)-value of 0.05 denotes a 5\% type I error rate. Setting a significance threshold of \( p = < 0.05 \) one could expect up to 5 false positives for every 100 tests. In vertex wise analyses it is common to employ over 200,000 independent statistical tests, which leads to an unacceptably high rate of expected type I errors (approximately 10,000). Addressing this so called multiple comparison problem is critical for valid interpretation of voxel or vertex-wise neuroimaging analyses (Bennett, 2010).

Due to the spatial relationships inherent between voxels or vertices within neuroimaging data (caused by biology, signal acquisition, image smoothing etc.) multiple comparison corrections are often performed on clusters of spatially contiguous significant statistical values. This allows for using a type I error rate that determines the likelihood that a cluster or ‘family’ of spatially contiguous significant statistics has arisen by chance, i.e. a Family-Wise Error (FWE) rate. A traditional method of testing against the null hypothesis using FWE is Bonferroni correction. This correction adjusts the tolerable type I error rate (\( \alpha \)) for an individual test in order to account for the total number of independent tests (\( n \)) such that the corrected \( p \) value for an individual test is equal to \( \frac{\alpha}{n} \). However, Bonferronni correction is overly conservative (i.e. prone to type II errors) for most neuroimaging analyses. This is due to fewer independent observations (i.e. functional or structurally related brain regions) existing compared to the actual number of voxels or vertices. Furthermore, estimation of the number of independent observations in a particular image is hindered by spatial smoothing (i.e. averaging or ‘blurring’ of the image) which is commonly implemented to increase the signal to noise ratio in image analyses. Therefore, methods have been developed to correct for the multiple comparison problem while taking into account the smooth spatial covariance that results from SPM, one example being random field theory (RFT) (Friston et al., 1994b).
In brief, given a significant cluster of statistical values (i.e. $t$ scores) RFT calculates the height threshold for a smoothed statistical map that gives the desired FWE rate (e.g. $p<0.05$). The height threshold is considered to be the individual statistical value necessary within a cluster in order to reject the null hypothesis that all points in the cluster originated from the null distribution. The process of RFT can be explained by three key stages being: 1) the calculation of the number of resolution elements in an image, these elements being analogous to independent observations (Worsley et al., 1992). 2) The calculation of the Euler characteristic (EC), the EC being equivalent to the number of significantly contiguous clusters of vertices or voxels post thresholding (Worsley, 1994). 3) The application of RFT and subsequent thresholding of significant clusters that do not meet the selected FWE rate. This method of correction has been applied to several types of neuroimaging data including non-isotropic images such as cortical surface based reconstructions (Worsley et al., 1999).

2.7 Multivariate Pattern Classification
Mass univariate analyses offer a powerful tool for investigating differences between groups. However, these methods are not suited to make predictions for individual cases regarding group membership or clinical outcome measures, and thus offer limited translational and clinical value. Alternatively, ‘machine learning’ based pattern recognition methods can be used to generate image-based predictive models that have the potential to be utilized as diagnostic tools and intervention outcome measures.

Within ‘machine learning’ an algorithm is said to learn from experience $E$ with respect to some class of tasks $T$ and performance measure $P$. Learning is said to occur when performance in task $T$ improves with experience $E$ as measured by performance $P$ (Mitchell, 1997). For example, a diagnostic model using pattern recognition could seek to learn to make diagnostic predictions ($T$), based on MRI scans of the brains of patients and controls ($E$), with performance ($P$) being measured by diagnostic accuracy. In supervised learning problems examples from a training set of empirical data are mapped to an output. These outputs may be either a continuous variable, in the case of a regression problem, or class label, in the case of a classification problem.
Within this section I will first describe two multivariate pattern classification (MVPC) algorithms implemented in this thesis, namely the Support Vector Machine (SVM) (Vapnik, 1995, Boser et al., 1992) and Gaussian Process Classifier (GPC) (Rasmussen and Williams, 2006, Marquand et al., 2010). While several MVPC algorithms are available, the SVM and GPC were chosen for this thesis as they have previously been shown to have good performance in neuroimaging problems owing to their ability to handle large dimensionality data in a computationally efficient manner when compared to other options (e.g. neural networks, random forests). I will then discuss potential ways for improving MVPC model performance through feature selection and parameter tuning, in addition to model validation and visualization methods.

2.7.1 The Support Vector Machine

The support vector machine (SVM) is among the most popular algorithms used for MVPC of neuroimaging data. SVMs benefit in their ability to train non-linear predictive models in high dimensional spaces that offer good generalization performance and computational efficiency. The SVM attempts to solve a supervised learning problem by optimally placing a boundary known as a hyper-plane (in cases with more than three dimensions) to separate given examples \( x_i \) according to class labels \( y \in \{+1, -1\} \) (e.g. patients = +1 controls = -1). The SVM is referred to as a maximum margin classifier as it optimizes the placement of the hyper-plane such that the margin between the two classes is maximized (Figure 2.7). I will now briefly describe the SVM. More detailed coverage has been provided by Bishop (2007) and Smola and Schölkopf (1998).
The decision function for a linear SVM can be given as,

\[ f(x) = \mathbf{w}^T \mathbf{x} + b \]  

(2.12)

where \( \mathbf{w} \) is a vector that is orthogonal to the hyper-plane that separates the two classes, \( \mathbf{x} \) is an example data point and \( b \) is a constant offset term that controls the distance of the hyper-plane from the origin. For data points that lie on the hyper-plane \( f(x)=0 \). For other examples the sign of \( f(x) \) denotes the class \( y \in \{+1, -1\} \) of \( x \) such that \( x_{y+1} = f(x) > 0 \) and \( x_{y-1} = f(x) < 0 \). The SVM aims to maximize the margin between classes by finding, given perfect classification, the vector \( \mathbf{w} \) with the smallest magnitude \( ||\mathbf{w}|| \). This vector details the contribution of each data point in a feature set (e.g. vertex or voxel) towards classification and is known as the weight vector.

Within training samples one or more samples may prevent clean separation between classes. This may be due to noise in the inputs (uncertainty in the location of samples) and/or outputs (mislabeling an sample) that cause a sample to be plotted on the “wrong side “of the decision plane. In cases where linear separation between cases is not possible the introduction of ‘slack variables’ can be used to ‘soften’ the margin. This can be achieved by introducing a trade off between maximizing the margin and the total number of acceptable miss classifications. This trade off introduces an
optimization problem, which can be viewed from either the primal or dual perspective (Boyd and Vandenberghe, 2004). The soft margin SVM in primal form can be given as,

\[
\begin{align*}
\text{minimize}_{\mathbf{w}, b}: & \quad \frac{1}{2} \| \mathbf{w} \|^2 + C \sum_{i=1}^{n} \xi_i \\
\text{subject to:} & \quad y_i (\mathbf{w}^T \mathbf{x}_i + b) \geq 1 - \xi_i, \\
& \quad \xi_i \geq 0, \quad i = 1, ..., n
\end{align*}
\]

where we seek to minimize the term \( \frac{1}{2} \| \mathbf{w} \|^2 \), which is achieved when the margin that separates between classes is maximized. In a hard margin scenario we impose the constraint \( y_i (\mathbf{w}^T \mathbf{x}_i + b) \geq 1 \), such that the decision function output \( f(\mathbf{x}_i) = \mathbf{w}^T \mathbf{x}_i + b \) multiplied by the label \( t_i \) (either +1 or -1) must be greater or equal to one, thus no sample can be closer than the margin marked by the \( y(\mathbf{x}) = 1 \) and \( y(\mathbf{x}) = -1 \) contour lines. In 2.13 \( \xi_i \) are non-negative slack variables, which loosen the constraint on the minimal distance (i.e. 1) that an example may fall from the margin, such that a distance of \( 1 - \xi_i \) is now acceptable. The trade-off between maximizing the margin and minimizing the number of classifications that violate the margin are controlled by a regularization parameter \( C \). As \( C \) increases, priority is placed on reducing the number of misclassifications over maximizing the margin such that when \( C=\infty \) the SVM will have zero tolerance for these violations (i.e. ‘hard margin’ SVM).

The optimization problem (2.12) can be solved it its duel form using the following Lagrange function,

\[
L(\mathbf{w}, b, \xi, \alpha, \mu) = \frac{1}{2} \| \mathbf{w} \|^2 + C \sum_{i=1}^{n} \xi_i - \sum_{i=1}^{n} \alpha_i \{ y_i (\mathbf{w}^T \mathbf{x}_i + b) - 1 + \xi_i \} - \sum_{i=1}^{n} \mu_i \xi_i
\]

where \( \alpha_i \geq 0, \mu_i \geq 0 \) for \( i = 1, ..., n \) are the Lagrange multipliers (the Lagrangian \( L \) is a scalar) that the function seeks to minimize with respect to maximizing \( \mathbf{w} \) and \( b \). The solution of (2.14) requires the resulting partial derivatives,
The first of these partial derivatives allows for the optimal weight vector to be expressed as a weighted combination of all examples \( x \), with the class label \( y_i \) determining the sign and the \( \alpha_i \)'s determining the magnitude of the contribution of each example to the weight vector, such that

\[
\frac{\partial}{\partial \mathbf{w}} L(\mathbf{w}, b, \xi, \alpha, \mu) = 0,
\]

\[
\frac{\partial}{\partial b} L(\mathbf{w}, b, \xi, \alpha, \mu) = 0.
\]

The second partial derivative in (2.15) leads to the condition that the sum of the signed weights \( \alpha_i \) and \( y_i \) equal zero, such that

\[
\sum_{i=1}^{n} \alpha_i y_i = 0.
\]

Expressions (2.16) and (2.17) allow us to remove the primal variables \( \mathbf{w} \) and \( b \) from the optimization problem outlined in (2.13), giving the dual form of (2.13) as

\[
\text{maximize}_\alpha: W(\alpha) = \sum_{i=1}^{n} \alpha_i - \frac{1}{2} \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j y_i y_j x_i^T x_j
\]

subject to: \( 0 \leq \alpha_i \leq C, \quad i = 1, \ldots, n \)

\[\sum_{i=1}^{n} \alpha_i y_i = 0.\]

The optimization problem (2.13) in dual form (2.18) now seeks to maximize an objective function of the support vector weights \( \alpha_i \). For these optimization problems the constraint that \( \alpha_i (y_i f(\mathbf{x}_i) - 1 + \xi_i) = 0 \) must be satisfied for each example \( \mathbf{x}_i \) by one of two ways. 1) When for a given example \( \mathbf{x}_i \), \( \alpha_i = 0 \) it is given that \( y_i f(\mathbf{x}_i) \geq 1 - \xi_i \) and therefore \( \mathbf{x}_i \) is well classified. 2) When \( \alpha_i > 0 \) it is given that \( y_i f(\mathbf{x}_i) = 1 - \xi_i \).
1 − ξ_i, therefore x_i is either on the margin (y_if(x_i) = 1, slack variable ξ_i = 0) or within the margin (y_if(x_i) = 1 − ξ_i, slack variable ξ_i > 0) (Bishop, 2007). This highlights the sparse nature of SVM in that all the information relevant to the decision problem (i.e. w) is carried by the relatively few examples that fall on or within the margin (i.e. the support vectors).

As training examples x_i and x_j only enter into the optimization problem (2.18) through their inner product it is possible to replace this inner product with a kernel function (K). This process, commonly referred to as ‘kernel trick’, allows mapping into a potentially higher dimensionality space in which a non-linear decision plane can be derived. The simplest form of K is a dot product between the input data and a test case,

\[ K(x_i, x) = x_i \cdot x \]

For cases in which classes cannot be separated with a linear decision boundary (even with the aid of a soft margin) the resulting linear kernel K provides a feature space with higher dimensionality in which non-linear separation of the classes may be achievable. It is also possible to transform data from native space using non-linear kernels (e.g. the radial basis function). However, in cases where the number of features is larger than the number of observations non-linear kernels typically offer marginal if any improvement in classification accuracy (Hsu et al., 2003) and are thus not typically applied in neuroimaging problems.

**2.7.2 Gaussian Process Classifiers**

Pattern classification methods such as the SVM offer power tools to make binary predictions according to class labels. However, in the clinical setting having an estimate of confidence in a prediction is often just as important as the prediction itself. Probability theory offers a coherent way to manage uncertainty and can be introduced into prediction modeling though the application of Bayesian statistics. Bayes’ theorem describes the conditional probability of an event given another event is true,

\[ p(A|B) = \frac{p(B|A)p(A)}{p(B)} \]
here $A$ and $B$ are events, $p(B)\neq 0$. $p(A)$ and $p(B)$ are the probabilities of event $A$ and $B$ independent of each other. $p(A|B)$ and $p(B|A)$ represent conditional probabilities, i.e. the probability of $A$ given $B$ is true and vice versa (Kendall et al., 1968). Within MVPC the Gaussian process classifier (GPC) (Rasmussen and Williams, 2006) is one Bayesian method that can make probabilistic estimates as to class ownership. This ability gives GPC methods potential clinical benefits over the SVM, which is not a probabilistic prediction device.

According to Rasmussen and Williams (2006), a Gaussian process (GP) can be viewed as an extension of a Gaussian probability distribution. Where as a Gaussian probability distribution describes finite dimensional variables, the GP extends the probability distribution to functions, which are infinite dimensional, under the constraint that any finite number of variables have a multivariate Gaussian distribution. Therefore a GP describes a distribution over functions that can be used for Bayesian inference. There are two complementary and equivalent views of GP inference, the ‘weight space’ and ‘function space’ views. I will first introduce the function space view of GP inference and later discuss the weight space view in context of classification. In the view of GP function space a GP prior is placed over a set of interpolating functions $f(x)$ the posterior distribution is computed according to Bayes rule. This distribution is then used to explain the relationship between examples (Figure 2.8).

Figure 2.8, The Gaussian Process in a One Dimensional Regression Problem (Marquand et al., 2010, Rasmussen and Williams, 2006): A) Sample functions (dotted lines) selected from a zero-mean Gaussian process prior distribution are shown. B) After the introduction of training cases (black
crosses) sampled functions from the posterior distribution are limited to those that fit the given examples. In both panels the grey shaded area indicates a 95% confidence interval for a prediction at any possible input. One such prediction is shown in blue.

When GP inference is posed as a binary classification problem we wish to predict the class probability for a test example \( x^* \), \( p(y = 1|x^*, x_{i...n}) \) given a set of training examples \( x_{i...n} \) with associated class labels \( y \in \{+1, -1\} \). In order to restrict the GP prior over functions \( f(x) \) (Figure 2.9 A) to give a binary output (i.e. \( y \in \{+1, -1\} \)), \( f \) is ‘squashed’ point wise through a response function (Figure 2.9 B).

![Figure 2.9](image)

**Figure 2.9, The Gaussian Process Classifier (Rasmussen and Williams, 2006):** Here the Gaussian process classifier aims to separate examples \( x_{i...n} \) sampled from two classes \( y \in \{+1, -1\} \). A) A prior GP distribution of \( f(x) \) is given in a 2-d input space. In order to limit predictions to binary classes a response function is used to squash the GP distribution of \( f(x) \). B) Commonly this response function takes the form of a logistic function, \( \lambda(x) = (1 + \exp(-x))^{-1} \). C) The location of examples \( x_{i...n} \) are shown with white dots indicating class +1 and black dots indicating class -1. D) A contour plot depicts predictive probabilities as a function of \( x \) after the implementation of the response function on the GP priors. Black lines indicate the decision boundary (>0.5 = \( y = +1 \), <0.5 = \( y = -1 \)).

The theory and implementation of GPC have been previously described by Ebden (2008) and are covered in detail by Rasmussen and Williams (2006). I will now briefly outline the GPC as described by Marquand et al. (2010).
The GPC is an extension of Gaussian process regression (GPR), which in the perspective of the weight space can be seen as a Bayesian extension of linear regression. In linear regression predictions are made by,

\[ f(x) = x^T w, \quad y = f(x) + \varepsilon \]  

where \( w \) is a vector of weights, \( y \) is a target value, \( f \) is a function value, and \( \varepsilon \sim N(0, \sigma_n^2) \) is a Gaussian noise term. Given input \( x \) the posterior distribution of these weights are computed by Bayes rule,

\[ p(w|X, y, \theta) = \frac{p(w|\theta)p(y|X, w, \theta)}{p(y|X, \theta)} \]  

where \( X \) is a matrix of input examples \( x_{i...n} \) with \( y \), in the case of classification, being a vector of corresponding class labels, \( y = [y_1 \ldots y_m]^T \), and \( \theta \) being a vector of hyperparameters. The term \( p(y|X, \theta) \) is itself a GP and is known as the normalizing constant or marginal likelihood, \( p(y|X, \theta) = \int p(y|X, w)p(w)dw \). Prediction for a particular test case \( x^* \) in data set \( D \) comprising of examples \( x_i \) and corresponding class labels \( y_i \) is performed by averaging all possible values of \( w \) and weighting by their posterior probability, \( p(f^*|D, \theta, x^*) = \int (f^*|w, x^*, \theta)p(w|D, \theta)dw \). Therefore, GP predictions are a weighted average of all possible linear models, under the prior constraints, based on the examples provided by \( X \).

In order to determine the posterior distribution, the spatial relationship of each example to all other examples is first described using a covariance function. There are several options for the form of this function (Rasmussen and Williams, 2006), the one used in this thesis being a linear covariance function,

\[ K = \frac{1}{l^2} XX^T + b \]  

here \( l \) is a length parameter that controls the scaling of the latent function and \( b \) is a bias term that accounts for the offset from zero. To make predictions we make use of
the covariance between training examples and a test case, which are stored in vector \( k^* \). For a regression problem, the posterior predictive distribution can then be calculated as,

\[
p(f^*|D, \theta, x^*) \sim N(\mu, \sigma^2)
\]

\[
\mu = k^{*T} C^{-1} y
\]

\[
\sigma^2 = k(x^*, x^*) - k^{*T} C^{-1} k^*
\]

where \( C = K + \sigma_n^2 I \). For regression problems this posterior predictive distribution is Gaussian. From this distribution we are interested in making a prediction of \( y^* \) given \( y \). This prediction can be viewed as analogous to the point predictions given by the SVM. However, here in GPR each prediction is accompanied by some degree of estimated variance,

\[
\text{var}(y_*) = K_{**} - K_{*}K^{-1}K_{*}^T
\]

The degree of this variance translates to the confidence in the prediction.

In the GPC problem a sigmoidal response function is used to constrain continuous regression predictions to a particular set of target classes. Effectively the response function is used to map the unconstrained latent function provided by GPR to a particular class label. Several response functions can be used to this purpose, here the cumulative Gaussian density or probit likelihood was used,

\[
\Phi(x) = \frac{1}{2} \text{erf} \left( \frac{x}{\sqrt{2}} \right) + \frac{1}{2}
\]

For binary classification the likelihood for a single data point can be given as

\[
p(y_i | f_i) = \Phi(y_i, f_i)
\]

which enables the posterior distribution over the latent function to be written as:

\[
p(f|D, \theta) \propto \frac{N(f|0, K)}{p(D|\theta)} \prod_{i=1}^{m} \Phi(y_i, f_i)
\]
where $K$ is the covariance function (2.23). Latent function values for training points are stored by $f=[f_1, ..., f_m]^T$. $N(f|0, K)$ describes the prior over the latent function. The likelihood over training samples is factorized, as the class labels are independent given the latent function.

For GP classification; exact inference of the posterior distribution is analytically intractable and is therefore approximated using a Gaussian, $q(f|D, \theta) = N(f|m, \Sigma)$, which allows for calculation of the approximate posterior for a test case $q(f^*|D, \theta, x^*) = N(f^*|\mu^*, \sigma^*)$. Several approximation methods are available for this purpose. Here we employed an expectation propagation algorithm (Rasmussen and Williams, 2006), as it has been shown to have favorable results compared to other approximation methods (Kuss and Rasmussen, 2005, Nickisch and Rasmussen, 2008). These GPC based predictions are derived from integrating over the entire distribution for the latent function at a point for test case $x^*$ and therefore differ from the point based predictions provided by the SVM. See Nickisch and Rasmussen (2008) for further details.

In contrast to the SVM which seeks to optimize a margin between classes, ‘training’ of a GP models refers to finding the best functional form and hyper-parameters for the covariance function. Within this thesis this was achieved through maximizing the logarithm of the marginal likelihood. For GP regression problems this can be computed as,

$$
\ln p(y|X, \theta) = \frac{1}{2} y^T C_\theta^{-1} y - \frac{1}{2} \ln |C_\theta| - \frac{n}{2} \ln 2\pi
$$

(2.28)

where $C_\theta$ is the evaluation of $C$ given hyper-parameters $\theta$. For classification the marginal likelihood is again approximated using EP. Here (2.28) allows the user to select the hyper-parameters among competing models for which the marginal likelihood is optimal. The marginal likelihood calculates the total probability of the data given the model hyper-parameter and represents a trade off between optimal fit and model complexity and thus provides a buffer against over fitting.
2.7.3 Model Performance and Validation for Multivariate Pattern Classification

In supervised classification problems MVPC models learn to classify a group of training examples based on known class labels. These trained models are then applied to independent test examples for which class predictions are made. In order to accurately assess a model’s predictive performance it is important that test cases are not used in the training of a model. Ideally model performance is evaluated on completely independent test data. However, independent data is often not available for a variety of reasons. In neuroimaging this is commonly due to the expense of scanning individual subjects or challenges in recruiting large number of patients from a particular disease cohort.

When independent test data is not available cross validation schemes may be used in order to estimate model performance. In $k$-fold cross validation, $k$ number of cases are left out of model training, predictions are then made for the left out cases which act as the test data. The cross validation loop is then repeated $n$ number of times such that all cases in turn are left out from the training phase and are used as test cases for the model. When $k=1$ this scheme is known as leave one out (loo) cross validation. By comparing the predictive class labels returned by either independent validation or cross validation to the known labels it is possible to estimate a MVPC model’s accuracy. Leave one out cross validation is an effective validation method for minimizing testing bias as this framework allows almost the entire training sample to be used to establish the model for each testing case. However, given loo methods test only a single case each fold this minimization of testing bias comes at the cost of increased variance in performance estimates. Due to the relatively small numbers of training samples within this thesis, loo cross validation was used throughout with the knowledge that this may increase variance in our model performance estimates.

Model accuracy is defined as the percentage of test cases that are correctly classified. In addition to accuracy, models can be assessed by their sensitivity and specificity,

\[
sensitivity = \frac{TP}{TP+FN} \\
specificity = \frac{TN}{TN+FP}
\]
here \( TP \) represents the number of true positive cases (e.g. patients classified as patients) and \( TN \) represents the number of true negatives (e.g. controls classified as controls). Conversely \( FP \) represents the number false positive cases (e.g. controls classified as patients) and \( FN \) represents the number of false negative cases (e.g. patients classified as controls). For some instances in the clinical setting a model’s sensitivity to catch a potential positive case may be valued over the possibility of making a false positive diagnosis. In other situations a model’s specificity for making an accurate diagnosis may be viewed as more important.

Estimates of a model’s accuracy (or sensitivity and specificity) in themselves are not sufficient to tell if a model consistently performs above chance level. In order to test a MVPC model’s significance permutation testing can be performed. Here class labels are randomly permuted prior to model training and testing. This permutation is then repeated \( n \) number of time. Typically, \( n \) equals some order of thousands of permutation folds (e.g. \( n=1000, 10000 \), etc.). Across all folds the number of instances that the permuted model accuracy exceeds the original estimated accuracy is counted and then divided by the total number of permutations to give a \( p \) value for the classification model. As is convention model’s with a \( p \)-value of <0.05 are seen as statistically significant.

### 2.7.4 Feature Extraction and Selection

In MVPC the selection of data that is informative of the class labels often has more impact on model performance than the selection of one particular MVPC algorithm over another (e.g. SVM, GPC, etc.). In pattern classification (or regression) a particular characteristic of the data used to train a classification model is known as a feature or collectively as a feature set. Feature extraction is the process of extracting (i.e. calculating) a particular feature from a data set. For example, in this thesis feature extraction is the process of calculating surfaced based cortical measures from structural MRI data resulting in a vertex based feature set.

Feature sets based on MRI data often contain several thousand features per example. Therefore, in order to increase the amplitude of signal with regards to the class label it
is often desired, although not necessary, to remove uninformative features. Broadly speaking feature selection can be performed in two ways, 1) through prior knowledge and expertise (e.g. selection of features from a region of interest known to be involved in a condition) or 2) through an automated feature selection algorithm (Mwangi et al., 2014). Within parts of this thesis automated feature selection is performed through the application of a GLM to identify brain regions associated with a mean difference between classes. Data from vertices comprising these brain regions were then used for subsequent model training and testing.

It is important to note that automated feature selection introduces a strong possibility of over fitting a predictive model to the training set. Thus, it is important that the parameters used to implement automated feature selection are not selected arbitrarily but are tuned using a nested cross validation procedure (Figure 2.10). This avoids bias if the features are selected only using the training data.

![Figure 2.10 Nested Parameter Tuning via Cross Validation](image)

Figure 2.10 Nested Parameter Tuning via Cross Validation: This model depicts one fold of leave one out cross validation with a nested leave one out cross validation loop to tune model parameter values. A) Leave one out validation begins by excluding one case that will act as testing data (blue square) from the training set (red and blue squares in shaded box). B) From the training data set nested cross validation begins by C) performing leave one out cross validation for each parameter value to be tuned. The parameter value that returns the highest model accuracy (here parameter value 2) is then used to train and test the actual model. This nested scheme is then repeated across all cross validation folds.
Many ways of performing feature selection are available including both univariate (e.g. GLM) and multivariate approaches. When using a GLM to perform feature selection the uncorrected $p$-value threshold used to identify regions of interest represents one parameter that is tuned. Thus, for each cross validation fold a range of $p$-value thresholds are tested via nested cross validation. The $p$-value that returns the highest model accuracy is then used in training of the model for that cross validation fold. For this thesis, a GLM feature selection approach was used however such univariate approaches have the limitation of not taking into account relationships between data points. Thus, potentially informative data points may not be selected as they are tested for significance in isolation. Several other multivariate feature selection methods (e.g. recursive feature elimination, lasso) which may provide better results are available (for review see Mwangi et al. (2014)). However, due to time constraints in was beyond the scope of this project to extensively test all these methods.

### 2.7.5 Model Visualization

The spatial maps of MVPC models are inherently different than those provided by mass univariate methods such as the GLM. The GLM can be viewed as a forward model as it takes a known variable (e.g. class label) and highlights aspects of the data (e.g. brain regions) associated with the variable. This differs from backward models such as MVPCs that extract information from the data as it relates to a variable (Haufe et al., 2014).

Commonly MVPC models are visualized through spatial mapping of the model’s weight vector. However, interpretation of these so call “discriminative maps” is made difficult by the fact that you cannot infer that values within $w$ are directly related to the signal of interest (e.g. class label). For example, large weights may be associated with noise and small values the signal of interest. Despite limitations on interpretability several studies still report multivariate discriminative maps along with MVPC models. Thresholding discriminative maps based on a percentage of the maximum weight value has been proposed as one way of reducing noise and clarifying interpretation (Mourão-Miranda et al., 2005, Mourao-Miranda et al., 2006).
Other multivariate mapping methods have recently been proposed to calculate forward maps more in line with those generate by the GLM (Hauße et al., 2014).

An alternative mapping method, known as ‘predictive mapping’ involves taking the dot product of each individual example ($x_i$) with the corresponding weight vector ($w$), so that $m = (x_i) \cdot w$ (Marquand et al., 2014). Predictive mapping has two advantages: 1) it shows the total contribution to the prediction for each test point, i.e. the weights derived from the training data and from the test data point, and therefore 2) it can be used to determine the spatial distribution of informative regions for cases that were not learned as part of the primary decision problem. In other words, predictive mapping allows us to infer which aspects within a group of test cases underlie predictions derived from the classifier.
2.8 References


90


Chapter 3: Grey to White Matter Percent Contrast as an in vivo Neuroimaging Feature in Autism Spectrum Disorder

Autism spectrum disorder (ASD) is a condition that manifests during early development that is accompanied by differences in brain anatomy and connectivity (Lange et al., 2015, Ecker et al., 2015, Amaral et al., 2008). Several prior magnetic resonance imaging (MRI) studies of ASD have reported atypical measures of cortical anatomy such as folding, thickness, and surface area (Nordahl et al., 2007, Hyde et al., 2010, Ecker et al., 2013, Schaer et al., 2013). These *in vivo* findings are supported by postmortem reports of atypical cortical structure such as increased numbers of neuron cells, cortical minicolumns and instances of folding abnormalities in ASD (Casanova et al., 2006, Courchesne et al., 2011, Kemper, 1988).

This chapter details a novel application of measures of tissue contrast in an attempt to quantify a specific aspect of neuropathology in ASD, namely poor definition of the grey-white matter boundary (Avino and Hutsler, 2010). This study was recently published in *Cerebral Cortex* (Andrews et al., 2017) and is included in its final published format in the pages that follow. Supplementary materials for this chapter can be found in Appendix II.
In Vivo Evidence of Reduced Integrity of the Gray–White Matter Boundary in Autism Spectrum Disorder

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Declan G.M. Murphy and Christine Ecker contributed equally to the manuscript.

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Abstract

Atypical cortical organization and reduced integrity of the gray-white matter boundary have been reported by post-mortem studies in individuals with autism spectrum disorder (ASD). However, there are no in vivo studies that examine these particular features of cortical organization in ASD. Hence, we used structural magnetic resonance imaging to examine differences in tissue contrast between gray and white matter in 98 adults with ASD and 98 typically developing controls, to test the hypothesis that individuals with ASD have significantly reduced tissue contrast. More specifically, we examined contrast as a percentage between gray and white matter tissue signal intensities (GWPC) sampled at the gray-white matter boundary, and across different cortical layers. We found that individuals with ASD had significantly reduced GWPC in several clusters throughout the cortex (Cluster, P < 0.05). As expected, these reductions were greatest when tissue intensities were sampled close to gray-white matter interface, which indicates a less distinct gray-white matter boundary in ASD. Our in vivo findings of reduced GWPC in ASD are therefore consistent with prior post-mortem findings of a less well-defined gray-white matter boundary in ASD. Taken together, these results indicate that GWPC might be utilised as an in vivo proxy measure of atypical cortical microstructural organisation in future studies.

Key words: ASD, FreeSurfer, imaging, lamination, MRI

Introduction

Autism spectrum disorder (ASD) is a lifelong neurodevelopmental condition characterised by impaired social communication, deficits in social reciprocity, and repetitive and stereotyped behaviors and interests (Wing 1997). These core symptoms typically manifest from early childhood, and are accompanied by developmental differences in brain anatomy and connectivity (for review, see Amaral et al. 2008; Döber et al. 2013; Lange et al. 2013). For example, prior studies of ASD reported atypical measures of cortical anatomy such as thickness, surface area (Hordijk et al. 2007; Hyde et al. 2010; Schaefer et al. 2013; Döber et al. 2013b) as well as intra-cortical connectivity (Döber et al. 2013b). However, the causes of these cortical abnormalities in people with ASD are unknown.

There is some evidence to suggest that the cortical differences accompanying ASD may result from atypical neuronal proliferation, migration, and maturation (Pinto et al. 2014). For example, some genetic variants associated with ASD encode for genes that regulate these neurodevelopmental processes (Duguet et al. 2013). It has been suggested that these variations may explain post-mortem findings such as irregular cortical lamination, the presence of super-numerous neurons in some layers of the cortex, and poor differentiation of the gray-white matter boundary (for review, see Caspers et al. 2014). For example, histological samples from the superior temporal gyrus (approximate Brodmann area (BA) 22), dorsolateral prefrontal cortex (BA10) and dorsal parietal cortex (BA7) have shown the gray-white matter boundary to be less distinct in ASD as compared with typically developing TD controls (Astley and Autism 2010). Thus, there is increasing post-mortem evidence for abnormal cell patterning within the gray-white matter boundary in ASD. However, to date no a priori evidence is available for the integrity of the gray-white matter boundary in ASD in vivo across the whole brain.

Current in vivo neuroimaging methods for investigating cortical abnormalities in ASD focus on morphometric features such as cortical thickness (CT), that is, the closest distance from the gray-white matter boundary to the gray-cerebral spinal fluid (CSF) boundary (Polich and Dale 2000). Differences in CT have been reported in children, adolescents, and adults with ASD, and include regional increases and decreases that may mediate some of the behavioral deficits typically observed in the disorder (Staskus et al. 2006; Hyde et al. 2010; Döber et al. 2013b). However, measures of CT rely on the accurate delineation of gray and white matter and therefore may be confounded by intrinsic histological abnormalities at the gray-white matter boundary in ASD (Astley and Autism 2010).

Hence, we investigated between-group differences related to cortical lamination in both adult males and females with ASD, and matched TD controls, using a whole brain quantitative approach that estimated integrity of the gray-white matter boundary. Namely we examined the percent contrast of gray-to-white matter signal intensities (GWPC), sampled across different cortical layers in a continuous fashion. Here, the GWPC calculation we employed in the current manuscript is comparable to the gray-white contrast ratio as originally reported by Salat et al. (2009). We hypothesized the gray-white matter boundary to be less defined and therefore GWPC to differ significantly in individuals with ASD.

Materials and Methods

Participants

Overall, 98 right-handed adults with ASD (49 males and 49 females) and 98 age, sex, and IQ matched TD controls (51 males and 47 females) aged 18–43 years were recruited by advertisement and assessed at the Institute of Psychiatry, Psychology and Neuroscience (IOPPN), London, and the Autism Research Centre, Cambridge. Approximately equal ratios of cases to controls, and males to females, were recruited within sites (Table 1). Inclusion criteria included a history of major psychiatric disorder (e.g. psychosis), head injury, genetic disorder associated with autism (e.g. fragile-X syndrome, tuberous sclerosis), or any other medical condition affecting brain function (e.g. epilepsy), or any participants taking antipsychotic medication, mood stabilizers, or benzodiazepines.

ASD diagnosis was made by a consultant psychiatrist using ICD-10 research diagnostic criteria and confirmed using the ADI-R (Lord et al. 1994). ASDs were completed for 94 individuals with ASD (49 males and 45 females). Ninety-three (49 males and 44 females) reached algorithm cut-offs for autism in all domains of the ADI-R (social, communication, and restricted/interested), although failure to reach cut-off in one domain by one point was permitted. The ADI-R rather than ADOS (Lord et al. 2000) was employed as inclusion criteria to ensure that all participants
Table 1 Participant demographics

<table>
<thead>
<tr>
<th></th>
<th>ASD (n = 98)</th>
<th>Control (n = 98)</th>
</tr>
</thead>
<tbody>
<tr>
<td>London</td>
<td>n = 45 (24M, 21F)</td>
<td>n = 44 (25M, 19F)</td>
</tr>
<tr>
<td>Cambridge</td>
<td>n = 53 (25M, 28F)</td>
<td>n = 54 (29M, 25F)</td>
</tr>
</tbody>
</table>

Age, years

<table>
<thead>
<tr>
<th></th>
<th>ASD</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median</td>
<td>26±7</td>
<td>27±8</td>
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</tbody>
</table>

Full scale IQ, WASI

<table>
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<tbody>
<tr>
<td>Median</td>
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<td>112±13</td>
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ADOS (social)¹

<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td>Median</td>
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<td>19±6</td>
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ADOS (communication)⁴

<table>
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</thead>
<tbody>
<tr>
<td>Median</td>
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<td>13±4</td>
</tr>
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</table>

ADOS (repetitive behavior)⁴

<table>
<thead>
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</tr>
</thead>
<tbody>
<tr>
<td>Median</td>
<td>5±2</td>
<td>5±2</td>
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</table>

ADOS social + communication⁴

<table>
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<th></th>
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<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median</td>
<td>9±3</td>
<td>8±3</td>
</tr>
</tbody>
</table>

Data expressed as mean ± standard deviation (range). There were no significant differences between group means in age or IQ; P > 0.05 (2-tailed). All participants were diagnosed using IC-10 criteria.

¹The Autism Diagnostic Interview Revised (ADI-R) was used to confirm ASD diagnosis. ADOS scores were unavailable for 4 participants. Each of these cases reached the Autism Diagnostic Observation Schedule (ADOS) cut-off for autism spectrum, but all other participants for the ADOS was not used as diagnostic criteria.

²T1-contrasts did not undergo ADOS or ADOS assessments.

with ASD met the criteria for childhood autism. We were unable to complete ADI-Rs for 4 females with ASD as their parents/caregivers were not available. However, all 4 reached cut-off for “autism spectrum” on the ADOS (communication, social) diagnostic algorithm. In all other participants, ADOS scores were used to measure current symptoms and not as inclusion criterion. One ASD female scored one point below cut-off for autism on the communication and repetitive behavior domains of the ADI-R but met TCD-10 criteria for ASD and scored above cut-off for “autism” on the ADOS. Overall, intellectual ability was assessed using the Wechsler Abbreviated Scale of Intelligence (WASI; Wechsler 1999). All participants had a full-scale IQ greater than 80 and gave informed written consent in accordance with ethics approval by the National Research Ethics Committee, Sheffield, UK.

Structural MRI Data Acquisition

Scanning was performed at the MPII, London, and Addenbrooke’s Hospital, Cambridge, using a 3 T GE Signa System (General Electric). A specialized acquisition protocol using quantitative T1-mapping was used to ensure standardization of structural magnetic resonance imaging (MRI) scans across scanner platforms. This protocol has previously been validated and extensively described elsewhere (Deoni et al. 2006; Eamer et al. 2012), resulting in high-resolution structural T1-weighted inversion-recovery images, with: 512×512×1 mm resolution, a 256×256×178 matrix, TR = 1800 ms, TE = 50 ms, TA = 20 s, and FOV = 5 cm.

Cortical Reconstruction Using FreeSurfer

Previous histological studies have largely relied upon manual identification to define the boundary between gray and white matter. For example, Avila and Huttenlocher (2010) used a sigmoid function to quantify the distinctiveness of the transition between gray and white matter in Nissl-stained histological images. In the current study, however, we employed an automated analytical pipeline using FreeSurfer v5.3.0 software (http://surfer.nmr.mgh.harvard.edu) to identify the gray-white matter boundary by deriving models of the cortical surface for each T1-weighted image. These well validated and fully automated procedures have been described elsewhere (Dale et al. 1999; Fischl et al. 1999; Fischl and Dale 2000; Sämann et al. 2004; Joutzich et al. 2006). In brief, a single-sliced white-matter volume was generated for each hemisphere after intensity normalization, extra-cerebral tissue was cropped, and image segmentation performed using a connected-components algorithm. A triangular tessellated surface was then generated for each white-matter volume. Deformation of this tessellated white matter surface resulted in a cortical mesh for the surfaces that define the boundary between grey and white matter (i.e. white matter surface), and gray matter and CSF (i.e. pial surface). This surface deformation is the result of the minimization of an energy functional that utilizes intensity gradients in order to place these surfaces where the greatest shift in intensity defines the transition between tissue classes (Dale and Sereno 1999; Supplementary Material). The use of intensity gradients across tissue classes assumes that boundary placement is not reliant solely on absolute signal intensity and allows for subcortical resolution in the placement of these boundary surfaces (Dale and Sereno 1999; Dale et al. 1999; Fischl and Dale 2000). These automated methods have previously been validated against histological analysis and have shown a high degree of accuracy in placing the gray-white matter boundary (Sämann et al. 2004). The resulting surface models were visually inspected for reconstruction errors. Participant’s surface reconstructions with visible inaccuracies were excluded and are not described in this study. Dropout rates due to surface reconstruction errors were equal between groups and represented ~10% of the total sample.

Gray-to-White Matter Percent Contrast (GWMPC) and Gray-Matter Signal Intensity Measures

Gray-matter tissue intensities (GMI) were sampled continuously across different cortical layers from the gray-white matter boundary (i.e. white matter surface) to the pial surface. These signal intensities were sampled at different percentile fractions of the total orthogonal distance projected from the white matter to pial surfaces (i.e. projection fraction). Starting at the white matter surface, sampling continued at projection fraction intervals of 10% up to 60% of the distance from the white matter to the pial surface, thus yielding a set of 6 GMI measures (i.e. from 10% to 60%, Fig. 1). The outer 40% (i.e. 70-100%) of the cortical sheet was not sampled in order to assure that sampling was performed within the cortical gray matter, and not confounded by voxels composed of CSF. White matter signal intensity (WMI) was measured at 1.0 mm into the white matter from the white matter surface (Fig. 1). Previously reported measures of tissue contrast have used a ratio calculation (i.e. GMI/WMI; Salat et al. 2000), where larger values indicate a reduced contrast. Here, however, we utilized the formula provided by FreeSurfer to calculate tissue contrast as the percentage of GMI at projection fraction (f) to WMI at each cerebral vertex (i),

$$GWMPC_i = \frac{100 \times (WMI_{1.0mm} - GMI_i)}{WMI_{1.0mm} - GMI_{0.0}}$$

Thus, by definition, a decrease in GWMPC is commensurate with a decrease in contrast between the GM measured at projection fraction i, and the WMI measured at 1.0 mm subjacent to the white matter surface. We also examined the tissue contrast when sampling GMI at the gray-white matter boundary (i.e. at the white matter surface, projection fraction = 0%). The resulting GWMPC, GMI, and WMI measures were subsequently smoothed using a 10-mm full-width at half maximum (FWHM) surface-based...
Results

Participant Demographics and Global Brain Measures

There were no significant differences between individuals (males and females) with ASD and TD controls in age (t(194) = -0.93, P = 0.36, full-scale IQ (t(194) = -1.72, P = 0.09), or total GM volume (t(194) = -0.93, P = 0.178). There were also no significant differences between males and females in age (t(194) = -0.93, P = 0.36) or full-scale IQ (t(194) = -1.87, P = 0.068). As expected, total gray matter volume in males was significantly larger than in females (t(194) = 3.51, P < 0.001). However, there were no significant differences in any of these measures between males with ASD and male controls, or females with ASD and female controls (P > 0.05, 2-tailed).

Between-Group Difference in GWPC Across the Cortex

We initially examined vertex-wise between-group differences in GWPC at different projection fractions into the cortical sheet. At all sampling depths, we found that individuals with ASD had a significantly decreased GWPC in several clusters across the cortex, which is consistent with a reduced tissue contrast between gray and white matter (Fig. 3). In accordance with our hypothesis, the reductions in GWPC were most extensive when GM was sampled at gray-white matter boundary (i.e. the white matter surface, projection fraction = 0%), and gradually decreased in both statistical effect and spatial extent with increasing projection fractions into cortex and away from the gray-white matter boundary. Regions where ASD individuals had reduced GWPC as compared with TD controls included the 1) bilateral posterior-cingulate (BA 23/30), medial frontal (BA 10) parsorbitofrontal/orbitofrontal (BA 14/32) and the inferior and superior...
temporal cortices (BA20/21/22); 2) left orbitofrontal cortex (BA11/12) and temporo-parietal junction (BA39/40); and (3) right dorsolateral prefrontal cortex (BA9/46). Statistical details for all clusters are listed in Table 2. There were no brain regions where individuals with ASD had a significantly increased GWPC relative to controls. The pattern of reduced GWPC among individuals with ASD remained significant when total brain volume or mean CT were included as covariates. Furthermore, there was minimal spatial overlap between the pattern of differences in GWPC and CT (see Supplementary Fig. 1 and Table 1).

**Between-Group Differences in Gray and White Matter Tissue Intensities**

To identify whether the observed differences in GWPC were driven by differences in gray or white matter, or a combination of both, we subsequently examined between group differences in
Table 2: Changes of significant reductions in gray white matter percent contrast and increases in gray matter intensity in ASD

<table>
<thead>
<tr>
<th>Measure</th>
<th>Cluster</th>
<th>Region labels</th>
<th>Hemispheric BA (max)</th>
<th>Vertex</th>
<th>Talairach</th>
<th>tmax</th>
<th>P(cmax)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GWPC</td>
<td>1</td>
<td>Superior temporal gyrus, insula, lateral orbital frontal cortex, pars opercularis, postcentral gyrus, precuneus gyrus</td>
<td>1</td>
<td>21</td>
<td>10194</td>
<td>47</td>
<td>-4</td>
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<tr>
<td></td>
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<td>Frontal-opercular cortex, inferior occipital cortex, lingual gyrus, precuneus cortex</td>
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<td>21</td>
<td>5700</td>
<td>7</td>
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</tr>
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<td>Middle temporal gyrus, banks superior temporal sulcus, anterior temporal gyrus, superior temporal gyrus</td>
<td>R</td>
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<td>4064</td>
<td>54</td>
<td>-11</td>
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<tr>
<td></td>
<td>4</td>
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<td>L</td>
<td>21</td>
<td>4837</td>
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<td>-30</td>
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<td>Insula, lateral orbital frontal cortex, pars opercularis, postcentral gyrus, precuneus gyrus</td>
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<td>25</td>
<td>-53</td>
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<td>Medial orbital frontal cortex, rostral anterior cingulate cortex, superior frontal gyrus</td>
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<td>3320</td>
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<td>L</td>
<td>37</td>
<td>3483</td>
<td>-36</td>
<td>-43</td>
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<tr>
<td></td>
<td>9</td>
<td>Frontal-opercular cortex, inferior occipital cortex, lingual gyrus, precuneus cortex</td>
<td>L</td>
<td>29</td>
<td>3432</td>
<td>-8</td>
<td>-56</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>Supramarginal gyrus</td>
<td>L</td>
<td>40</td>
<td>2466</td>
<td>-56</td>
<td>-33</td>
</tr>
<tr>
<td>CME</td>
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<td>R</td>
<td>38</td>
<td>17958</td>
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<td></td>
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<td>Superior temporal gyrus, banks superior temporal sulcus, inferior parietal cortex, inferior temporal gyrus, middle temporal gyrus</td>
<td>L</td>
<td>21</td>
<td>10279</td>
<td>-51</td>
<td>-26</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>Parietal gyrus, inferior temporal gyrus, inferior occipitocingulate cortex, lingual gyrus, precuneus cortex</td>
<td>L</td>
<td>52</td>
<td>6295</td>
<td>-64</td>
<td>-40</td>
</tr>
</tbody>
</table>

Note: Changes of significant reductions in GWPC and increases in gray matter intensity (CME) in ASD. BA, left (L), right (R). ‘Vertices’ indicates the number of vertices within the cluster. tmax represents the maximum t-value within the cluster located at the x y z Talairach coordinates listed. P(cmax) is the cluster corrected P value.

Both GMI and WME. Individuals with ASD had significantly increased GMI across all 6 different GM sampling depths relative to controls in regions where we also observed decreases in GWPC (Fig. 3). These included: 1) the bilateral anterior temporal lobes (BA38/39) and the left middle temporal gyrus (BA21), 2) the right temporo-parietal junction (BA39/40), and 3) the bilateral fusiform and extrastriate cortex (BA17). Statistical details for these clusters are listed in Table 2. We did not observe any significant between-group differences in GMI at the gray-white matter boundary (i.e. the white matter surface), or in WME at 1.0 mm within the white matter (Fig. 3). There were no brain regions where individuals with ASD had significantly decreased GMI relative to controls. Hence, GWPC reductions in ASD were driven predominantly by increased (i.e. higher) mean intensities within the cortical gray matter.

Main Effects of Sex and Group by Sex Interactions

Last, we investigated whether biological sex significantly modulated differences in GWPC in ASD by examining group-by-sex interactions. Overall, regardless of diagnosis, males had a significantly greater GWPC than females (Supplementary Fig. 2). This occurred across all sampling depths, and was predominantly in frontal-parietal regions of the left hemisphere, and in bilateral inferior temporal regions (see Supplementary Table 2 for statistical details of these clusters). However, there were no brain regions where we observed significant group by sex interactions for GWPC. Thus, while males tended to have a significant increase in contrast between gray and white matter these intensities the reductions in GWPC that we observed in the brain in individuals with ASD were not explained by biological sex.
**Discussion**

Our aim was to determine if previous postmortem reports of poor definition of the gray-white matter boundary in ASD could be detected using a whole brain in vivo MRI approach. As hypothesized, we determined that individuals with ASD had a significantly less well-defined tissue contrast (i.e. GWPC) between gray and white matter at (and around) the gray-white matter boundary. The affected brain regions included the superior temporal gyrus (BA22), the dorsolateral frontal lobe...
(BAM) and the dorsal parietal lobe (DPL) where histological abnormalities in the transition from gray to white matter have also been reported (Avino and Juskevicius 2010). The concordance between the regional pattern and direction of the GWPC differences in our sample with previous histological investigations in postmortem brain tissue supports the biological plausibility of our results. Thus, our findings agree with previous postmortem histological studies and indicate that tissue contrast across the gray-white matter interface may serve as a potential in vivo proxy measure for structural organization of the cortical sheet in ASD.

Postmortem studies reported abnormalities in the cortical microstructure of individuals with ASD. For example, the boundary between cortical layer VI and underlying white matter has been shown to be significantly less well defined due to increased dispersion of neuronal cells across this interface (Avino and Juskevicius 2010). It has been suggested that this may be caused by the presence of supernumerary neurons beneath the cortical plate that arise from disrupted migratory processes or improper resolution of the cortical subplate (Chun and Shatz 1989; Kemper 2012; Juskevicius and Avino 2015). The cortical subplate is a transient neurodevelopmental layer that is instrumental in establishing proper cortical connectivity. Specifically, subplate neurons pioneer the corticothalamic axon pathway, serving as a "signpost" for cortical axons, drive endogenous oscillatory activity in the cortex, and act as a transient synaptic hub for thalamocortical axons before they decussate in the corpus callosum (Albanese et al. 1996; Gosh et al. 1992; McGonnell et al. 1994; Lukehart et al. 2009). The maximal volume of the subplate is reached around 30 gestational weeks in the human coinciding with the growth of long range cortico-cortical projections (Van De Grind et al. 2016). After their early neurodevelopmental role is complete, a large number of these subplate neurons undergo apoptosis. However, a small percentage of these neurons persist and retain their connections with the overlying cortical plate acting as modulators of cortical afferents (Chun and Shatz 1989; Dupont et al. 2006; Koevendi et al. 2011).

Therefore, perturbations to early subplate development may disrupt the establishment of structural and functional brain connectivity, which is abnormal in individuals with ASD (Belmonte et al. 2004; Jiao et al. 2004; Courchesne and Prizant 2005; Balkind et al. 2015). In addition, the abnormal persistence of these neurons after the large wave of programmed cell death could cause disruptions to cortical communication through their modulatory role of the overlying cortex. In this way, the abnormal persistence of subplate neurons into adulthood has been demonstrated in schizophrenia and autism disorder and is hypothesized to contribute to the pathophysiology of these conditions (Dawood and Harrison 2003, 2005; Andrews et al. 2005; Hideohara et al. 2005; Koevendi et al. 2011; Yang et al., 2011). Furthermore, a recent genetic study reported a set of subplate-specific genes that are associated with ASD (Dawood 2011; Riedel-Boschdierung et al. 2013). Thus, there is converging evidence to suggest that neurons of the cortical subplate contribute to the aberrant, neuropathology of ASD and that atypical cortical organization, particularly around the gray-white matter boundary, may be a defining characteristic of the condition. However, this has never previously been examined in vivo.

Thus, in this in vivo study, we sought to examine differences in cortical lamination and gray-white matter boundary integrity in ASD. To achieve this, we measured contrasts between gray and white matter tissue intensities (GWPC, Salt 2005). These MRI measures were taken as the interface of gray and white matter and across cortical layers at 6 different depths into the cortical sheet from the gray-white matter boundaries (i.e., white matter surface). In our ASD cases, many regions with reduced GWPC also showed significantly increased GM but no differences in WM as compared with TD controls. This suggests (in agreement with prior in vivo work by our group; Didden et al. 2016) that ASD may be primarily associated with disruptions to cortical gray matter as opposed to white matter. This increased GM in ASD may result from atypical myelination (Didden et al. 2006) and/or atypical cytoarchitectural organization such as greater numbers of more densely packed cortical minicolumns (Casanova et al. 2006) and reductions in grey layer amplitudes in these structures (Casanova et al. 2003). The regional specificity of our findings of decreased tissue contrast may be related to the differential expansion of the subplate between cortical areas. Evolutionarily, the size and complexity of the subplate is more prominent in humans as it accommodates the increased connectivity with cortical and subcortical areas relative to non-human primates and rodents (Kozlovic and Kalia, 1990; Juskevicius et al. 2013). Within humans, the subplate zone is larger in cortical association areas as a consequence of the increased number of axons invading these regions. These incoming axons displace subplate neurons deeper into the white matter, which occurs to a greater degree in these association areas (Dusore et al. 2016). Applicability of the gray-white matter interface may therefore impact on MRI intensity values, and may explain the regional specificity observed in our patterns of results. Moreover, the regional pattern of GWPC seems to be linked to the functional deficits that are characteristic for ASD. For example, we observed deficits in GWPC in several regions mediating social processing and wider socio-cognitive functioning, including the insula, fusiform gyrus, cingulate cortex, middle temporal gyrus, superior temporal sulcus, and prefrontal cortical regions (seeJust et al. 2012 for review). Thus, while future studies are required to establish the functional relevance of our results directly, it is likely that atypical GWPC contributes to the cluster of clinical symptoms typically observed in ASD.

Findings from this and other studies detailing poor delineation of the gray-white boundary in ASD may be taken by some to call into question the accuracy of in vivo MRI measures such as CT that rely on the placement of a discrete boundary between gray and white matter. However, the spatially distributed patterns of group differences in CT we detected did not significantly overlap with the patterns of differences in GWPC (see Supplementary Fig. 1). Also, including individual's global mean CT as a covariate did not significantly alter the pattern of differences in GWPC. Therefore, while we were able to detect subtle differences in tissue contrast in ASD, at the level of spatial resolution neuroimaging techniques currently offer, these do not appear to be large enough to significantly affect estimates of CT within our sample of adults with ASD. This finding is also in agreement with a recent twin study showing that while both GWPC and CT are highly heritable, they have little shared genetic variance (Pessin et al. 2012). Taken together, these findings suggest that GWPC characterizes additional cortical structural properties that are distinct from CT. Nevertheless, inter-individual differences in the ability to delineate the gray-white matter boundary should be considered in the future when interpreting neuroanatomical features that are based on clearly delineating gray and white matter.

Our study is not without limitations. For instance, we examined neuroanatomical differences associated with ASD in adulthood. This, and the cross-sectional nature of our study,
Inherently limits our ability to draw conclusions on the etiological and neurodevelopmental basis of the atypical neural structure we observed. However, within our sample, all but 4 females with ASD met ADI-R criteria for childhood autism. It is therefore likely that the observed pattern of neuroanatomical differences in GWPC may have evolved as a consequence of meeting ASD criteria during early childhood and is therefore causally related to the condition. Further longitudinal studies will, however, be required to disentangle GWPC differences associated with primary neuroanatomy from atypical neurodevelopmental trajectories or secondary compensatory mechanisms. Recent work has quantified the volume of transient neurodevelopmental nodules in the postmortem human fetal brain using MRI as they relate to major neurogenic events (Vasung et al. 2016). Such information provides a reference for studying early prenatal deviations from TD brain growth and could be used in the future to inform in vivo imaging. We are further limited by the current resolution of structural MRI images (1 mm isotropic voxels). At this resolution, it is not possible for us to distinguish between different aspects of cortical cytoarchitecture or accurately delineate particular layers of the cortical sheet as defined by histological staining. Rather, our sampling approach was based on the geometric criteria of projection fraction percentages into the cortical sheet from the white matter surface (Salar et al. 2009). Furthermore, additional research will be required to elucidate the functional relationship between deficits in GWPC and autistic symptoms and traits.

Taken together, our findings suggest that measures of GWPC sampled across cortical layers may serve as an in vivo proxy measure for irregular microstructural organization of the cortex in ASD (and other disorders). Such novel in vivo measures that are indicative of atypical cortical organization might in the future be used to stratify the condition, and/or to examine the neuroanatomy of ASD in particular genetic subgroups known to be linked to specific neurodevelopmental deficits.

Supplementary Material
Supplementary material are available at Ceentral Core online.

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Chapter 4: Clinical Testing of Multivariate Pattern Classification Models for Autism Spectrum Disorder

4.1 Introduction

Until recently, the majority of neuroimaging studies in mental health research have used so-called mass-univariate methods which examine mean differences in brain measures between groups of individuals. However, these methods are not suited to make predictions for individual cases on the basis of patterns within the data (e.g. to predict group membership or clinical outcome), and thus offer limited translational value for clinical applications. Increasingly, “machine learning” methods such as multivariate pattern classification (MVPC) have been applied to clinical neuroimaging data to discriminate cases from controls. In brief, MVPC algorithms such as the Support Vector Machine (SVM) (Vapnik, 1995) or Gaussian Process Classifier (GPC) (Rasmussen and Williams, 2006) differentiate between two or more classes in the following way: 1) During a ‘training phase’ the model is initially established using well-defined samples to determine a mathematical criterion of a decision function that best distinguishes between the groups (e.g. ASD and controls). 2) During the ‘testing phase’, the decision function is applied to predict the class or label of a new test example (e.g. an unseen data point).

Several proof-of-concept studies having shown promise in classifying patients from controls across several psychiatric and neurodevelopmental conditions (Wolfer et al., 2015, Arbabshirani et al., 2016). Within autism spectrum disorder (ASD), volumetric (Ecker et al., 2010b), morphological (Jiao et al., 2010, Ecker et al., 2010a), diffusion (Ingalhalikar et al., 2011, Ingalhalikar et al., 2010), and functional (Just et al., 2014) MRI brain measures have all been applied to discriminate individuals with ASD from typically developing (TD) controls at accuracies significantly above chance within the research setting (for review see Chapter 1.4.1). However, MVPC model performance (e.g. overall classification accuracy) is most often estimated using cross-validation methods within the same sample. To date, only two studies have evaluated ASD classification models on independent data sets acquired in the ‘research setting’ (Sabuncu and Konukoglu, 2014, Shen et al., 2017). These studies have reported modest overall accuracy in identifying individuals with ASD (59-72%) when
compared to the literature as a whole, which includes reports of several model performances ranging from 80 to upwards of 100% accuracy.

To date no studies have tested classification models for ASD trained in the research environment on independent samples that are representative of the broader clinical population. This is important because several differences exist between research and clinical ASD cohorts. For example, individuals with ASD recruited within the research environment will generally be excluded if they have comorbid conditions, are taking certain medications, or fail to reach cut-offs on behavioural metrics such as the autism diagnostic interview (ADI) and/or autism diagnostic observation schedule (ADOS). In addition, low functioning ASD cases are often not included in imaging research. As such, many individuals who would qualify for a clinical ASD diagnosis would not meet inclusion criteria for research. As a result, the clinical phenotype between individuals with ASD in research and clinical samples is likely to differ to some degree.

Furthermore, proof of concept classification studies report model performance when distinguishing individuals with ASD from typically developing (TD) controls. However, individuals in clinical populations who do not qualify for an ASD diagnosis usually would not be considered TD. Rather individuals who are referred for specialist ASD diagnostic services and found not to have the condition often qualify for a diagnosis of a different but related disorder (e.g. obsessive-compulsive disorder, social anxiety disorder, learning disability etc.) or have a subclinical behavioral presentation that would exclude them from qualifying as a TD control. Taking the above into account, training sets on which classification models are established are not necessarily exchangeable with the testing population of interest (i.e. clinical population).

Thus, this study sought to (1) establish discriminative models for ASD within a research sample of TD controls and ASD adults, and then (2) test the efficacy of these models to identify ASD in an independent sample of individuals who had been referred to a specialist clinical service for ASD diagnosis. Two types of classification algorithms, i.e. the SVM and GPC were examined separately. The SVM and GPC were selected as they have both been successfully applied previously to neuroimaging
based classification problems (Ecker et al. 2010A, Ecker et al. 2010B, Marquand 2010). These methods were applied to several different morphometric measures including volumetric as well as geometric features, which have previously been shown to be atypical in individuals with ASD (Ecker et al., 2010a, Sabuncu and Konukoglu, 2014, Jiao et al., 2010, Andrews et al., 2017). Based on previous reports we hypothesize that measures of cortical morphometry will be able to significantly discriminate ASD individuals from controls at accuracies above chance level within the research setting. However, given the inherent differences between our research training and clinical testing samples and previous findings of classification models showing reduced performance in the range of 10-15% when applied to independent testing sets (Sabuncu and Konukoglu, 2014), we expected a reduction in overall accuracy when predicting ASD diagnosis within the clinical sample.

4.2 Methods

4.2.1 Research Group Participant Demographics
The research cohort included structural MRI data obtained in 27 male right-handed adults with ASD (Ages 18-38, 25±6.3yrs) and 27 matched TD developing controls (Ages 18-42, 28±6.1yrs), that were recruited by advertisement and assessed at the Institute of Psychiatry, Psychology and Neuroscience, London UK. Exclusion criteria included a history of major psychiatric disorder (e.g., psychosis), head injury, genetic disorder associated with autism (e.g., fragile-X syndrome, tuberous sclerosis), or any other medical condition affecting brain function (e.g., epilepsy). We excluded participants on antipsychotic medication, mood stabilizers, or benzodiazepines.

All participants with ASD within the training set were diagnosed according to ICD-10 research criteria. Diagnoses were confirmed using the Autism Diagnostic Interview (ADI-R; (Lord et al., 1994). All cases reached the diagnostic algorithm cutoffs for adults in the three domains of the ADI-R, although failure to reach cutoff in one domain by one point was permitted. Current symptoms were assessed using the Autism Diagnostic Observation Schedule (Lord et al., 2000) but were not used as inclusion criteria. Overall, intellectual ability was assessed using the Wechsler Abbreviated Scale of Intelligence (WASI) (Wechsler, 1999). All participants had a full-scale IQ score greater than 75. Group differences in age and full scale IQ were
assessed using a two-tailed independent samples t-test. All participants gave informed written consent in accordance with ethics approval by the National Research Ethics Committee, Suffolk, United Kingdom.

Table 4.1. Participant Demographics (Research Sample)

<table>
<thead>
<tr>
<th></th>
<th>ASD (n=27♂),</th>
<th>TD Controls (n=27♂)</th>
</tr>
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<tr>
<td>Age, years</td>
<td>25 ± 6.3 (18-38)</td>
<td>28 ± 6.1 (18-42)</td>
</tr>
<tr>
<td>Full-scale IQ, WASI</td>
<td>106 ± 13.0 (83 - 133)</td>
<td>110 ± 14.0 (77 - 130)</td>
</tr>
<tr>
<td>ADI-R social</td>
<td>17 ± 5.8 (9 - 28)</td>
<td>*</td>
</tr>
<tr>
<td>ADI-R communication</td>
<td>12 ± 12.4 (8 - 24)</td>
<td>*</td>
</tr>
<tr>
<td>ADI-R repetitive behavior</td>
<td>4 ± 1.7 (1 - 8)</td>
<td>*</td>
</tr>
<tr>
<td>ADOS social + communication</td>
<td>11 ± 3.6 (7 - 21)</td>
<td>*</td>
</tr>
</tbody>
</table>

Note: Values given as Mean ± (Range), TD controls did not undergo ADI or ADOS assessment.

There were no significant differences between individuals with ASD and TD controls within the training set in terms of age (t=1.703, p=0.095), full-scale IQ (t=.946, p=0.348), or total GM volume (t=.399 , p=0.691).

4.2.2 Clinical Group Demographics

In addition to the participants described above, 62 right handed male individuals who had received referrals for a specialist ASD assessment were recruited from the Autism Assessment and Behavioral Genetics Service at the Bethlem Royal Hospital, South London and Maudsley NHS Foundation Trust. All individuals within this sample were diagnosed according to ICD-10 research criteria. Of the 62 individuals, 14 did not receive an ASD diagnosis, 17 met ICD-10 criteria for ASD but not ADI-R or ADOS cutoffs (i.e. ‘clinical’ diagnosis for ASD), and 31 met ICD-10 and ADI-R and/or ADOS cutoffs for the condition (i.e. obtained a ‘research’ diagnosis of ASD). Individuals within this sample were representative of patients assessed in the clinical setting who often present with comorbid conditions. High rates of obsessive compulsive disorder (OCD), attention deficit disorder (ADD), phobias, depression, general and social anxiety disorders were observed across each subgroup within the clinical sample (Figure 4.2).

Table 4.2. Participant Demographics (Clinical Sample)

<table>
<thead>
<tr>
<th></th>
<th>No ASD (n=14)</th>
<th>Clinical ASD (n=17)</th>
<th>Research ASD (n=31)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>34 ± 9.19 (20-45)</td>
<td>27 ± 5.67 (19-36)</td>
<td>29 ± 7.65 (18-44)</td>
</tr>
</tbody>
</table>
Within the clinical sample; individuals who did not receive an ASD diagnosis were significantly older than those with a clinical diagnosis of ASD ($t=2.574, p=0.015$) but not those with a research diagnosis of ASD ($t=1.865, p=0.015$). No significant difference in full scale IQ was observed between individuals with no ASD diagnosis and those with a clinical ($t=1.017, p=0.318$) or research diagnosis of ASD ($t=1.930, p=0.060$). Furthermore, individuals with clinical and research diagnoses did not significantly differ ($p<0.05$) in terms of age or FSIQ.

In terms of ASD symptom severity as measured by the ADI-R and ADOS, individuals who received a clinical referral but did not meet ICD-10 criteria for an ASD diagnosis did not show a significant difference in any of the ADI-R subdomains or combined ADOS social and communication scores from individuals who met criteria for a clinical ASD diagnosis but did not reach ADI-R and ADOS cutoffs for a research diagnosis ($p<0.05$). Those with a research diagnosis showed statistically significant increases across all these measures compared to both the no ASD and clinical ASD groups ($p<0.05$).

### 4.2.3 MRI Data Acquisition:

All MRI scanning was conducted at the Institute of Psychiatry, Psychology and Neuroscience, King’s College London, UK, using a 3T GE Signa System (General-Electric). A quantitative T1-mapping sequence (see Chapter 2.4; Deoni et al., 2008) was used to derive synthetic high-resolution structural T1-weighted inversion-recovery images, with $1 \times 1 \times 1$ mm resolution, a $256 \times 256 \times 176$ matrix, repetition time = 1,800 ms, inversion time = 50 ms, flip angle = $20^\circ$, and field-of-view = 25 cm.
4.2.4 Cortical Surface Reconstruction

The FreeSurfer software package (http://freesurfer.net/, v5.3.0) was used to derive models of the cortical surface for each T1-weighted image (see Chapter 2.5). In brief, a single filled white-matter volume was generated for each hemisphere after intensity normalization, the removal of extra cerebral tissue, and image segmentation using a connected components algorithm. A triangular surface tessellation was then generated for each white-matter volume by fitting a deformable template, resulting in a cortical mesh for the white and grey/pial (i.e., outer) surfaces with ~150,000 vertices (i.e., points) per hemisphere. Prior to surface reconstruction a radiologist screened scans to exclude participants with clinically significant abnormalities or excessive motion artifacts. A secondary quality check was then performed to identify cortical surface reconstructions of insufficient quality (i.e. with visible reconstruction errors), which were excluded from the analysis and are not described here. Overall the drop out within the training set was equal between groups and was <10% of the total available sample. In order to avoid the introduction a subjective bias no further manual edits were performed.

Each individual’s surface reconstructions were used to calculate vertex-wise measures of cortical morphometry, which included i) point-wise estimates of cortical volume (VOL), ii) cortical thickness (CT), iii) pial surface area (SA), iv) the local gyrification index (lGI), v) average convexity/concavity (SULC, i.e. sulcal depth), and vi) percent contrast of grey-white matter signal intensity (GWPC), where grey matter intensity was sampled at the grey-white matter boundary (i.e. 0% projection fraction) and contrasted with white matter intensity sampled at 1mm subjacent to the grey-white matter boundary. Further details on the calculation of these measures are provided in Chapter 2.5 (for CT, SA, lGI, and SULC) and the methods section of Chapter 3 (for GWPC). Prior to further statistical analyses, each morphometric feature (excluding lGI) was smoothed using a 15mm FWHM surface based Gaussian kernel. As lGI utilizes cortical SA measures sampled from a 15mm diameter spherical ROI, and is hence an inherently smooth feature, no additional smoothing was applied to measures of lGI.

4.2.5 Pattern Classification

Classification of individuals with ASD and TD controls was performed using two
pattern classification algorithms, the support vector machine (SVM) (Vapnik, 1995) and Gaussian process classification (GPC) (Rasmussen and Williams, 2006, Marquand et al., 2010). Whole brain vertex-wise measures of the six morphometric features described above were each used independently of each other as feature sets for classification. Details of SVM and GPC methods are provided in Chapter 2.7. In brief, SVMs operate by identifying a decision boundary (linear or non-linear) known as a hyperplane. The hyperplane separates the experimental classes with a maximum margin. Test cases are then classified as cases or controls depending on which side of the hyperplane they are projected to (Boser et al., 1992). GPC is a Bayesian classification method, which provides probabilistic predictions of class ownership. These predictive class probabilities make GPCs particularly well suited for clinical contexts where the probability of an individual having a particular condition (i.e. psychiatric disorder) is of primary interest. In brief, GPCs can be understood as a distribution over functions, with GP inference consisting of applications of Bayes’ rule to find the posterior function distribution that best approximates the training data.

The performance of each SVM and GPC model within the research sample was assessed via leave one out cross validation (LOO-CV). Within each LOO-CV fold, data from all but one individual was used to train the classifier. The resulting pattern recognition model was then tested on the individual who was excluded from the training phase. This process was repeated \( S \) times (where \( S \) is the total number of subjects). Classification accuracy was then defined as the percentage of subjects correctly assigned to the ASD or control group. For each classification model prediction sensitivity (SE) and specificity (SP) were also calculated (see Chapter 2.7.3). Model significance was then assessed via \( n=1000 \) permutations of the class labels to derive a null distribution of model accuracies by which to test significance against.

Following the training phase, the established models that showed the highest overall accuracies within the research training sample were used to predict the class membership of individuals within the independent clinical sample. This sample contained both ASD positive (clinical or research criteria) and ASD negative cases, which for purposes of classification were labeled as analogous to TD controls in the training sample. Within the clinical sample, in addition to overall classification
accuracy, prediction accuracy within the ASD negative, clinical and research ASD subgroups was calculated.

4.2.6 Classification Using Feature Selection

In addition to the models above we retrained each classifier using a general linear model (GLM) based feature selection technique (see Chapter 2.7.4). This method sought to limit vertices used to train models to only those that had a significant mean difference between groups, and thus increase the amplitude of signal in the data with regards to the class labels (i.e. ASD/TD). For each cortical feature the main effect of group was identified by a regression of a GLM using only the training data. Here, we used diagnostic group as a categorical fixed-effects factor and age and full-scale IQ as continuous covariates:

\[ Y_i = \beta_0 + \beta_1 \text{Group} + \beta_2 \text{Age} + \beta_3 \text{FSIQ} + \epsilon_i \]

where \( \epsilon_i \) is the residual error at vertex \( i \). A nested LOO-CV procedure was then applied to select the \( p \)-value used to threshold the vertex-wise statistical maps, such that only vertices with a statistically significant effect of group (\( \beta_1 \)) that exceeded this threshold were included as input features for classification.

This nested LOO-CV procedure followed the following steps. i) For each LOO-CV fold across the research sample (S) one individual was left out to act as a test case, with the remaining individuals (S-1) acting as training data for that fold. ii) Within each of these LOO-CV folds a second nested LOO-CV loop was performed within the training data (i.e. S-1) for that particular fold. iii) For each of these nested LOO-CV folds another individual was left out and acted as a test case within the nested fold. iv) Using only the remaining training data within the nested fold, the above GLM was fit and the statistical maps for the main effect of group (\( \beta_1 \)) thresholded separately for a range of \( p \)-values (\( p > 0.4, 0.3, 0.2, 0.1, 0.05, 0.01, 0.001 \)). v) Within the same nested training group, classification models were trained for each \( p \)-value using only vertices where the effect of group (\( \beta_1 \)) exceeded the significance threshold. vi) Model accuracy was estimated for each of these classifiers using LOO-CV within the nested loop. vii) The \( p \)-value threshold that yielded the highest classification accuracy within the nested LOO-CV loop was then selected and applied to the initial left out training
Thus, only vertices that showed a main effect of group that exceeded the \( p \)-value selected through nested LOO-CV to provide the highest overall accuracy were used to train the classifier within the initial training sample (S-1) to make a prediction on the initial left out test case.

To apply this procedure within the independent clinical validation data set \( i \) the \( p \)-value threshold that yielded the highest estimated CV accuracy on average across all LOO folds in the research training set was selected. \( ii \) This \( p \)-value threshold was then applied to statistical maps of the main effect of group across the entire research training set. \( ii \) Vertices exceeding this statistical threshold were then used to retrain classifiers on the entire research training set. \( iv \) These established models were then used to make predictions within the independent clinical data set.

4.3 Results

4.3.1 Classification of ASD and TD individuals in the research sample
Using whole brain vertex wise morphometric measures SVM was not able to classify individuals with ASD from TD controls within the research sample at accuracies above what would be expected statistically by chance (VOL 50% Accuracy [48/52% SE/SP], \( p=0.56 \); CT 50% [48/52%], \( p=0.53 \); SA 54% [52/56%], \( p=0.11 \); lGI 41% [37/44%], \( p=0.88 \); SULC 56% [56/56%], \( p=0.44 \); GWPC 50% [48/52%], \( p=0.48 \), Figure 4.1.
Figure 4.1: Support Vector Machine Predictions Within the Training Set: Class predictions for individuals within the training set derived by leave one out cross validation are shown in scatter plots for each cortical feature. A = cortical volume, B = cortical thickness, C = surface area, D = local gyrification index, E = sulcal depth, D= grey-white matter percent contrast. ‘Projection onto Weight Vector’ represents unthresholded SVM predictions, normalized between -1 (TD) and 1 (ASD) for visualization.

Similar model performance was achieved using GPC (VOL 52% [52/52%], $p=0.41$; CT 50% [48/52%], $p=0.46$; SA 57% [59/56%], $p=0.13$; lGI 41% [33/48%], $p=0.81$; SULC 52% [48/56%], $p=0.33$; GWPC 52% [52/52%], $p=0.44$), Figure 4.2.
When limiting the feature set to include only vertices that showed a significant group difference according to a traditional GLM both SVM (VOL 44% [37/52%]; CT 52% [52/52%]; SA 50% [56/44%]; lGI 41% [37/44%]; SULC 61% [67/56%] \( p = 0.345 \); GWPC 48% [52/44%]; Figure 4.3) and GPC (VOL 50% [48/52%]; CT 54%...
[48/59%]; SA 50% [56/44%]; lGI 43% [37/48%]; SULC 61% [59/63%], $p=0.074$; GWPC 56% [56/56%], $p=0.197$) Figure 4.4) overall classification accuracies remained modest.

Figure 4.3: Support Vector Machine Predictions within the Training Set using GLM based Feature Selection: Class predictions for individuals within the training set derived by leave one out cross validation are shown in scatter plots for each cortical feature. A = cortical volume, B = cortical thickness, C = surface area, D = local gyrification index, E = sulcal depth, D= grey-white matter percent contrast. ‘Projection onto Weight Vector’ represents unthesholed SVM predictions, normalized between -1 (TD) and 1 (ASD) for visualization.
Figure 4.4: Gaussian Process Classification Predictions within the Training Set using GLM based Feature Selection: Predictive class probabilities for individuals within the training set derived by leave one out cross validation are shown in density and scatter plots for each cortical feature. Predicted class probability >0.5=ASD, <0.5=TD. A = cortical volume, B = cortical thickness, C = surface area, D = local gyrification index, E = sulcal depth, D= grey-white matter percent contrast
4.3.2 Predicting ASD Diagnosis within Clinical Sample

The best performing classification models within the research training sample showed relatively modest overall classification accuracy (61%). While this is in line with previous reports in large ASD cohorts using measures of CT (Sabuncu and Konukoglu, 2014), given the relatively small size of our training sample we were not able to show these results were statistically significant relative to chance. Thus, interpretation these classifiers ability to make predictions within the clinical testing set is made difficult as it is uncertain if this predictive value is reflective of biological differences between the groups. However, as an exploratory step we applied the best performing models (>55% overall accuracy) to make class predictions within the clinical sample.

Across all SVM models, those trained on SULC measures with (61%, \( p=0.345 \)) and without feature selection (56%, \( p=0.44 \)) showed the highest classification accuracies within the research training sample. When these models were applied to the clinical sample they showed overall classification accuracies below chance levels with (39%) and without feature selection (39%), Table 4.5. These classifiers performed equally at or below chance level across all patient subgroups including those with no ASD diagnosis, a clinical, or a research diagnosis of ASD, Figure 4.3.

![Figure 4.5: Support Vector Machine Predictions within the Testing Set: Class predictions for individuals within the testing set are shown in scatter plots for SVM classifiers trained on vertex wise measures of sulcal depth both with (A) and without (B) GLM based feature selection. 'Projection onto Weight Vector' represents unthresholed SVM predictions, normalized between -1 (TD) and 1 (ASD) for visualization.](image-url)
Table 4.3 Classification Accuracies within Subgroups of the Clinical Sample

<table>
<thead>
<tr>
<th>Classifier</th>
<th>Overall</th>
<th>No ASD</th>
<th>Clinical ASD</th>
<th>Research ASD</th>
</tr>
</thead>
<tbody>
<tr>
<td>SVM SULC</td>
<td>24/62 (39%)</td>
<td>7/14 (50%)</td>
<td>6/17 (35%)</td>
<td>11/31 (35%)</td>
</tr>
<tr>
<td>SVM SULC*</td>
<td>24/62 (39%)</td>
<td>6/14 (43%)</td>
<td>6/17 (35%)</td>
<td>12/31 (39%)</td>
</tr>
<tr>
<td>GPC SA</td>
<td>25/62 (40%)</td>
<td>4/14 (29%)</td>
<td>11/17 (65%)</td>
<td>10/31 (32%)</td>
</tr>
<tr>
<td>GPC SULC*</td>
<td>26/62 (42%)</td>
<td>7/14 (50%)</td>
<td>6/17 (35%)</td>
<td>13/31 (42%)</td>
</tr>
<tr>
<td>GPC GWPC*</td>
<td>50/62 (81%)</td>
<td>12/14 (86%)</td>
<td>15/17 (88%)</td>
<td>23/31 (74%)</td>
</tr>
</tbody>
</table>

Note: Classification accuracies for models applied to the clinical sample are given as number of correctly labeled / total number of individuals overall and for each subgroup. SVM=Support Vector Machine, GPC=Gaussian Process Classifier, SULC=Sulcal Depth, SA=Surface Area, GWPC=grey:white matter percent contrast. *Classifier used general linear model based feature selection.

Among GPC models trained within the research sample without GLM based feature selection, measures of SA were found to return the highest overall classification accuracy (57%, $p=0.13$), but showed below chance levels when applied to the clinical sample (40%), Figure 4.6. Two additional GPC models trained using feature selection on measures of SULC (61%, $p=0.074$) and GWPC (56%, $p=0.197$) showed overall classification accuracies above 55% within the research sample. When applied to the clinical sample the model based on measures of SULC showed overall accuracies below chance levels (42%), Figure 4.6. Accuracies for these models within the clinical subgroups are given in Table 4.3.

Figure 4.6: Gaussian Process Classification Predictions within the Testing Set: Predictive class probabilities for individuals within the test set are shown in density and scatter plots for the GPC trained on measures of (A) surface area and (B) sulcal depth when using GLM based feature selection (B).
However, the GPC trained on GWPC measures that were restricted to vertices that showed a significant effect of group (\(p\) value threshold of 0.05) within the training set showed a high overall accuracy (81%) within the clinical set, across all clinical subgroups (Table 4.3). However, we were unable to show this model performed above chance at a statistically significant level within the training set (57%, \(p=0.25\)).

![Image](image.png)

**Figure 4.7**: Gaussian Process Classification Predictions within the Testing Set using measures of Tissue Contrast and GLM based Feature Selection: Predicted class probabilities for individuals within the test set are shown in density and scatter plots for measures of grey-white matter percent contrast using GLM based feature selection

### 4.4 Discussion

The first aim of this study was to establish predictive models for ASD based on different morphometric cortical features within a training set of TD controls and individuals with ASD acquired in a research setting. Secondly, this study aimed to test the ability of these classifiers to predict ASD within an independent sample of individuals who had been referred for a clinical ASD diagnostic assessment, thus representing a ‘real-world’ clinical setting. While previous reports suggest that measures of cortical structure can be used to classify ASD individuals (Ecker et al., 2010a, Wee et al., 2014, Jiao et al., 2010, Hazlett et al., 2017, Sabuncu and Konukoglu, 2014), the current study was not able to replicate these findings in our research training sample. As an exploratory step, we applied the best performing classifiers within the research sample to make predictions as to ASD diagnosis within an independent clinical sample. In this context, one model showed the ability to discriminate individuals with and without ASD diagnoses at high overall accuracy.
(81%). However, given this model’s performance in the research setting this result must be considered as preliminary and suggestive only.

While a number of classifiers for ASD have been proposed and reported to show significant classification accuracies when discriminating individuals with ASD from TD controls within research settings, we were not able to replicate these results within our sample. A likely key contributor to this is the highly heterogeneous nature of the ASD phenotype. Within a heterogeneous cohort such as the ASD population, maximal classification accuracy based on measures of brain structure is likely to be rather modest. For example, classification studies reporting validation on independent ASD research samples have reported accuracies in the range of 59-72% (Sabuncu and Konukoglu, 2014, Shen et al., 2017). Whilst some of the accuracies we report here are in this range, we were not able to establish the statistical significance of these classifiers due to the small size of our research sample.

When classifiers are trained on relatively small samples (i.e. n<100) drawn from a heterogeneous population (e.g. ASD) a high degree of variance in model performance is to be expected. Furthermore, cross validation methods that test a small number of cases for at each fold (i.e. n=1) may also increase variance in estimates of model performance. Such variance has been shown across classification studies in heterogeneous psychiatric and neurodevelopmental conditions along with a trend towards decreasing model performance with increasing training sample sizes (Wolfers et al., 2015) (Figure 4.8). Thus, the development of approaches that directly address phenotypic heterogeneity within ASD will be of critical importance for the successful delivery of classification methods that have translational value in clinical settings. Methods that seek to parse heterogeneity based on one particular source of variance (e.g. biological sex) could be of particular benefit and are a major theme in the chapters that follow in this thesis.
Figure 4.8: Accuracy as a Function of Number of Included Patients in the Classification of Psychiatric and Neurodevelopmental Conditions (Wolfers et al., 2015): This figure shows a plot of classification accuracies for studies based on brain imaging data as a function of the number of patients included in those studies across a variety of heterogeneous psychiatric and neurodevelopmental conditions, including ASD. Two trends are of note, i) A high degree of variance in accuracies of studies including less than 100 patients can be observed. ii) Classifier performance is also seen to decrease with increasing numbers of patients within each study, which likely is attributable to increased sample heterogeneity. SCZ = schizophrenia; BPD = bipolar disorder; MDD = major depressive disorder; OCD = obsessive compulsive disorder; SAD = social anxiety disorder; PTSD = post-traumatic stress disorder; SP = specific phobia; ADHD = attention-deficit/hyperactivity disorder; ASD = Autism spectrum disorder; C = controls.

When comparing the current results to previous ASD classification studies in small research samples that have reported good model performance (e.g. 70-100% accuracy) it is important to carefully note if and how feature selection has been applied. Broadly speaking, feature selection incorporates a variety of methods used to identify and include only features (e.g. MRI data points) that are informative to the classification problem at hand (Cuongnet et al., 2011, Chu et al., 2012, Mwangi et al., 2014). While feature selection offers a valid way to increase signal within data to allow for better classification between groups, it also introduces the possibility of overfitting one’s model to the training set. To avoid overfitting it is essential to perform feature selection in a non-biased fashion such as through a nested LOO-CV framework like was conducted in the current study or to ensure so called ‘expert
based’ feature selection (e.g. ROI selection) is fixed a-priori. Ultimately, the gold standard test for over fitting is the application of a trained classifier to a completely independent test set. Thus, while previous studies reporting exceptional model performance through applications of feature selection in ASD are valuable for their ability to identify features that can distinguish cases from controls within individual research samples, testing of these models on independent research or clinical samples is required in order to establish their translational ability.

Whilst our trained classifiers did not statistically exceed chance, as an exploratory step we applied our best performing models to predict ASD diagnoses within our independent clinical sample. This is based on the rationale that these classifiers may still have learned useful discriminating information, despite not exceeded chance owing to the small size of the training set. We found the majority of these models showed decreased overall classification performance within the clinical testing set. This result was expected based on previous reports that overall classification accuracy decreases when models are applied to independent samples (Sabuncu and Konukoglu, 2014) and was likely further enhanced by the clinical nature of our testing sample, which introduces differences in both class frequencies (Hahn et al., 2013) and the clinical ASD phenotype between the research and clinical samples.

One notable exception to the above results was the GPC trained on measures of grey-white matter percent contrast (GWPC) taken at the grey-white matter boundary (Andrews et al. 2017) which showed high overall accuracy within the clinical sample (81%) and across all diagnostic sub-categories (74-88%). This is potentially of interest as this feature was designed with intent to investigate a particular aspect of ASD neuropathology, namely poor definition of the grey-white matter boundary (Avino and Hutsler, 2010) and thus may have increased sensitivity to detect the condition compared too more general measures of cortical morphology. However, interpretation of this finding is limited in light of our inability to show this model performed significantly above chance level within the research sample, or within the clinical sample via LOO-CV. The possibiility of this result being attributed to chance is further increased given the relatively large number of models trained within this study (n=24). If this model was found to identify individuals with ASD in a statistically significant fashion, correction for multiple comparisions would be
critically important before the results could be confidently attributed to a informative biological pattern in the data.

It is important to note that if the above limitations can be addressed and an efficacious imaging biomarker for ASD is developed, the current high cost of MRI scanning makes the use of imaging as a population screening tool for ASD implausible at this time. However, MRI based screenings may still be able to offer benefits to populations at high risk for the condition. For example, within the current study ASD prevalence was found to be 77% among the sample of individuals referred for a clinical ASD assessment. This is important due to the fact that models trained on artificially balanced groups will suffer from high rates of false positives (type I errors) when applied to a general population with prevalence rates in the low single digits, regardless of initial model performance (i.e. high SE and SP). Accordingly, once a classification model has been established it is important to take into account any differences in class frequencies that may exist between training and test groups (Hahn et al., 2013). Taking the above factors into account, if an imaging based biomarker is to be successfully implemented targeting screening within appropriate high-risk populations with elevated ASD prevalence will be as equally as important as the identification of the biomarker in the first place.

4.5 Conclusions
Autism spectrum disorder (ASD) is a highly heterogeneous condition that poses several hurdles in the development of a MRI based biomarker for the condition. While a large body of evidence suggests that ASD is accompanied by changes in brain structure, the current study was not able to replicate previously reported findings that measures of cortical morphometry can identify individuals with ASD with significant accuracy in the research setting. Accordingly, we were not able to test the hypothesis that models trained within the research setting translate to clinical samples. Future classification studies would benefit by recruiting large scale ASD clinical samples that allow for the development of methodologies that directly address issues of heterogeneity in the condition though biological subtyping. This along with the identification of imaging features with increased sensitivity to detect ASD phenotypes should improve upon maximal classification accuracies within the condition.
References


Chapter 5: Association Between the Probability of Autism Spectrum Disorder and Normative Sex-Related Phenotypic Diversity in Brain Structure

Male preponderance is an understudied hallmark of autism spectrum disorder (ASD) and many other neurodevelopmental and psychiatric conditions. While some degree of this male preponderance is likely attributable to sex differences in the ASD phenotype and male biased diagnostic tools, it has been purposed that variance in biology attributable to normative sex differences may modulate ASD risk (Werling, 2016).

The following chapter details a study providing an initial ‘proof of concept’ for a novel use of multivariate pattern recognition to characterize normative phenotypic diversity associated with biological sex, and how this aspect of neuro-diversity is associated with probability for ASD. This study was recently published in JAMA Psychiatry (Ecker, et al 2017) and is included in its final published format in the pages that follow. Contributions to this publication by the current author included significant input on the study concept and design, data analysis and interpretation, and drafting of the manuscript. Supplementary materials for this chapter can be found in Appendix III.
Association Between the Probability of Autism Spectrum Disorder and Normative Sex-Related Phenotypic Diversity in Brain Structure

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for the Medical Research Council Autism Imaging Multicentre Study (MRC-AIMS) Consortium

IMPORTANCE: Autism spectrum disorder (ASD) is 2 to 5 times more common in male individuals than in female individuals. While the male preponderant prevalence of ASD might partially be explained by sex differences in clinical symptoms, etiological models suggest that the biological male phenotype carries a higher intrinsic risk for ASD than the female phenotype. To our knowledge, this hypothesis has never been tested directly, and the neurobiological mechanisms that modulate ASD risk in male individuals and female individuals remain elusive.

OBJECTIVES: To examine the probability of ASD as a function of normative sex-related phenotypic diversity in brain structure and to identify the patterns of sex-related neuroanatomical variability associated with low or high probability of ASD.

DESIGN, SETTING, AND PARTICIPANTS: This study examined a cross-sectional sample of 98 right-handed, high-functioning adults with ASD and 98 matched neurotypical control individuals aged 18 to 42 years. A multivariate probabilistic classification approach was used to develop a predictive model of biological sex based on cortical thickness measures assessed via magnetic resonance imaging in neurotypical controls. This normative model was subsequently applied to individuals with ASD. The study dates were June 2005 to October 2009, and this analysis was conducted between June 2015 and July 2016.

MAIN OUTCOMES AND MEASURES: Sample and population ASD probability estimates as a function of normative sex-related diversity in brain structure, as well as neuroanatomical patterns associated with low or high ASD probability in male individuals and female individuals.

RESULTS: Among the 98 individuals with ASD, 49 were male and 49 female, with a mean (SD) age of 26.88 (7.18) years. Among the 98 controls, 51 were male and 47 female, with a mean (SD) age of 27.39 (6.44) years. The sample probability of ASD increased significantly with predictive probabilities for the male neuroanatomical brain phenotype. For example, biological female individuals with a more male-typical pattern of brain anatomy were significantly (i.e., 3 times) more likely to have ASD than biological female individuals with a characteristically female brain phenotype (P = .02; 95% CI, 0.69-0.86; P < .001, difference in P-values, 0.48; 95% CI, 0.09-0.68). This finding translates to an estimated variability in population prevalence from 0.2% to 1.3%, respectively. Moreover, the patterns of neuroanatomical variability carrying low or high ASD probability were sex-specific (e.g., inferior temporal regions, where ASD has different neurological underpinnings in male individuals and female individuals).

CONCLUSIONS AND RELEVANCE: These findings highlight the need for considering normative sex-related phenotypic diversity when determining an individual’s risk for ASD and provide important novel insights into the neurobiological mechanisms mediating sex differences in ASD prevalence.


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Autism spectrum disorder (ASD) is a complex neurodevelopmental condition that is 2 to 5 times more common in male individuals than in female individuals. While the male preponderance prevalence of ASD might partially be explained by sex differences in clinical symptoms, etiological models suggest that the biological male phenotype (e.g., in general) carries a higher risk for ASD than the female phenotype. However, despite the growing number of studies examining sex differences in the brain in ASD, those by Lat et al\(^\text{40}\) and by Scher et al\(^\text{41}\), this hypothesis has never been tested directly to our knowledge, and the neurobiological mechanisms that underpin the male preponderance prevalence of ASD remain elusive.

This study examined the probability of ASD as a function of normative sex-related phenotypic diversity in brain structure. To do so, we initially developed a predictive model of biological sex based on multi-variant differences in brain structure in a sample of typically developing (TD) male- and female-control individuals. This normative model was subsequently applied to male individuals and female individuals with ASD. Unlike recent studies examining sexual dimorphism of the brain\(^\text{42}\) and sex differences in brain structure\(^\text{38}\), we used a probabilistic pattern classification approach\(^\text{38}\), which allowed us to accommodate inter-individual phenotypic diversities within and across the binary categories dictated by biological sex. As a result, we were able to examine the probability of ASD along a normative phenotypic: sexes ranging from the characteristic female to male brain phenotype and C2 to identify the patterns of sex-related neuroanatomical variability associated with low or high probability of ASD.

We based our analysis on measures of cortical thickness (CT) because these measures have previously been shown to be highly variable between neurotypical male individuals and female individuals\(^\text{43}\) and tend to be significantly altered in individuals with ASD\(^\text{44}\), as shown by Echer et al\(^\text{45}\) and by Wallace et al\(^\text{46}\). Moreover, measures of CT are less sensitive to global confounds associated with biological sex (e.g., differences in total brain volume) than other morphometric features (e.g., cortical surface area).\(^\text{47}\) Last, we investigated whether the cortical underpinnings of ASD are significantly modulated by biological sex, and whether ASD has common or distinct neuroanatomical underpinnings in male individuals and female individuals.

**Methods**

**Participants**

Ninety-eight right-handed adults with ASD (49 male and 49 female; mean [SD] age, 26.88±7.18 years) and 98 matched neurotypical controls (50 male and 48 female; mean [SD] age, 27.39±6.44 years) aged 18 to 42 years were recruited locally by advertisement and assessed at the Institute of Psychiatry, Psychology and Neuroscience (IOPPN), London, England, and the Autism Research Centre, Cambridge, England (Table 1). Approximately equal ratios of cases to controls and male individuals to female individuals were recruited within sites (Table 1 in the Supplement). Exclusion criteria included a history of major psychiatric disorder (e.g., psychosis), head injury, ASD-associated genetic disorders (e.g., fragile X syndrome and tuberous sclerosis), or any other medical condition affecting brain function (e.g., epilepsy). We also excluded individuals taking antipsychotic medication, mood stabilizers, or benzodiazepines.

All participants with ASD were diagnosed according to the Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition criteria. The clinical diagnosis was confirmed using the Autism Diagnostic Interview-Revised (ADI-R)\(^\text{20}\) or the Autism Diagnostic Observation Schedule (ADOS).\(^\text{21}\) All participants diagnosed with the ADI-R reached algorithm cutoffs in the total score (cutoff of 80), score (cutoff of 6), social communication (cutoff of 10), and repetitive behaviors and stereotypes (cutoff of 6). Although failure to reach the cutoff in the repetitive domain by maximally 7 points was permitted. Because reliable informants were unavailable, we were unable to obtain ADI-R diagnostic data from 5 female individuals with ASD and confirmed diagnostic status using ADOS cutoffs. In other participants, the ADOS scores were used to assess the severity of current symptoms. One female individual with ASD fell short by 1 point on the ADI-R communication and repetitive behavior domains but met ADOS criteria. All participants had a full scale IQ greater than 70 assessed using the Wechsler Abbreviated Scale of Intelligence.\(^\text{21}\) Participants gave informed consent in accord with ethics approval by the National Research Ethics Committee, South East England. The study dates were June 2003 to October 2009, and this analysis was conducted between June 2015 and July 2016.

**Magnetic Resonance Imaging Data Acquisition**

Imaging took place at the IOPPN and the Addenbrooke's Hospital, Cambridge, using a 3-T system (Siemens, GE Medical Systems). A specialized acquisition protocol was used, consisting of T1-weighted sagittal T2 T1 images, which resulted in high-resolution structural 3T-weighted T1-weighted recovery images, with 1 x 1 x 1 mm resolution, a 256 x 256 x 1° pixels matrix, repetition time of 1100 milliseconds, inversion time of 30 milliseconds, flip angle of 30°, and field of view of 25 cm.
Cortical Reconstruction Using FreeSurfer
A software program (FreeSurfer, version 5.1.0) [https://surfer
mmgh.harvard.edu] was used to derive cortical surface models for each T1-weighted image. These well-validated and fully automated procedures have been extensively described elsewhere.10,32 All surface models were visually inspected for errors, and scans with visible reconstruction inaccuracies were excluded from the statistical analysis (dropout = 10%). We based our analysis on measures of CT (ie, the closest distance from the gray matter (GM) or white matter boundary to the GM or subcortical fluid boundary at each cortical vertex).10 Cortical thickness maps were smoothed with a 15-mm surface-based gaussian kernel.

Gaussian Process Classification
Gaussian process classification (GPC)14,19 was used for the probabilistic prediction of biological sex based on neuroanatomical variability in CT (details are available in the eMethods in the Supplement). In brief, we initially developed a normative model of biological sex by predicting the binary class labels y, e (-1, +1) for female and male T1 controls, respectively, using CT measures estimated at approximately 120,000 vertices on the cortical surface (Gaussian Processes for Machine Learning: Matlab toolbox [http://www.gaussianproces
s.org/gpml/code/matlab/doc/]). In addition to the overall model accuracy (ie, proportion of biological male individuals or female individuals correctly classified as phenotypic male individuals or female individuals), GPC returned a set of posterior-class probabilities indicating the probability of an individual belonging to the male or female category. Class probabilities ranged from 0 to 1 for the characteristic female to male brain phenotype, respectively, so that a class probability of 0.5 represents a binary cutoff separating both classes. Classifier performance was validated by cross-validation and tested for statistical significance via 1000 permutations of class labels. The normative model was subsequently applied to male individuals and female individuals with ASD.

A predictive mapping approach10 was used to identify (1) the spatially distributed patterns of neuroanatomical variability in CT characteristics for the neurotypical female (or male brain phenotype and (2) the set of brain regions associated with low or high probability of ASD. Normative (ie, low or high probability) patterns were derived by quantifying the extent to which the individual’s neuroanatomy reflects the spatial representation of the decision function, ie, the weight vector w separating neurotypical male individuals from female individuals phenotypically, the product of w and CT. The high ASD probability pattern included all male individuals (or female individuals) with a male neuroanatomical phenotype (ie, class probability = 0.5). The low ASD probability pattern included all male individuals (or female individuals) with a female neuroanatomical phenotype (ie, class probability = 0.5). We also identified patterns across all individuals along the normative axis of predictive class probabilities. At each vertex, the resulting maps were summarized by the mean value across all individuals within each examined group.
Estimation of ASD Probability

To estimate the probability of ASD as a function of normative-phenotypic variability in brain structure, we converted the continuous axis of class probabilities into a set of discrete bins (e.g., from 0 to 1 in steps of 0.05). Within each bin, the sample probability of ASD was determined as the ratio of individuals with ASD relative to the total number of individuals per bin. Moreover, these data allowed us to estimate the population prevalence of ASD given that an individual exhibits a male or female neuroanatomical phenotype in addition to being biologically male or female (details are available in the Methods in the Supplement). In brief, using Bayes theorem, we combined our sample prevalence estimates with previously published population prevalence rates of ASD for biological male and female individuals (i.e., 1:42 for male individuals and 1:89 for female individuals). In this way, we were able to estimate the population prevalence of ASD (12 = 1) given a male (or female) neuroanatomical brain phenotype (M) for biological male individuals (or female individuals) (S), that is, P(12 = 1 | M, S). Confidence intervals were determined using an exact binomial test implemented in a software package (R Project for Statistical Computing; https://www.r-project.org).

Verbena Between Group Comparison of CT

Verbena statistical analysis of CT measures was conducted using the R2004a Matlab toolbox (StatSoft; http://www.math.mcgill.ca/keith/unifstat). For the comparison between male and female controls, parameter estimates were obtained by regression of an age-normed linear model at each vertex i, with biological sex and center as categorical fixed-effects factors and total GM volume as a continuous covariate. To examine whether the neuroanatomy of ASD is significantly modified by biological sex, we also included a main effect of diagnostic group and biological sex and a group x sex interaction as follows:

\[ y_i = \beta_0 + \beta_1 \text{Group} + \beta_2 \text{Sex} + \beta_3 (\text{Group} \times \text{Sex}) + \varepsilon_i \]

where \( y_i \) refers to the variable that is predicted by the model (i.e., the CT at voxel \( i \)) and \( \varepsilon_i \) is the residual error. Effects were estimated from the coefficients \( \beta_3 \), normalized by the corresponding standard error. Corrections for multiple comparisons were performed using a random field theory-based cluster analysis for nonneurotological images at a cluster threshold of 2-tailed \( P < .05 \) 30. We also confirmed our findings using a nonparametric clustering approach (2-tailed \( P < .05 \)) using 10,000 permutations of the original data.

Results

Prediction of Biological Sex Based on Normative Variability in CT

Overall, our CT-driven probabilistic classifier predicted biological sex at an accuracy of 71% under cross-validation in neuroontological controls, which was significantly higher than what would be expected by chance (P = .001 obtained via permutation testing). In total, 83.3% (32 of 39) of all biological female individuals were correctly allocated to the category of phenotypic female individuals and 74.4% (34 of 46) of all biological male individuals to the category of phenotypic male individuals (Figure 1A, lower panel). Across individuals, class probabilities ranged continuously from 0 to 1 for the characteristic female/male brain phenotype, respectively (Figure 1A, upper panel). Moreover, the patterns of CT variability associated with the phenotypic shift of the brain from a female to male presentation resembled the set of brain regions where female individuals significantly differed from male individuals (Figure 1C and Figure 2 in the Supplement). Therefore, while a perfect phenotypic differentiation between biological sexes could not be achieved, we were able to separate male individuals from female individuals to some extent in most cases, based on moderate differences in brain structure.

Phenotypic Prediction of Biological Sex for Individuals With ASD

When applying our normative predictive model of biological sex to the independent sample of individuals with ASD, we found that, in female individuals with ASD, predictive class probabilities for the male neuroanatomical brain phenotype were significantly increased relative to female controls (OR = 6.0, P < .001) (Figure 1B). Overall, 39 of 49 female individuals (79.6%) with ASD were allocated to the category of phenotypic male individuals, which was significantly more frequent than expected based on the typical ratio of subclassification for female controls (39/49; 80.4%) (15 of 87, respectively, \( \chi^2 = 20.26, P < .001 \)). No such differences were observed in male individuals with ASD, who were correctly allocated to the male category in 80.6% (40 of 50) of all cases and also did not differ significantly from male controls in predictive class probabilities (95% = 1.35, 2-tailed \( P > .37 \)). Therefore, when representing individuals along a single axis of normative sex-related phenotypic diversity in brain structure, female individuals with ASD displayed a CT pattern that resembled more closely the neuroanatomical male rather than female (pseudophenotype). This phenotypic shift in female individuals is likely to influence the probability of ASD.

ASD Probability as a Function of Normative Sex-Related Phenotypic Variability in Brain Structure

In female individuals with ASD, the sample probability of ASD increased with increasing predictive probabilities for the male neuroanatomical brain phenotype (Figure 3A in the Supplement). More specifically, female individuals with class probabilities exceeding the binary sex cutoff of 0.5 (i.e., biological female individuals falling into the category of phenotypic male individuals) were significantly more likely to have a diagnosis of ASD than biological female individuals with a lower class probability (Figure 3B). For example, for the 95th percentile (25 vs. 24), \( \chi^2 = 20.26, P < .001 \), and additional differences in values (0.48 and 0.39) (\( \chi^2 = 0.68, P = .02 \)). Reciprocally, male individuals with class probabilities below 0.5 (i.e., biological male individuals falling into the category of phenotypic female individuals) were 2.3 times less likely to have ASD than male individuals allocated to the category of phenotypic male individuals (0.41 vs. 0.38, respec-
Figure 1. Gaussian Process Classification of Biological Sex

A. Gaussian process classification between male and female control individuals

B. Predicted posterior for male individuals and remote controls with AS

The results show that the model accurately predicts the sex of individuals based on their genetic characteristics. The predicted posterior for male individuals and remote controls with AS is shown in panel B. The model's performance is validated by the high accuracy rates in panel A.

Biological Sex Significantly Modulates the Cortical Anatomy of ASD

Our study reveals significant differences in the cortical anatomy of individuals with ASD based on their biological sex. The model accurately identifies these differences by analyzing the cortical thickness and surface area of various brain regions. The results suggest that sex-specific traits play a crucial role in the development and manifestation of ASD.

The model's performance is validated by the high accuracy rates in panel A. The predicted posterior for male individuals and remote controls with AS is shown in panel B. The model's accuracy in predicting the sex of individuals is further supported by the high correlation coefficients in panel A.

In conclusion, our findings highlight the importance of considering biological sex in the study of ASD. Further research is needed to explore the mechanisms underlying these sex-specific differences and to develop more effective interventions for individuals with ASD.
Supplement. The clusters with significant group × sex interactions also retained significant when controlling for variability in verbal and performance IQ (Table 5 and eFigure 7 in the Supplement) and thus do not seem to reflect a simple functional difference in these clinical or neurocognitive measures (further details are available in eFigure 8 and the eAppendix in the Supplement).

Discussion

Our study demonstrates that normative sex-related phenotypic diversity in brain structure affects the prevalence of ASD in addition to biological sex alone, with male neuroanatomical characteristics carrying a higher intrinsic risk for ASD than female characteristics. This increase in risk was predominantly driven by female individuals with ASD, who displayed a pattern of co-occurring anatomical variability in CT that resembled more closely the neurotypical male rather than female neuromorphotype overall. Moreover, our study links low and high risk (i.e., probability) of ASD to particular patterns of sex-related neuroanatomical variability, thus providing important novel insights into the neurobiological mechanisms underpinning the male, preponderant prevalence of ASD.

An important feature of GPC is that it is possible to summarize the highly complex and multidimensional pattern of normative sex-related phenotypic diversity in brain structure.
Figure 5. Significant Group × Sex Interactions in Cortical Thickness (CT) as examined by a conventional general linear model-type approach. In these regions, the difference in CT between female individuals with autism spectrum disorder (ASD) and female control individuals significantly exceeded the difference between male individuals with ASD and male control individuals statistical details are available in Table 4 in the Supplement. The group × sex interactions were driven by reduced CT in female individuals with ASD relative to female controls and increased CT in male individuals with ASD relative to male controls (Figure 6 in the Supplement). B. Clusters with significantly increased CT in male individuals with ASD relative to male controls (random field theory-based, cluster corrected, P < 0.05). C. Clusters with significantly reduced CT in female individuals with ASD relative to female controls (random field theory-based, cluster corrected, P < 0.05). Figure 5 in the Supplement shows results of the permutation-based (cluster thresholding of the same magnitude) T3 tests typically developing.
lization of interindividual phenotypic diversity across the binary categories dictated by biological sex. This study is of importance because it has recently been suggested that, on a phenotypic scale, the brain should be considered a "mosaic" of regions, each of relative maleness or femaleness, resulting in significant interindividual variability both within and between sexes. Therefore, the multivariate patterns derived from the classification between neurotypical male and female controls may be interpreted as being representative of such a mosaic, which drives the brain toward a female or male phenotypic end point overall (see also the study by Check and others). Furthermore, we demonstrated that the normative patterns of sex-related neuroanatomical variability predictive of ASD probability, particularly in female individuals, and that low and high ASD probability can be linked to regional differences in C1. These patterns are not only included in some of the brain regions where women typically differ more in men in C1 but also highlight brain areas that have previously been linked to the local behavioral deficits characteristic of ASD (reviewed by Amaral and others) and by Ecker and others. Notably, some of these brain regions (e.g., inferior temporal lobes) carry high or low probability of ASD in women but not in men. Hence, we found that the degree of cortical abnormality in female individuals with ASD significantly exceeded the degree of abnormality in male individuals compared with their respective normative populations. Yet, female individuals with ASD were not more severely impaired than male individuals on the level of autistic symptoms. Therefore, our findings agree with previous reports suggesting that (1) biological sex significantly mediates the neurobiological basis of ASD (e.g., the studies by Lee and others and by Schen and others) and (2) female individuals may have a higher threshold of minimum liability sufficient to cause ASD for reaching affection status than male individuals (reviewed by Westling and others) and may need to deviate more neurobiologically from their normative population to demonstrate a clinical ASD phenotype.


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REFERENCES
Chapter 6: Normative Sex-Related Phenotypic Diversity Across Multiple Measures of Brain Anatomy is Associated with Probability for Autism Spectrum Disorder

This chapter represents a critical extension of the study detailed in chapter 5 regarding the association between sex-related phenotypic diversity in measures of cortical thickness (CT) and the probability for autism spectrum disorder (ASD). As has been noted by others (Cahill 2017) it is unknown if and how the previous results detailed in Chapter 5 (Ecker, et al 2017) will extend to other measures of neuro-anatomy. Here, we extended our initial findings to consider multiple aspects of brain anatomy including both volumetric and geometric cortical features in order to (i) provide a more comprehensive characterization of normative sex related differences in brain structure, (ii) determine how diversity in sex based neurophenotypes are associated with ASD probability, and (iii) estimate how much ASD prevalence in the general population might vary as a function of normative sex-related phenotypic diversity.

This study is currently under review with *The American Journal of Psychiatry* and is presented in the pages that follow in its most recent format as of time of writing. Supplementary materials for this chapter are presented in Appendix IV, the current manuscript proof is presented in Appendix V.
Title: Normative Sex-Related Phenotypic Diversity Across Multiple Measures of Brain Anatomy is Associated with Probability for Autism Spectrum Disorder

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* The Medical Research Council Autism Imaging Multicentre Study Consortium (MRC AIMS Consortium) is a UK collaboration between the Institute of Psychiatry, Psychology and Neuroscience at King’s College, London, the Autism Research Centre, University of Cambridge, and the Autism Research Group, University of Oxford. The Consortium members in alphabetical order are: Anthony J. Bailey (Oxford), Simon Baron-Cohen (Cambridge), Patrick F. Bolton (IoP), Edward T. Bullmore (Cambridge), Sarah Carrington (Oxford), Marco Catani (IoPPN), Bhismadev Chakrabarti (Cambridge), Michael C. Craig (IoPPN), Eileen M. Daly (IoPPN), Sean C. L. Deoni (IoPPN), Christine Ecker (IoPPN), Francesca Happé (IoPPN), Julian Henty (Cambridge), Peter Jezzard (Oxford), Patrick Johnston (IoPPN), Derek K. Jones (IoPPN), Meng-Chuan Lai (Cambridge), Michael V. Lombardo (Cambridge), Anya Madden (IoPPN), Diane Mullins (IoPPN), Clodagh M. Murphy (IoPPN), Declan G. M. Murphy (IoPPN), Greg Pasco (Cambridge), Amber N. V. Ruigrok (Cambridge), Susan A. Sadek (Cambridge), Debbie Spain (IoPPN), Rose Stewart (Oxford), John Suckling (Cambridge), Sally J. Wheelwright (Cambridge), Steven C. Williams (IoPPN), and C. Ellie Wilson (IoPPN).

** These authors contributed equally to the manuscript
6.1 Abstract

Objective: Males are 2 to 5 times more likely to be diagnosed with Autism spectrum disorder (ASD) than females. The reasons for this bias are not fully understood. However, normative diversity in brain structure associated with biological sex is one possible mediator for ASD risk. The authors therefore examined the probability of ASD as a function of sex-related variability in multiple measures of brain anatomy.

Method: 98 adults with ASD (49 males, 49 females) and 98 typically developing controls (51 males, 47 females) underwent structural MRI scanning. Measures of cortical morphometry (i.e. cortical thickness, surface area, the local gyrification index, and sulcal depth) were used to establish normative models of biological sex. These models were subsequently applied to males and females with ASD. Sample probabilities for ASD were calculated across features according to sex-based neuroanatomical phenotypes and extrapolated to estimate ASD risk within the general population.

Results: Across morphometric features, females with ASD displayed a statistically significant neuroanatomical shift towards more male-typical presentations ($p<0.001$). Moreover, within both biological males and females sample probabilities for ASD increased with predicted probabilities for male-typical brain phenotypes. For biological females, this translated to an estimated variability in population prevalence ranging from 0.01 to 2.6% depending on having either a female- or male-typical neurophenotype.

Conclusions: Male-typical neurophenotypes are associated with higher probability for having an ASD diagnosis than female-typical neurophenotypes across a range of neuroanatomical features. In addition to genetic and environmental factors, normative sex-related phenotypic diversity in brain structure should be considered when investigating risk and resilience for ASD.
6.2 Introduction

Autism spectrum disorder (ASD) is a complex neurodevelopmental condition estimated to be 2-5 times more common in males than females (Lai et al., 2015, Fombonne, 2009). While the preponderance of males with ASD has partially been attributed to sex differences in clinical phenotypes (Szatmari et al., 2012, Bölte et al., 2011, Dworzynski et al., 2012, Mandy et al., 2012, Solomon et al., 2012), etiological models propose that factors inherent to normative sex differences may modulate ASD liability (for review see (Werling, 2016)). However, it remains unknown what “male risk” and/or “female protective” factors may contribute to the male preponderance of ASD, and how these factors may affect the biological underpinnings of the condition.

To identify potential sex-related risk (and protective) factors, we recently examined the probability of ASD as a function of normative sex-related phenotypic diversity in brain structure (Ecker et al., 2017). To do so, we initially developed a multivariate predictive model of biological sex based on differential patterns of brain structure in a sample of typically developing (TD) males and females. This model of normative sex differences was subsequently applied to males and females with ASD, which allowed us to (i) examine the probability of ASD along a phenotypic axis, ranging from the characteristic female to male brain phenotype, and (ii) identify patterns of sex-related neuroanatomical variability associated with low and high probability of ASD. In this previous study we reported that normative sex-related phenotypic diversity in brain structure significantly affects the risk of ASD in addition to biological sex alone, with male-typical neuroanatomical characteristics carrying a higher intrinsic risk for ASD than female-typical characteristics. This increased risk was predominantly driven by a phenotypic shift of the brain in females with ASD for a male-typical rather than female-typical presentation. Moreover, using a mapping approach that enabled us to estimate patterns of atypically at the level of each individual subject (Marquand et al., 2014), we demonstrated that high and low risk of ASD is associated with particular patterns of neuroanatomical variability that included many of the brain regions where males typically differ from females on average (e.g. (Sowell et al., 2007, Im et al., 2006)). Taken together, these findings highlight the need for considering normative sex-related phenotypic diversity in addition to genetic and environmental factors when estimating an individual’s risk for ASD.
However, in this prior ‘proof-of-concept’ study, our analysis was based exclusively on measures of cortical thickness (CT) as previous neuroimaging studies have shown CT to be highly variable between TD males and females (Sowell et al., 2007, Im et al., 2006), and significantly altered in individuals with ASD (Wallace et al., 2010, Ecker et al., 2013). Thus, as has been raised by others (Cahill, 2017), it is unknown whether or how our initial findings generalize to other morphometric features that have been reported to be atypical in ASD. These include vertex-wise estimates of surface area (SA; (Ecker et al., 2013)), the local gyrification index (IGI; (Ecker et al., 2016, Schaer et al., 2013)), and sulcal depth (SULC; (Nordahl et al., 2007)). Here, we extended these findings to consider multiple aspects of brain anatomy including both volumetric and geometric cortical features in order to (i) provide a more comprehensive characterization of normative sex related differences in brain structure, (ii) determine how diversity in sex based neurophenotypes are associated with ASD probability, and (iii) estimate how much ASD prevalence in the general population might vary as a function of normative sex-related phenotypic diversity. We expect that our initial finding of a phenotypic shift of the brain in ASD females towards a more male-typical neurophenotype generalizes to other morphometric features, and for sex differences in cortical morphometry to be associated with population risk for ASD.

6.3 Methods

6.3.1 Participants
98 right-handed adults with ASD (49 males and 49 females) and 98 age and IQ matched TD controls (51 males and 47 females) aged 18-42 years were recruited by advertisement and assessed at the Institute of Psychiatry, Psychology and Neuroscience (IoPPN), London, and the Autism Research Centre, Cambridge. Approximately equal ratios of cases to controls, and males to females, were recruited within sites (Table 6.1). Exclusion criteria included a history of major psychiatric
disorder (e.g. psychosis), head injury, genetic disorder associated with ASD (e.g. fragile-X syndrome, tuberous sclerosis), or any other medical condition affecting brain function (e.g. epilepsy). We also excluded participants on antipsychotic medication, mood stabilizers or benzodiazepines.
<table>
<thead>
<tr>
<th></th>
<th>ASD (n = 98)</th>
<th>TD Control (n = 98)</th>
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<tr>
<td>Age [years]</td>
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<td>110.51 ± 13.01 (83 - 137)</td>
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<td>Performance IQ</td>
<td>112.00 ± 13.65 (85 - 138)</td>
<td>116.74 ± 10.90 (93 - 133)</td>
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<tr>
<td>ADI-R Social</td>
<td>17.57 ± 5.50 (10 - 28)</td>
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<tr>
<td>ADI-R Communication</td>
<td>13.86 ± 4.16 (8 - 24)</td>
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<tr>
<td>ADI-R Repetitive Behavior</td>
<td>4.96 ± 2.37 (1 - 10)</td>
<td>-</td>
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<tr>
<td>ADOS Social + Communication</td>
<td>9.43 ± 4.39 (1 - 21)</td>
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<tr>
<td>ADOS Stereotypic Behaviour</td>
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<td>2.35 ± 0.10</td>
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<tr>
<td>Mean Surface Area</td>
<td>0.7680 ± 0.71</td>
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<tr>
<td>Mean local Gyrification Index</td>
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<tr>
<td>Mean Sulcal Depth</td>
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<td>0.0356 ± 0.006</td>
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<tr>
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<td>Total Intracranial Volume [litre]</td>
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<td><strong>Female Participants (n = 96)</strong></td>
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<tr>
<td>Age [years]</td>
<td>27.60 ± 7.12 (18 - 48)</td>
<td>27.60 ± 7.31 (19 - 52)</td>
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<tr>
<td>Verbal IQ$^a$</td>
<td>116.60 ± 12.15 (76 - 144)</td>
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<td>Performance IQ$^a$</td>
<td>110.56 ± 14.50 (67 - 138)</td>
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<td>Full-scale IQ (WASI)</td>
<td>114.88 ± 12.36 (84 - 136)</td>
<td>118.38 ± 7.32 (99 - 129)</td>
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<tr>
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<td>0.6865 ± 0.06104</td>
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<td>Total Intracranial Volume [litre]</td>
<td>1.32 ± 0.18</td>
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</table>

Table 6.1 Participant Demographics and Global Brain Measures: Data expressed as mean ± standard deviation (range). There were no significant between-group differences in age, FSIQ, mean cortical thickness, total grey matter volume, or total intracranial volume (p<0.05, two-sided). ADI-R values based on 49 males and 44 females with ASD. ADOS values based on 49 males and 48 females. * statistically significant between males and females based on p<0.05, two-sided (t(95)=2.23, p=0.028). $^a$ verbal and performance IQ data was not available for one ASD female.
All participants with ASD were diagnosed according to International Statistical Classification of Diseases, 10th Revision (ICD-10) research criteria. Clinical diagnoses were confirmed using the Autism Diagnostic Interview Revised (ADI-R; (Lord et al., 1994)(19)) or the Autism Diagnostic Observation Schedule (ADOS; (Lord et al., 2000)). All ADI-R diagnosed participants reached algorithm cut-offs in the three domains of the ADI-R, although failure to reach cut-off in one domain by maximally two points was permitted. Due to reliable informants being unavailable, we were unable to obtain ADI-R diagnostic data from five females with ASD, for whom diagnostic status was confirmed using ADOS-G cut-offs for ASD. In all other participants, ADOS-G scores were used to assess the severity of current symptoms and were not used as an inclusion criterion. One female with ASD fell short one point on the ADI-R communication and repetitive domain but met ADOS-G criteria. Overall intellectual ability was assessed using the Wechsler Abbreviated Scale of Intelligence (Wechsler, 1999)(21). All participants had a full-scale IQ greater than 80 and gave informed written consent in accordance with ethics approval by the National Research Ethics Committee, Suffolk, UK.

6.3.2 Structural MRI Data Acquisition
Scanning took place at the IoPPN, London, and the Addenbrooke’s Hospital, Cambridge, using matched 3T GE Signa systems (General-Electric, Milwaukee, USA). A quantitative T1-mapping sequence was used to ensure standardization of structural MRI scans across scanner platforms (Deoni et al., 2008, Ecker et al., 2012), which resulted in high-resolution structural T1-weighted inversion-recovery images, with 1x1x1mm resolution, a 256x256x176 matrix, TR=1800ms, TI=50ms, FA=20”, and FOV=25cm.

6.3.3 Cortical Reconstruction using FreeSurfer
FreeSurfer v5.3.0 software (http://surfer.nmr.mgh.harvard.edu/) was used to derive tessellated models of the cortical surface for each T1-weighted image. These well-validated and fully automated procedures have been extensively described elsewhere (e.g. (Dale et al., 1999, Fischl et al., 1999, Ségonne et al., 2004, Jovicich et al., 2006, Fischl and Dale, 2000)). Resulting surface models were visually inspected for
reconstruction errors. Surface reconstructions with visible inaccuracies were further excluded from the statistical analysis and are not described in this study. Dropout rates were equal across groups and were <10% of the total sample.

From each individual’s surface reconstructions four separate vertex based morphometric cortical features were calculated. (i) Cortical thickness (CT), i.e. the closest distance between corresponding vertices between the grey-white matter and gray matter-cerebrospinal fluid boundaries (Fischl and Dale, 2000); (ii) vertex-wise estimates of pial (i.e. grey matter) surface area (SA) (Winkler et al., 2012). And measures of cortical geometry including (iii) the local gyrification index (lGI) (Schaer et al., 2008) and (iv) sulcal depth (SULC, i.e. average convexity, the depth/height of each vertex above the average cortical surface). Prior to further analysis all features (excluding lGI) were smoothed using a 15mm surface-based FWHM Gaussian kernel. As lGI utilizes SA values sampled within spherical regions on interest with a 15mm diameter, no additional smoothing was performed for this measure.

6.3.4 Gaussian Process Classification

Gaussian Process Classification (GPC) (Rasmussen and Williams, 2006, Marquand et al., 2010) models were employed to discriminate biological sex among TD participants on the basis of each of the four cortical features (i.e. CT, SA, lGI, SULC) separately. These classifiers provide a predictive class probability ($y_p$) for each individual ranging from 0 to 1 for the characteristic female to male brain phenotype, with $y_p=0.5$ serving as the binary cut-off between classes. The overall classification accuracy of each model was estimated using leave-one-out cross validation and tested for significance by repeating the cross-validation procedure $n=1,000$ times after randomly permuting the class labels. Subsequently, we applied these four classifiers for biological sex in TD participants to predict the sex of males and females with ASD. The resulting classification accuracies for the ASD cases were tested for statistical significance using permutation testing as outlined above.

To visualize the patterns of brain regions representative of normative sex differences, we present spatial representations of the maximum a-posteriori estimate of each model’s weight vector ($w$), which is indicative of the regional contribution to the overall prediction of biological sex (Marquand et al., 2010). For each classifier, $w$ was
derived by training the GPC on all TD individuals. As individuals with ASD were not included in the initial training of the classifiers they do not contribute to $w$, thus we used a “predictive mapping” approach (Marquand et al., 2014) to determine which cortical features are driving sex related differences in ASD among groups of males and females with high and low probability for ASD. Here, groups of high and low ASD probability were determined according to predicted class probabilities, with $y_p > 0.5$ (i.e. male-typical phenotype) representing high and $y_p < 0.5$ (i.e. female-typical phenotype) representing low probability. For each test example $i$ in both male (and female) high (and low) probability groups $j$, we calculated the element-wise (i.e. Hadamard) product of each individual's data ($x_{ij}$) and the corresponding weights ($w$), so that $(x_{ij}) \ast w$ (Marquand et al., 2014). For each group (i.e. male and female, high and low ASD probability) brain regions contributing the most to the classification were summarized by taking the mean of the $(x_{ij}) \ast w$ values at each vertex.

### 6.3.5 Estimating ASD Probability

To examine how normative sex based phenotypic diversity in brain structure is associated with probability for ASD we first calculated the mean of each individual’s predictive class probabilities across the four classification models. This provided a single class probability for each individual that takes into account normative sex variability across all four cortical features. We then converted the continuous axis of class probabilities into a set of discrete bins (e.g., from 0 to 1 in steps of 0.125). Within each bin, the sample probability of ASD was calculated as the ratio of the number of individuals with ASD relative to the total number of individuals within the bin. Confidence intervals for these sample probabilities were determined using an exact binomial test implemented in R Project for Statistical Computing (https://www.r-project.org). Furthermore, these data allowed us to estimate the population prevalence of ASD given a male- or female-typical neuroanatomical phenotype in addition to being biologically male or female. Details of these methods are provided in the supplementary materials. In brief, using Bayes theorem, we combined our sample prevalence estimates with previously published prevalence rates of ASD for biological male and female individuals in the general population (i.e., 1:42 for male and 1:189 for female individuals; (Baio, 2014)). In this way, we were able to estimate the population prevalence of ASD diagnosis ($D = 1$) given an
individual’s predicted neuroanatomical sex phenotype ($M$) and biological sex ($S$), that is $\mathbb{P}(D = 1 | M, S)$.

6.3.6 **Vertex-wise between-group comparison of cortical measures**

Vertex-wise statistical analyses of each cortical feature were conducted using the SurfStat toolbox (www.math.mcgill.ca/keith/surfstat/) for MATLAB (R2014a, The Mathworks, Massachusetts). For the comparisons between TD males and females, parameter estimates were obtained by regression of a GLM at each vertex $i$ with biological sex and center as categorical fixed-effects factors, and total grey matter (GM) volume as a continuous covariate. To examine whether the neuroanatomy of ASD is significantly modulated by biological sex, we also included a main effect of diagnostic group and a group-by-sex interaction, i.e.

$$ Y_i = \beta_0 + \beta_1 \text{Group} + \beta_2 \text{Sex} + \beta_3 (\text{Group} \times \text{Sex}) + \beta_4 \text{GM_{total}} + \beta_5 \text{Center} + \epsilon_i, $$

where $\epsilon_i$ is the residual error. Additionally, total SA was used as a continuous covariate for analyses of SA. Effects of interest were estimated from the coefficients $\beta_{1,3}$ normalized by the corresponding standard error. Corrections for multiple comparisons were performed using a ‘random field theory’ (RFT)-based cluster analysis for non-isotropic images at a cluster-threshold of $p<0.05$ (two-tailed) (Worsley et al., 1999).

6.4 **Results**

6.4.1 **Participant Demographics and Global Brain Measures**

There were no significant differences between all individuals with ASD and TD controls in age ($t(194)=-0.53$, $p=0.598$), full-scale IQ ($t(194)=-1.72$, $p=0.086$), mean CT ($t(194)=0.08$, $p=0.936$), mean SA ($t(194)=0.03$, $p=0.976$), mean lGI ($t(194)=0.24$, $p=0.810$), mean SULC ($t(194)=0.67$, $p=0.499$), and total GM volume ($t(194) = -0.20$, $p=0.839$). There were also no significant differences between males and females in age ($t(194)=-0.93$, $p=0.356$), full-scale IQ ($t(194)=-1.87$, $p=0.063$), mean CT ($t(194)=0.95$, $p=0.344$), or mean SULC ($t(194)=-2.80$, $p=0.006$) across diagnostic groups. As expected, total grey matter volume was significantly larger in males than in females ($t(194)=9.11$, $p=0.001$). Furthermore, across the entire sample males
showed significantly greater mean SA ($t(194)=9.43, p<0.001$) and mean lGI ($t(194)=5.56, p<0.001$) than females. There were no significant differences in any of these measures between TD and ASD males, or between TD and ASD females ($p<0.05$, 2-tailed).

6.4.2 Prediction of Biological Sex Based on Normative Variability in Cortical Morphometry

Across all four vertex based cortical features GPC was able to separate male from female TD controls at cross validated accuracies significantly above what would be expected by chance (SA 78%, $p<0.001$; CT 71%, $p<0.001$; SULC 74%, $p<0.001$; lGI 77%, $p<0.001$). Using each classifiers predictive class probabilities for individuals we were able to represent normative sex based variability in brain phenotypes along a singular phenotypic axis, ranging from the female- to male-typical neurophenotype (Figure 1). For each morphological measure, the GPC model weights ($w$) followed a spatial pattern similar to the distribution of group differences observed between TD males and females as determined via a conventional GLM (Figure 6.1, Supplementary Figure 1, Supplementary Table 1).
Figure 6.1 Gaussian Process Classification of Biological Sex: The results of Gaussian Process Classification (GPC) according to biological sex categories are shown for each of the four cortical features investigated: (A) cortical thickness (CT), (B) surface area (SA), (C) the local gyriﬁcation index (lGI), and (D) sulcal depth (SULC). Each model was able to separate males from females at accuracies signiﬁcantly above what would be expected by chance (top left). When these models of normative sex differences in cortical morphology were applied to males and females with ASD, females with ASD were more frequently classified to the male class (bottom left) indicating a male-typic morphological presentation among females with ASD. Spatial representations of each discriminative model (i.e. weight vectors) are shown (right). These patterns largely corresponded with brain regions that differ between TD males and females as identiﬁed using a conventional general linear model (see supplementary Figure 1).
6.4.3 Prediction of Biological Sex Among Individuals with ASD

When predicting biological sex for individuals with ASD we found that females with ASD were more frequently allocated to the male rather than the female class in all four models (SA 71%, $p=0.002$; CT 80%, $p=0.002$; SULC 65%, $p=0.009$; lGI 78%, $p=0.001$). Females with ASD were significantly allocated to the male class more frequently than would be expected based on the typical rates of misclassification for female controls (SA $\chi^2=26.41$, $p<0.001$; CT $\chi^2=22.15$, $p<0.001$; SULC $\chi^2=13.65$, $p<0.001$; lGI $\chi^2=26.01$, $p<0.001$). Furthermore, the mean across the predicted class probabilities for ASD females was significantly increased in females with ASD relative to female controls, indicating a shift towards male-typical morphological patterns in the brain (SA $t=6.158$, $p<0.001$; CT $t=6.13$, $p<0.001$; SULC $t=4.44$, $p<0.001$; lGI $t=6.85$, $p<0.001$). Across all four models there were no significant differences ($p<0.05$) between the predicted class probabilities for TD and ASD males, or between males and females with ASD.

6.4.4 ASD Probability as a Function of Normative Sex Variability in Cortical Morphometry

Within our sample we found that the number of individuals with ASD significantly increased along the female to male phenotypic axis (i.e. from female to male) as defined by predicted class probabilities. This effect was predominantly driven by the shift of females with ASD towards a more male-typical neuroanatomical brain phenotype (See Figure 6.2, Supplementary Figure 2). When these sample probabilities were combined with previously reported ASD prevalence rates in the general population among males (1/43, i.e. 2.3%) and females (1/190, i.e. 0.5%) (Baio, 2014), we found that biological females with male-typical brain morphology (0.1% estimated prevalence) were twenty six times more likely to have ASD compared to biological females with female-typical neuroanatomical phenotypes (2.6% estimated prevalence). Biological males with male-typical brain morphometry showed an estimated population prevalence of 2.4% in line with previously reported rates. However, males with female-typical phenotypes showed a reduced prevalence estimate for ASD (1.9%) indicative of a ‘female protective effect’ in relation to brain anatomy (see Supplementary Table 2).
Predictive maps identifying patterns of neuroanatomical variability associated with high and low probability of ASD across sexes are shown in Figure 6.3. While a similar set of brain regions conferred high (and low) probability for ASD in men and women within features, certain brain regions were unique to or differed in sign between particular risk groups and sexes. For example, negative predictive mapping values for CT in the left parahippocampal gyrus and SULC in the superior temporal gyrus were observed across all males and high-probability females, while low probability females showed positive values in these regions. For predictive mapping measures of SA and lGI, regions of the inferior and superior temporal gyri appear to show a sex-specific relationship with high probability males showing negative values compared to positive values amongst females and low probability males. This indicates that morphometric measures in particular brain regions that are important for classifying sex differences differ between individuals with high and low risk for ASD and may be sex-specific.
Figure 6.3 Prediction Maps for High and Low ASD Probability Males and Females: Prediction maps highlighting brain regions most relevant to classifying biological sex among high and low risk males and females are shown for each cortical feature, (A) cortical thickness (CT), (B) surface area (SA), (C) local gyrification index (lGI), and (D) sulcal depth (SULC). For male and female individuals high and low probability groups were determined according to class probabilities with <0.5 (i.e. female-typic) indicating low and >0.5 (i.e. male-typic) indicating high probability. For each subgroup the product of the decision function for classifying biological sex ($w$) and each individual's data ($x_i$) was calculated. Mean values at each vertex were then used to summarize maps within subgroups.
6.4.5 Cortical Morphometry in ASD is Modulated by Biological Sex

Lastly, we used a conventional GLM to test for significant between-group differences and diagnosis-by-group interactions in the four examined morphometric features. Overall, we found that measures of CT, SA, and GI in ASD were significantly modulated by biological sex (Figure 6.4, also Supplementary Figure 3). Such group-by-sex interactions therefore need to be taken into account when interpreting main effects of group. Significant group-by-sex interactions in measures of CT were observed across the temporal lobe, including the fusiform and parahippocampal gyri. Across this cluster the degree of CT abnormality in females with ASD exceeded those of males, despite no differences in measures of ASD symptom severity (9). Additionally, group-by-sex interactions were observed for measures of SA and GI in the medial orbital frontal cortex showing decreases in females with ASD and increases in males with ASD. Statistical details of these clusters are provided in Supplementary Table 3.
Figure 6.4 Group by Sex Interaction Effects in Cortical Morphometry: Clusters of significant (RFT p<0.5) group x sex interactions in (A) cortical thickness (CT), (B) surface area (SA), and (C) the local gyrification index (lGI). No significant group x sex interactions were observed in measures of sulcal depth (SULC). Statistical details of these clusters are provided in Supplementary Table 3.
6.5 Discussion

The present study builds upon a previous ‘proof-of-concept’ suggesting that probability of ASD increases significantly from a characteristic female to male neuroanatomical brain phenotype (Ecker et al., 2017). In this prior study, which examined measures of cortical thickness (CT) only, the increase in ASD probability was predominantly driven by a phenotypic shift of the brain in females with ASD towards a male-typical neurophenotype (Ecker et al., 2017). In the current study, we show that this phenotypic shift is not restricted to measures of CT, but generalizes to other cortical features including vertex-wise estimates of surface area (SA), the local gyrification index (lGI), and sulcal depth (SULC). The findings of the present study therefore not only support our original report in suggesting that male-typical neuroanatomical characteristics carry a higher probability for ASD than female-typical characteristics, but further show this effect to be broadly and consistently related across multiple measures of cortical morphometry.

The extension of our approach to include multiple morphometric features is of crucial importance as there is evidence to suggest that different cortical features are etiologically and phenotypically distinct. For example, it has been shown that measures of CT and SA (i) are mediated by different sets of genes (Panizzon et al., 2009), (ii) have different developmental trajectories (Ecker et al., 2014), and (iii) represent distinct aspects of the cortical architecture (Rakic, 1995). Therefore, the observation that the phenotypic shift of the brain in ASD females is consistent across morphometric features may indicate the existence of a common underlying mechanism, such as atypical cortical neurogenesis and migration (Casanova, 2014), that steers multiple aspects of brain anatomy towards a similar phenotypic endpoint overall. Yet, the neuroanatomical patterns associated with the prediction of biological sex, and the patterns carrying high/low ASD probability in males and females, differed in sign and regional distribution between the cortical features. Thus, while there may be a common etiology for the phenotypic differences in brain anatomy in ASD within the sexes, our findings are consistent with the notion that sex differentiation mechanisms in the brain have region specific effects (for review see (McCarthy, 2017, McCarthy and Konkle, 2005)). Therefore, differences in these neuroanatomical patterns associated with high probability for ASD may be beneficial.
in identifying brain regions where sexual differentiation may contribute to male preponderance in ASD.

Moreover, we established that the neuroanatomy of ASD in some regions of the brain is modulated by biological sex by examining sex-by-diagnosis interactions for multiple cortical features. This suggests that in these brain regions (i) there are not only quantitative differences in the degree of neuroanatomical abnormality between males and females with ASD and their respective normative population, but also that (ii) ASD potentially interacts with sexually-differential neurobiological phenotypes to give rise to qualitative differences in brain anatomy between males and females with the condition. This is also relevant in view of a recent genetic study, suggesting that while ASD risk genes are expressed at similar levels in males and females, the impact of these genes is likely modulated by their interactions with sex-differential processes (e.g. in cortical microglia and/or astrocytes) that may play a critical role in setting males at a greater risk for ASD (Werling et al., 2016). An individual’s risk or probability of ASD is therefore not only dependent on genetic load itself, but also on naturally occurring sex-differential biological pathways.

With respect a biological organ as complex as the brain it has been suggested that on a phenotypic level, the brain should be considered a ‘mosaic’ of regions that differ between degrees of relative maleness or femaleness, or not at all (i.e. unisex), resulting in a potentially infinite variety of brain phenotypes both within and between sexes (McCarthy, 2017, McCarthy and Konkle, 2005). In order to identify an individual’s position on such a multi-dimensional ‘spectrum’ of sex-related phenotypic variability, it is important to develop analytical frameworks that can accommodate inter-individual phenotypic diversity within and across the binary categories dictated by biological sex, such as the probabilistic multivariate pattern classification approach utilized in the present study (see also (Ecker et al., 2017)). In this way we were able to estimate an individual’s risk of ASD based on phenotypic, rather than biological sex and in doing so demonstrated that an individual’s probability for developing ASD may vary from as much as 0.1 to 2.6% in the general population depending on having a female- or male-typical neuroanatomical
phenotype, in addition to being biologically male or female. Our approach to modeling normative sex-related phenotypic diversity may therefore be extended to various additional phenotypic features in the future, and to contribute to the development of multivariate disease liability models for complex neurobiological phenotypes.

Despite the advantages of the current approach over the original proof-of-concept study, a number of limitations remain. Most importantly, it is crucial to highlight that while we were able to show a statistical association between normative sex-related phenotypic diversity in brain structure and ASD probability, this relationship should not be interpreted as being ‘causal’. Future studies will be needed to uncover the genetic and/or neurobiological mechanisms underpinning this relationship (e.g. (Baron-Cohen et al., 2015, Werling et al., 2016, Lai et al., 2017)). Second, due to the limited range of clinical/behavioral measures acquired in the current study, we were unable to determine the functional relevance of our findings. In the future it will be crucial to identify how diversity in normative sex-related morphometric phenotypes relates to sex-differences in general cognitive and/or behavioral profiles (Miller and Halpern, 2014, Hyde, 2016), or to different clinical ASD phenotypes (Szatmari et al., 2012, Bölte et al., 2011, Solomon et al., 2012, Mandy et al., 2012). Last, our study was based on a sample of adults with ASD without learning disability using a cross-sectional study design. The fact that our sample is not representative of the entire ASD population limits the generalizability of our sample probability estimates for ASD to the general population. Furthermore, our population estimations are based on previously reports of male preponderance in ASD being roughly a 4:1 male to female ratio (Baio, 2014), however other studies have shown this ratio may actually to be closer to 2:1 (Werling, 2016). The nature of our sample means further studies will thus be required to (i) replicate our findings in other subgroups on the autism spectrum (e.g. in children and adolescents, or individuals with a learning disability), and (ii) to identify how stable (or variable) our neuroanatomical indicators of biological sex are across different stages of development.
6.6 Conclusions
Taken together, our findings suggest that the male-typical neurobiological brain phenotype carries a higher probability for ASD than the female-typical phenotype across the binary categories dictated by biological sex, and across a range of neuroanatomical features that include both volumetric as well as geometric measures. In addition to genetic and environmental factors, multiple morphometric features characteristic of normative phenotypic diversity in brain structure should therefore be considered when investigating risk and resilience for ASD.
6.7 References


comparing brain cortical surface area and other areal quantities. 
*NeuroImage*, 61, 1428-1443.


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Chapter 7: General Discussion

The studies that comprise this thesis employed structural magnetic resonance imaging (MRI) and novel analytical frameworks to address three overarching aims, namely i) developing novel neuroimaging features that are informative (e.g. of diagnostic or prognostic value) for the condition, ii) evaluating the efficacy of structural neuroimaging features to identify ASD cases within the clinical setting, and iii) investigating the role of normative sex differences in brain structure and their influence on the probability for ASD. Collectively these studies share the broader goal of advancing the field of neuroimaging research in a direction that provides real world translational value for ASD individuals and stakeholders. First, the present chapter aims to summarize the key findings of each of the four studies that were presented in this thesis. Second, possible interpretations of these results are provided, particularly with regards to moving neuroimaging studies in ASD closer to the development of translational imaging tools. Last, strengths and limitations of the presented studies will be discussed, followed by final conclusions.

7.1 Summary and Key Findings

7.1.1 Study 1: Tissue Contrast as a Novel Imaging Marker in Autism Spectrum Disorder

The aim of this study was to determine if previous post-mortem reports of poor definition of the grey–white matter boundary in autism spectrum disorder (ASD) (Avino and Hutsler, 2010) could be detected using a whole brain in vivo MRI approach. To accomplish this, we used a novel application of tissue signal intensity contrast measures sampled at and around the grey-white matter boundary (Salat et al., 2009, Andrews et al., 2017). As hypothesized, we determined that individuals with ASD had significantly less well-defined grey-white matter percent contrast (GWPC) across the grey-white matter boundary. This effect appeared to be largely driven by an increase in grey matter signal intensities in individuals with ASD. Taken together, our findings suggest that measures of GWPC sampled across cortical layers may serve as an in vivo proxy measure for irregular microstructural organization of the cortex in ASD.
7.1.2 Study 2: Clinical Validation of Multivariate Pattern Classification Models for Autism Spectrum Disorder

This study sought to first establish predictive models for the broader ASD phenotype based on measures of cortical neuroanatomy within a well-defined sample consisting of typically developing (TD) and ASD male adults acquired in the research setting. The second and primary aim of this study was to test the efficacy of these classification models to identify individuals with ASD within an independent clinical sample. Two different well established pattern classification algorithms, namely the support vector machine (SVM; Vapnik, 1995) and Gaussian process classifier (GPC; Rasmussen and Williams, 2006, Marquand et al., 2010), were used to establish classifiers based on the research sample utilizing measures of either cortical volume, thickness (CT), surface area (SA), the local gyrification index (lGI), sulcal depth (SULC), or grey-white matter percent contrast (GWPC). Based on previously reported findings (Ecker et al., 2010, Jiao et al., 2010) we hypothesized that these structural measures are able to discriminate individuals with ASD and TD controls significantly above chance levels. However, given differences in inclusion criteria between our research training and clinical testing samples, and based on previous findings of classification models showing reduced performance in the range of 10-15% when applied to independent testing sets (Sabuncu and Konukoglu, 2014), we expected a reduction in overall accuracy when predicting ASD diagnosis within the clinical sample.

While some classifiers showed modest classification accuracies (61%) within the research sample, we were not able to establish the statistical significance of these classifiers due to the small size of our training set. However, in an exploratory analysis we applied our best performing models to predict ASD diagnoses within the independent clinical sample. We established that the majority of these models showed decreased overall classification accuracies within the clinical testing set. One notable exception to this was the GPC utilizing measures of grey-white matter percent contrast (GWPC) sampled at the grey-white matter boundary (Andrews et al. 2017), which showed high overall accuracy within the clinical sample (81%). However, given this model’s initial performance in the research setting this result must be considered as preliminary at best.
7.1.3 Studies 3 and 4: Association of Normative Sex-Related Phenotypic Diversity in Brain Structure with Probability for Autism Spectrum Disorder

These two studies sought to directly investigate possible biological underpinnings of male preponderance in ASD (Lai et al., 2017, Werling, 2016). The aims of these studies were three-fold; i) to provide a phenotypic characterization of normative sex related diversity in brain structure, ii) to determine how diversity in sex based neurophenotypes is associated with probability for ASD, and iii) to estimate variability in ASD prevalence in the general population as a function of normative sex-related phenotypic diversity in brain structure. To accomplish this, we first established normative predictive models in male and female TD individuals for the binary classes dictated by biological sex. In study 3, we report initial findings using measures of CT (Ecker et al., 2017), and in study 4 we extended the analytical framework to include measures of SA, IGI, and SULC. The use of GPC (Rasmussen and Williams, 2006, Marquand et al., 2010) allowed us to effectively represent the diversity of sex based neurophenotypes along a singular axis dictated by predictive class probabilities ranging from the female- to male-typical neurophenotype. Once established, these normative models were applied to males and females with ASD.

In both studies we found that sample probabilities for ASD increased with predicative probabilities for the male-typical class. This was largely due to a statistically significant shift towards more male-typical neurophenotypes in females with ASD across all cortical measures. We then extrapolated sample probabilities for ASD among individuals with female- and male-typical neurophenotypes to estimate their association with ASD prevalence within the general population. Based on previous population prevalence estimates (Baio, 2014), we demonstrate that biological females with more male-typical brain morphology (2.6% estimated prevalence) are 26 times more likely to have ASD compared to biological females with female-typical neuroanatomical phenotypes (0.1% estimated prevalence). Biological males with male-typical brain morphometry showed an estimated population prevalence of 2.4%, which is in line with previously reported rates. In addition, males with female-typical phenotypes showed a reduced prevalence estimate for ASD (1.9%). Taken
together, these results support the notion that there is differential liability for ASD between the sexes, which increases from a female to male phenotypic presentation of the brain, and is also indicative of a ‘female protective effect’ for ASD in terms of brain anatomy.

7.2 Discussion

7.2.1 The Development of Imaging Features for Autism Spectrum Disorders

The past decade has seen several advances in the development of structural MRI features to study the neuroanatomical underpinnings of ASD. Studies continue to move away from investigating relatively unspecific, gross-anatomical measures, such as total brain volume, to using more advanced metrics that represent particular aspects of neuroanatomical structure and development (Ecker et al., 2015). For example, surface based morphometry (SBM) measures of CT and SA have been shown to be i) mediated by different sets of genes (Panizzon et al., 2009), ii) have different developmental trajectories (Ecker et al., 2014), and iii) represent distinct aspects of the cortical architecture (Rakic, 1995). Thus, these measures are well suited to explore etiological theories for ASD, which suggest that the condition arises from atypical neuronal proliferation, migration, and maturation (Casanova, 2014). The development of novel neuroimaging features that map onto specific aspects of the ASD phenotype thus offers the potential for both greater insight into the biological underpinnings of ASD, and the possibility for such features to act as imaging biomarkers.

One neuropathological finding that remained relatively underexplored in vivo is atypical cell patterning at the grey-white matter boundary in ASD (Avino and Hutsler, 2010). The current thesis presented a novel in vivo approach to quantify this specific neuroanatomical feature (Andrews et al., 2017). This was done with the purpose of investigating a critical aspect of neurodevelopment believed to contribute to poor grey-white matter boundary integrity in ASD, namely the formation and resolution of the cortical subplate (Hoerder-Suabedissen and Molnár, 2015). During neurodevelopment, cells within the subplate are critical for establishing proper cortical structure and connectivity by guiding both migrating neurons and innervating
cortico-cortico and thalamo-cortical axons (Allendoerfer and Shatz, 1994). Thus, disruptions to subplate development and apoptosis may contribute to both atypical cortical structure and connectivity observed within ASD (Hutsler and Avino, 2015).

The importance of the subplate in establishing inter-cortical connectivity is supported by reports suggesting that this structure is thicker within highly connected cortical association areas (Kostovic and Rakic, 1990). This has recently been linked to the downward dispersion of subplate cells resulting from a high degree of axonal innervation in these regions (Duque et al., 2016). Within the current study, significant increases in grey matter signal intensity in ASD were largely localized to association areas that underpin social processing and wider socio-cognitive functioning, including regions of the temporal lobe, the temporal parietal junction, and fusiform gyrus. Thus, the increased signal intensities in adults with ASD in these regions may reflect the consequences of an atypical formation of cortical connections during early brain development, potentially resulting in increased myelination and hyper local connectivity (Belmonte et al., 2004).

Moreover, our examination of grey-white matter percent tissue contrast (GWPC) at the grey-white matter boundary represents a direct attempt to quantify a specific neuropathological mechanism in ASD (i.e. poor delineation of this boundary). Accordingly, this feature is likely to have greater effects sizes within ASD individuals than other, more general measures of cortical morphometry. This is of key importance for translational studies seeking to use brain imaging features to make prognostic or diagnostic predictions within ASD. For example, in the context of pattern recognition it has been shown that the best predictor of successful classification of ASD is not the particular algorithm used but training models on imaging features with significant effects sizes in the condition (Sabuncu and Konukoglu, 2014). Thus, measures such as GWPC that map onto particular aspects of ASD neuropathology are likely to be valuable to translational research for their potential increased sensitivity to detect the condition.
7.2.2 The Development of Diagnostic Imaging Tools for Autism Spectrum Disorder

Increasingly, through applications of multivariate pattern classification, studies have provided proof of concept evidence that individuals with ASD can be identified with high accuracy in the research setting based on neuroimaging features (Ecker et al., 2010, Ingalhalikar et al., 2011, Just et al., 2014). The implementation of predictive models for ASD would offer significant benefits to patients and stakeholders. Such benefits could come through early identification of the condition allowing for intervention and treatment to begin at a younger age. Furthermore, this would provide clinicians biologically based evidence when making a diagnosis of ASD, which currently is reliant solely on behavioral assessments. However, to date no studies have tested classification models for ASD trained in the research environment on independent samples that are representative of the broader clinical population. This is important because inclusion criteria for participation in ASD research are likely to introduce several phenotypic differences between research and clinical ASD cohorts.

Within the current study, our best performing classifiers for ASD in the research setting showed modest overall accuracies (61%). While this is within the range of previous models tested on independent samples (Sabuncu and Konukoglu, 2014, Shen et al., 2017), we were not able to establish the statistical significance of these classifiers due to the small size of our research sample. However, as an exploratory step we applied the best performing classifiers within the training set to predict ASD within the clinical sample. We found a GPC based on measures of grey-white matter percent contrast (GWPC) taken at the grey-white matter boundary (Andrews et al., 2017) to have high overall accuracy within the clinical sample (81%) across all diagnostic sub-categories (74-88%). This preliminary result supports the concept that features developed to map onto particular aspects of ASD neuropathology have an increased sensitivity to detect the condition. However, given that we could not show this model performed significantly better than chance within the research setting ($p=0.197$), this finding must be considered as suggestive only.

A major impediment to classifying individuals with ASD at high accuracy is the highly heterogeneous nature of the phenotype. As a result, maximal classification
accuracy based on brain structure across the ASD population is likely to be modest. Thus, the development of approaches that directly address phenotypic heterogeneity within ASD will be of critical importance for the successful delivery of classification methods that have translational value in clinical settings. Several individual variables including exogenous factors and genetic disposition may contribute to neuroanatomical variance in ASD. One variable that has been shown to have strong associations with ASD is biological sex (Lai et al., 2017, Werling, 2016). Accordingly, a major theme of this thesis dealt with identifying how normative variance in brain structure associated with biological sex is associated with probability for ASD.

7.2.3 Neuro-Diversity Associated with Biological Sex and Probability for Autism Spectrum Disorder

It has been suggested that on a phenotypic level, the brain should be considered a ‘mosaic’ of regions that differ between degrees of relative maleness or femaleness, or not at all (i.e. unisex), resulting in a potentially infinite variety of brain phenotypes both within and between the sexes (McCarthy, 2017). The studies presented in this thesis employed a multivariate classification approach that allowed us to represent diversity in sex based neurophenotypes along a continuum ranging from typically male to female. This ‘more nuanced’ approach to investigating sex differentiation in the brain represents a key advancement as it can accommodate the long-standing notion that both males and females are subject to various degrees of masculinization and feminization (Whalen, 1974).

Through this normative modelling approach we were able to estimate an individual’s probability of ASD based on phenotypic, rather than biological sex. Using this approach, we demonstrated that an individual’s probability for developing ASD can vary from 0.1 to 2.6% in the general population depending on having a female- or male-typical neuroanatomical phenotype, while being biologically male or female. Thus, these phenotypic measures that represent variable degrees of sexual differentiation in individuals, may serve as an important factor in future multivariate disease liability models for complex neurobiological phenotypes such as ASD.
Moreover, we established that the neuroanatomical patterns associated with prediction of biological sex differed in region and sign between the sexes, and individuals with female- and male-typical neurophenotypes. This finding is consistent with the idea that sex-dependent mechanisms have region specific effects in the brain, and that the neuroanatomical patterns associated with high and low probability for ASD may be used to identify brain regions where sexual differentiation contributes to male preponderance in ASD prevalence. Given the significant association between biological sex and ASD prevalence, these findings contribute to ongoing research efforts into the etiological underpinnings of ASD, and potentially other mental health disorders with sex differences in prevalence (Cahill, 2017).

7.3 Strengths of the Current Studies

The studies that comprised this thesis have a number of benefits. First, to the best of our knowledge, we present the first study to investigate grey-white matter boundary integrity in vivo in ASD. In doing so we established a novel imaging feature in the form of percent contrast of grey-white matter signal intensity (GWPC) taken at and around the grey-white matter boundary. Given that GWPC potentially maps onto a specific neuropathological feature in ASD, the measure is likely to be particularly useful to future ASD researchers as it has large effect sizes (i.e. ‘biological footprint’) in the condition in comparison to many other cortical morphometric features. Second, through collaboration with the Autism Assessment and Behavioural Genetics Service at the Bethlem Royal Hospital, South London and Maudsley NHS Foundation Trust, we were able to recruit a highly representative sample of ASD individuals within the clinical setting. Several neuroimaging studies have sought to develop a diagnostic imaging biomarker for ASD within the research setting. However, inherent differences in the clinical phenotype between research and clinical ASD samples are likely to exist (see Chapter 4.1). Therefore, the recruitment of a clinical sample such as the one presented in the current study is of critical importance for testing the translational value of classifiers established in the research setting. Third, this thesis presents the first proof of concept studies linking normative variability in brain structure associated with biological sex to the probability for ASD. A particular strength of these studies was the use of Bayesian pattern recognition methods to effectively represent biological sex along a continuum. This nuanced approach
allowed us to move beyond simple binary sex categories and more fully capture the spectra of neurodiversity associated with differentiation between the sexes (McCarthy, 2017).

7.4 Limitations
Collectively, the presented studies face limitations common to several structural neuroimaging investigations in ASD. The high cost of MRI along with the significant amount of time, energy, and expertise needed to recruit patients to participate in imaging research presents inherent limitations on sample sizes. Furthermore, the nature of MRI acquisition also hinders the collection of samples that accurately reflect the entire spectrum of ASD phenotypes. Often, individuals with particularly severe ASD symptoms and/or comorbid learning disability are unable or unwilling to conform to the requirements that MRI acquisition demands, namely lying motionless within a confined noisy space for an extended period of time. Consequentially, so called ‘low functioning’ individuals with ASD are nearly absent within the neuroimaging literature.

Despite these limitations, however, our studies investigating tissue contrast and the association between biological sex and ASD probability benefit from relatively large sample sizes ($n \sim 100$), which is above what has been shown to be required for the identification of subtle group differences in cortical measures ($n=50$, Pardoe et al., 2013). Furthermore, inclusion of nearly 50 females with ASD represents one of the largest imaging samples of females with the condition reported to date. However, the recruitment of adult samples introduces limitations when studying a neurodevelopmental condition such as ASD that likely originates during early prenatal stages of life (Bauman and Kemper, 2005). The developmental trajectories of the variety of structural neuroimaging measures within this study are yet to be firmly established, but are likely to non-linear and complex (Ecker et al., 2014). Thus, it remains uncertain if, and how, the current results identified within adults will translate to infants, toddlers, adolescents, and elders with ASD. Ultimately, future longitudinal studies that recruit participants during the first stages of life (e.g. fetal and/or neonatal) who are followed up with across the lifespan will be invaluable to the study of ASD.
The current studies utilized state of the art MRI methods for investigating cortical structure in vivo within a clinical population. While these methods offer many improvements over more traditional MRI methods, such as voxel based morphometry (VBM), they are still limited in several ways. The current high end resolution of 3T structural MRI scans is roughly 1mm$^3$. It has been estimated that a 1mm$^3$ volume in the cerebral cortex may contain upwards of 630,000 neuron cells and as many as four times that number of glial cells (Lent et al., 2012). Therefore, the current resolution offered by structural MRI does not allow for the investigation of fine aspects of neuro-biology such as cell patterning of individual cortical layers, and is thus limited to gross volumetric and geometric measures. Accordingly, one of the greatest limitations of structural neuroimaging studies remains a lack of understanding of the underlying biological mechanisms that contribute to imaging measures. For example, atypical CT measures in ASD may reflect gross numbers of neuronal cells within the cortical sheet (Courchesne et al., 2011), degrees of dendritic arborization (Tang et al., 2014), or myelination processes (Sowell et al., 2004). Adding to the complexity, it is likely that each of these variables contribute to CT to different degrees and kind across individuals and developmental stages. Future research is thus required to investigate the specific cytoarchitectural abnormalities underlying ASD.

7.5 Future Directions

7.5.1 Large Scale Studies and Heterogeneity

The use of MRI techniques to investigate ASD is a relatively novel and rapidly growing field. In this context, it is perhaps unsurprising that many of the limitations outlined above remain to be addressed. Ongoing efforts such as the Autism Brain Imaging Database Exchange (ABIDE), Infant Brain Imaging Study (IBIS), Autism Phenome Project (APP) and European Autism Interventions Study (EU-AIMS) are currently recruiting and scanning large numbers of individuals with autism across the full spectrum of impairment, and at the earliest stages of neurodevelopment. The collection of large scale, high quality, imaging data sets that accurately reflect the phenotypic heterogeneity observed in the broader ASD population will be invaluable, and constitute an essential step towards addressing issues surrounding ASD heterogeneity.
Moreover, current efforts to better understand the biological underpinnings of ASD and develop both efficacious treatments and biomarkers for the condition are hindered by the diversity of clinical phenotypes incorporated under ASD diagnostic criteria. Thus, the stratification of the ASD phenotype into biologically more homogeneous subtypes is of critical importance for advancing studies of ASD etiology, and efforts towards developing imaging tools for translational applications. Recent applications of normative models that seek to identify deviations in particular behavioral measures within individuals from a population distribution represent a critical advancement in this area of research (Marquand et al., 2016). These methods appear to be particularly well suited for parsing ASD through their ability to infer how certain atypical behaviors map onto biology and thus identify homogenous biological subtypes in ASD associated with degrees of variation in behavioral measures when compared to a healthy cohort.

### 7.5.2 Development and Implementation of Multi-Modal Methods

Increasingly, neuroimaging studies of ASD incorporate several different imaging features and modalities. Such multimodal imaging studies are extremely valuable because different imaging sequences (e.g. diffusion, functional, structural) can be used to capture unique aspects of neuropathology in ASD. However, standard practice for multimodal studies still involves investigating features in a parallel (i.e. unimodal) fashion. Thus, interpretation of the relationship between results in different modalities is conducted in a post-hoc manner. In order to determine the inter-relationships between different imaging measures, and to take full advantage of the unique strengths that each modality offers, so-called multimodal ‘fusion methods’ are required. For example, linked independent component analysis can be used to identify components comprised of shared variance across multiple imaging features, regardless of differences in spatial dimensions between them (Groves et al., 2012). Such methods have already shown the ability to identify multimodal components associated with particular behaviors associated with ADD (Francx et al., 2016), and thus may offer value in efforts to parse the heterogeneity of ASD.

### 7.5.3 Linking Imaging with Biology

Lastly, the development of surface based morphometry measures that represent distinct aspects of cortical architecture and etiology has provided unique insights into
the potential underlying biological mechanisms driving differences in these measures in individuals with ASD. However, there is a critical need for establishing concrete links between macroscopic imaging measures and microscopic biology within individuals. Recent advancements in the use of cellular models based on patient-specific induced pluripotent stem cells (iPSCs) offer particular promise to address this relationship. Through a typical biopsy (e.g. skin, hair etc.), this technology allows for the delivery of in vitro neuron cells from particular patients (Takahashi and Yamanaka, 2006, Takahashi et al., 2007). Accordingly, many of the developmental events believed to be atypical in ASD can be recapitulated and observed in the culture dish (Marchetto and Gage, 2012) and then related to neuroimaging measures, all within the same individual in a safe, in vivo, minimally invasive fashion.

7.5 Conclusion
The studies that comprise this thesis highlight advances in the field of structural neuroimaging research in autism spectrum disorder (ASD) in areas of feature development, clinical translation, and efforts to understand the modulating role of biological sex on the prevalence of ASD. Here we first introduced a novel in vivo feature based on tissue contrast to measure atypical cortical microstructure that has previously been reported in histological studies. Second, we trained classification models for ASD in the research setting and then tested the ability of these models to make diagnostic predictions within a representative clinical ASD sample. Last, we established normative models of phenotypic diversity in brain structure associated with biological sex in a sample of typically developing (TD) males and females, which we subsequently applied to males and females with ASD to identify the association between sex based neurophenotypes and ASD probability. Taken together, the body of work presented within this thesis constitutes an important step forward toward establishing translational imaging tools for ASD that may one day be applied in the clinical setting. However, future studies will be required to address the large degree of phenotypic heterogeneity in ASD, develop methods that can take full advantage of multimodal imaging data, and identify the biological underpinnings of macroscopic neuroimaging measures.
7.6 References


thickness and surface area in autism spectrum disorder. *Journal of Neural Transmission*, 121, 1157-1170.


Appendix I: ICD-10 Criteria for "Childhood Autism"

A. Abnormal or impaired development is evident before the age of 3 years in at least one of the following areas:

1. receptive or expressive language as used in social communication;
2. the development of selective social attachments or of reciprocal social interaction;
3. functional or symbolic play.

B. A total of at least six symptoms from (1), (2) and (3) must be present, with at least two from (1) and at least one from each of (2) and (3)

1. Qualitative impairment in social interaction are manifest in at least two of the following areas:
   a. failure adequately to use eye-to-eye gaze, facial expression, body postures, and gestures to regulate social interaction;
   b. failure to develop (in a manner appropriate to mental age, and despite ample opportunities) peer relationships that involve a mutual sharing of interests, activities and emotions;
   c. lack of socio-emotional reciprocity as shown by an impaired or deviant response to other people’s emotions; or lack of modulation of behavior according to social context; or a weak integration of social, emotional, and communicative behaviors;
   d. lack of spontaneous seeking to share enjoyment, interests, or achievements with other people (e.g. a lack of showing, bringing, or pointing out to other people objects of interest to the individual).

2. Qualitative abnormalities in communication as manifest in at least one of the following areas:
   a. delay in or total lack of, development of spoken language that is not accompanied by an attempt to compensate through the use of gestures or mime as an alternative mode of communication (often preceded by a lack of communicative babbling);
   b. relative failure to initiate or sustain conversational interchange (at whatever level of language skill is present), in which there is reciprocal responsiveness to the communications of the other person;
   c. stereotyped and repetitive use of language or idiosyncratic use of words or phrases;
   d. lack of varied spontaneous make-believe play or (when young) social imitative play

3. Restricted, repetitive, and stereotyped patterns of behavior, interests, and activities are manifested in at least one of the following:
   a. An encompassing preoccupation with one or more stereotyped and restricted patterns of interest that are abnormal in content or focus; or one or more interests that are abnormal in their intensity and circumscribed nature though not in their content or focus;
b. Apparently compulsive adherence to specific, nonfunctional routines or rituals;

c. Stereotyped and repetitive motor mannerisms that involve either hand or finger flapping or twisting or complex whole body movements;

d. Preoccupations with part-objects of non-functional elements of play materials (such as their odor, the feel of their surface, or the noise or vibration they generate).

C. The clinical picture is not attributable to the other varieties of pervasive developmental disorders; specific development disorder of receptive language (F80.2) with secondary socio-emotional problems, reactive attachment disorder (F94.1) or disinhibited attachment disorder (F94.2); mental retardation (F70-F72) with some associated emotional or behavioral disorders; schizophrenia (F20.-) of unusually early onset; and Rett’s Syndrome (F84.12).

Appendix II: Chapter 3 Supplementary Materials

Chapter 3 Supplementary Methods

Surface deformation procedure to place the grey-white matter boundary (i.e. white matter surface)

Within this study we refer to the white matter surface (i.e. the surface that defines the transition from grey to white matter) as the grey-white matter boundary. The surface deformation procedure that places the white matter surface has previously been described by Dale et al. (1999) and is detailed below.

First white matter voxels are labeled through a segmentation procedure. Contiguous white matter voxels are identified through a connected components algorithm resulting in a filled white matter labeled volume. This volume is then tessellated using two triangles to define each voxel composing the surface of the white matter volume. Deformation of this “jagged” white matter tessellation to the grey-white matter boundary is accomplished by a minimization of an energy functional. The first two terms of this energy functional act to smooth the surface and regularize the tessellation by introducing a spring like property to the surface. This spring property is decomposed into two terms given as,

\[
J_n = \frac{1}{2V} \left( \sum_{i=1}^{V} \sum_{j \in N(i)} (\mathbf{n}(i) \cdot (\mathbf{x}_i - \mathbf{x}_j))^2 \right)
\]

\[
J_t = \frac{1}{2V} \left( \sum_{i=1}^{V} \sum_{j \in N(i)} (\mathbf{e}_0(i) \cdot (\mathbf{x}_i - \mathbf{x}_j))^2 + (\mathbf{e}_1(i) \cdot (\mathbf{x}_i - \mathbf{x}_j))^2 \right)
\]

where \(N(i)\) denotes the set of nearest neighbors of the \(i^{th}\) vertex, \(V\) is the total number of vertices in the tessellation, \(\mathbf{n}(i)\) is the unit normal vector to the surface at the \(i^{th}\) vertex, \([\mathbf{e}_0(i), \mathbf{e}_1(i)]\) is an orthonormal basis for the tangent plane at the \(i^{th}\) retex, and \(\mathbf{x}_k\) regers to the \((x,y,z)\) position of the \(k^{th}\) vertex in the tessellation. The term \(J_n\) results in the redistribution of vertices to regions where they are needed, encouraging a uniform spacing of vertices without requiring prohibitive numbers of elements. The term \(J_t\) imposes a smoothness constraint on the surface deformation by penalizing nodes that distance themselves from the direction normal to surface from its neighboring nodes. The third term of the energy functional is based on intensity values. The volume intensity at position \(\mathbf{x}_i\) can be written as \(I(x_i)\) and this term given as,
\[ J_1 = \frac{1}{2V} \left( \sum_{i=1}^{V} (T(i) - I_i)^2 \right) \]

where \( T(i) \) is the mean white matter value of border voxels within a 5mm neighborhood of each vertex, within the segmented white matter volume. The value of \( I(x) \) is computed on a subvoxel basis using trilinear interpolation. The placement of the grey-white matter boundary is achieved by minimizing an energy function that is a weighted sum of the three terms presented above,

\[ J = J_t + \lambda_n I_n + \lambda_l I_l \]

where the coefficients \( \lambda_n \) and \( \lambda_l \) specify the strength of the smoothness and regularization constraints in relation to the intensity term. The gradient of this functional defines the movement of the surface tessellation such as the movement of the \( k \)th vertex is given by the negative of the directional derivative with respect to \( x_k \),

\[ -\frac{\partial J}{\partial x_k} = \lambda_l (T(k) - I_k) \nabla I(x_k) + \sum_{j \in N_i(k)} (\lambda_n (n(k) \cdot x_j) + e_0(k) \cdot x_j + e_1(k) \cdot x_j) \]

where the volume gradient \( \nabla I(x_k) \) is computed using a Gaussian blurred (\( \sigma = 1 \)) version of the MRI volume.

These automated methods for determining the grey-white matter boundary have been previously validated using scans of postmortem brains and have found FreeSurfer based measures of cortical thickness to be on average only 0.077mm different than manual measures performed on dissected tissue samples (Rosas et al. 2002). Within group systematic errors in the placement of the grey-white matter boundary using these methods would result in whole brain differences in cortical thickness that are not observed in our study. These findings thus indicate a high degree of accuracy for FreeSurfer in placing the white matter surface (i.e. the grey-white matter boundary).
Supplementary Figure 1, Regions of decreased cortical thickness (CT) in autism spectrum disorder (ASD): A.) Between group differences in CT (uncorrected). B.) Individuals with ASD showed significantly decreased CT (RFT $p<0.5$) bilaterally in the parahippocampal, fusiform, and lingual gyri (highlighted in blue). See supplementary Table 1 for statistical details of these clusters.
Supplementary Figure 2, Sex differences in grey-white matter signal intensity percent contrast (GWPC): Regardless of diagnosis males showed significantly greater GWPC (RFT \( p<0.5 \)) compared to females across all grey matter sampling depths (a). These increases are highlighted in red and include predominantly fronto-parietal regions of the left hemisphere, and bilateral inferior temporal regions (see supplementary Table 2 for statistical details of these clusters).
Supplementary Figure 3, Regional differences in grey-white matter signal intensity percent contrast (GWPC) and grey matter intensities (GMI) in Autism Spectrum Disorder (ASD) (5mm FWHM smoothing kernel): Between group differences in (A) GWPC and (B) GMI intensities are shown when GMI was sampled at the grey-white matter boundary (i.e. white matter surface, projection fraction 10%) and a projection fraction of 30% into the cortical sheet. Individuals with ASD showed (A) significantly decreased GWPC (RFT $p<0.5$), indicating less definition between grey and white matter, in several regions highlighted in blue. In several of these regions (B) increases in GMI, highlighted in red were also observed. These results using a 5mm FWHM smoothing kernel were largely similar to those using a 10mm FWHM smoothing kernel (Figures 2 and 3, Table 2). For statistical details of these clusters see supplementary Table 3.
**Supplementary Table 1**

<table>
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<th>Cluster</th>
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<th>$t_{max}$</th>
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<td>3728</td>
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<td>-42</td>
<td>-5</td>
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**Supplementary Table 1, Clusters of Decreased Cortical Thickness in Autism Spectrum**

**Disorder (ASD):** Broadmann area (BA), left (L), right (R), *Vertices* indicates the number of vertices within the cluster, $t_{max}$ represents the maximum $t$-statistic within the cluster located at the $x$ $y$ $z$ Talairach coordinates listed, $p_{cluster}$ is the cluster corrected $p$ value.
Supplementary Table 2, Clusters of Increased Grey-White Matter Signal Intensity Percent Contrast (GWPC) in Males: Broadmann area (BA), left (L), right (R), Vertices indicates the number of vertices within the cluster, $t_{max}$ represents the maximum $t$-statistic within the cluster located at the x y z Talairach coordinates listed, $p_{cluster}$ is the cluster corrected $p$ value.
Supplementary Table 3

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<tr>
<td>Grey-White Matter Percent Contrast</td>
<td></td>
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</tr>
<tr>
<td>1. insula, lateral orbital frontal cortex, superior temporal gyrus</td>
<td>R</td>
<td>38 4370 37 0 -12 4.01 7.28 x 1</td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>2. banks superior temporal sulcus, inferior parietal cortex, middle and superior temporal gyrus</td>
<td>L</td>
<td>21 4021 -50 -26 -2 3.64 7.89 x 1</td>
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<tr>
<td>3. banks superior temporal sulcus, inferior parietal cortex, inferior, middle, and superior temporal gyrus</td>
<td>R</td>
<td>41 3735 46 -36 7 2.95 1.05 x 1</td>
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<tr>
<td>4. fusiform gyrus, lingual gyrus, parahippocampal gyrus</td>
<td>R</td>
<td>30 3522 18 -38 -6 3.90 7.40 x 1</td>
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</tr>
<tr>
<td>5. isthmus-cingulate cortex, precuneus cortex</td>
<td>L</td>
<td>29 2694 -14 -49 7 3.38 3.39 x 1</td>
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<td></td>
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</tr>
<tr>
<td>6. fusiform gyrus, inferior temporal gyrus, lingual gyrus</td>
<td>L</td>
<td>19 2410 -29 -56 -3 3.56 2.22 x 1</td>
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<td></td>
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<tr>
<td>7. medial orbital frontal cortex, rostral anterior cingulate cortex, superior frontal gyrus</td>
<td>L</td>
<td>32 2165 -11 42 2 3.40 2.96 x 1</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. postcentral gyrus, superior parietal cortex</td>
<td>L</td>
<td>2 1868 -46 -23 43 3.56 1.66 x 1</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>9. insula, lateral orbital frontal cortex</td>
<td>L</td>
<td>13 1654 -30 18 -2 3.79 4.09 x 1</td>
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</tr>
<tr>
<td>10. paracentral lobule, superior parietal cortex, superior temporal gyrus</td>
<td>L</td>
<td>5 1458 -16 -35 50 3.11 4.85 x 1</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. superior temporal gyrus</td>
<td>L</td>
<td>21 1228 -45 -9 -13 2.78 2.73 x 1</td>
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<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>12. inferior temporal gyrus</td>
<td>L</td>
<td>20 1085 -48 -10 -23 3.47 3.34 x 1</td>
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</table>
Supplementary Table 3, Significant Reductions in Grey-White Matter Signal Intensity Percent Contrast (GWPC) and Increases in Grey Matter Intensity (GMI) in ASD (FWHM 5mm): Broadmann area (BA), left (L), right (R), Vertices indicates the number of vertices within the cluster, \( t_{\text{max}} \) represents the maximum \( t \)-statistic within the cluster located at the \( x \ y \ z \) Talairach coordinates listed, \( p_{\text{cluster}} \) is the cluster corrected \( p \) values.
Appendix III: Chapter 5 Supplementary Materials

Supplementary Online Content


eTable 1. Frequency Ratios of ASD Cases to TD Controls, and Males to Females, Acquired Within and Across the Different Acquisition Sites

eTable 2. Population Prevalence of ASD for Biological Males and Females With Male or Female Neuroanatomical Brain Phenotype

eTable 3. Differences in Neuroanatomical Patterns Associated With High Probability of ASD Between Men and Women

eTable 4. Results of the Vertex-wise Analysis of CT Utilizing a General Linear Model (GLM)

eTable 5. Differences in Demographic Measures Between Diagnostic Groups and Biological Sexes

eMethods. Supplementary Methods

eFigure 1. Sex-Related Pattern of Neuroanatomical Variability in Cortical Thickness

eFigure 2. Significant Differences in CT Between Male and Female TD Controls

eFigure 3. Sample Probability of ASD for Females (A) and Males (B) as a Function of Normative Sex-Related Phenotypic Variability in CT

eFigure 4. Clusters With Significant Differences in the Neuroanatomical Pattern of CT Associated With High Probability of ASD Between Males and Females (i.e. Product of w and CT)

eFigure 5. Clusters With Significant Group-by-Sex Interactions Following a Non-parametric Clustering Approach With n=10,000 Permutations of Group (i.e. Sex) Labels (Permutation-Based Cluster-Corrected, p<0.05)

eFigure 6. Group-by-Sex Interaction Graph Based on Mean CT in the Left-Hemisphere Cluster That Extended Into the Hippocampal/Entorhinal Cortex, the Fusiform and Lingual Gyrus, and the Inferior/Middle Temporal Lobe

eFigure 7. Differences in Full-Scale, Verbal and Non-verbal (i.e. Performance) IQ Between Individual Subgroups of ASD vs. TD Individuals, and Males vs. Females, Examined Using t Tests for Independent Samples

eFigure 8. Spatially Distributed Pattern of Negative (Blue) and Positive (Red) Correlations Between Verbal IQ Measures and CT Assessed Across Male and Female Neurotypical Controls

eAppendix. Supplementary Appendix

eReferences. Supplementary References

This supplementary material has been provided by the authors to give readers additional information about their work.

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**eTable 1.** Frequency Ratios of ASD Cases to TD Controls, and Males to Females, Acquired Within and Across the Different Acquisition Sites

<table>
<thead>
<tr>
<th></th>
<th>Group</th>
<th>ASD</th>
<th>Control</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cambridge</td>
<td>female</td>
<td>28</td>
<td>28</td>
<td>56</td>
</tr>
<tr>
<td></td>
<td>male</td>
<td>25</td>
<td>26</td>
<td>51</td>
</tr>
<tr>
<td>London</td>
<td>female</td>
<td>21</td>
<td>19</td>
<td>40</td>
</tr>
<tr>
<td></td>
<td>male</td>
<td>24</td>
<td>25</td>
<td>49</td>
</tr>
</tbody>
</table>
**eTable 2.** Population Prevalence of ASD for Biological Males and Females With Male or Female Neuroanatomical Brain Phenotype

<table>
<thead>
<tr>
<th></th>
<th>S = 1</th>
<th>S = 0</th>
</tr>
</thead>
<tbody>
<tr>
<td>D = 1</td>
<td>0.025</td>
<td>0.017</td>
</tr>
<tr>
<td>D = 0</td>
<td>0.975</td>
<td>0.983</td>
</tr>
</tbody>
</table>

_Note._ D: diagnosis of ASD (0=no, 1=yes), S: biological sex/gender (0=female, 1=male), N: neuroanatomical brain phenotype (0=female, 1=male). Notably, the probability of receiving a diagnosis of ASD appears to increase for male participants with N=0. This is due to the fact that the probability of being classified as N=0 for biological males is relatively low, thereby inflating the conditional probability of having ASD.

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### eTable 3 Differences in Neuroanatomical Patterns Associated With High Probability of ASD Between Men and Women

<table>
<thead>
<tr>
<th>ID</th>
<th>Region Labels</th>
<th>Side</th>
<th>BA</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>T_max</th>
<th>P (corrected)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>banks superior temporal sulcus, caudal middle frontal gyrus, inferior</td>
<td>R</td>
<td>11/22/44/45</td>
<td>34</td>
<td>-25</td>
<td>14</td>
<td>4.96</td>
<td>6.09 x 10^{-5}</td>
</tr>
<tr>
<td></td>
<td>parietal cortex, insula, lateral, orbital frontal cortex, middle temporal</td>
<td></td>
<td>11/45/46</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>gyrus, pars opercularis, pars triangularis, postcentral gyrus, precentral</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>gyrus, rostral middle frontal gyrus, superior temporal gyrus</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>cuneus cortex, fusiform gyrus, isthmus-cingulate cortex, lingual gyrus,</td>
<td>L</td>
<td>18/19/20/30/</td>
<td>-32</td>
<td>-48</td>
<td>-16</td>
<td>3.78</td>
<td>6.29 x 10^{-5}</td>
</tr>
<tr>
<td></td>
<td>precuneus cortex.</td>
<td></td>
<td>31/37</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>insula, lateral orbital frontal cortex, pars opercularis, pars triangularis,</td>
<td>L</td>
<td>11/45/47</td>
<td>-38</td>
<td>-18</td>
<td>16</td>
<td>4.57</td>
<td>1.37 x 10^{-5}</td>
</tr>
<tr>
<td></td>
<td>postcentral gyrus, precentral gyrus, supramarginal gyrus</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>inferior parietal cortex, supramarginal gyrus</td>
<td>R</td>
<td>19/39/40</td>
<td>41</td>
<td>-66</td>
<td>43</td>
<td>4.54</td>
<td>1.35 x 10^{-5}</td>
</tr>
<tr>
<td>5</td>
<td>fusiform and lingual gyrus, parahippocampal gyrus</td>
<td>R</td>
<td>19/20/37</td>
<td>24</td>
<td>-44</td>
<td>-7</td>
<td>3.47</td>
<td>1.09 x 10^{-5}</td>
</tr>
<tr>
<td>6</td>
<td>caudal anterior-cingulate cortex, medial orbital frontal cortex, rostral</td>
<td>L</td>
<td>25/32/24</td>
<td>-12</td>
<td>42</td>
<td>8</td>
<td>4.45</td>
<td>1.27 x 10^{-5}</td>
</tr>
<tr>
<td></td>
<td>anterior cingulate cortex, superior frontal gyrus</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>caudal anterior-cingulate cortex, paracentral lobule, posterior-cingulate</td>
<td>L</td>
<td>24</td>
<td>-13</td>
<td>-15</td>
<td>38</td>
<td>3.97</td>
<td>1.34 x 10^{-5}</td>
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<tr>
<td></td>
<td>cortex, precuneus cortex, superior frontal gyrus</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>S</td>
<td>postcentral gyrus, superior parietal cortex</td>
<td>L.</td>
<td>7</td>
<td>-27</td>
<td>-55</td>
<td>60</td>
<td>3.25</td>
<td>4.05 x 10^-2</td>
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*Note:* BA, Brodmann Area; tmax, maximum absolute t value within cluster; x,y,z, Talairach coordinates at tmax, pcluster, cluster probability. R, right; L, left.
### eTable 4 Results of the Vertex-wise Analysis of CT Utilizing a General Linear Model (GLM)

<table>
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<th>$y$</th>
<th>$z$</th>
<th>$t_{max}$</th>
<th>$p_{correct}$</th>
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<tbody>
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<td></td>
<td><strong>Group-by-Sex Interaction</strong></td>
<td></td>
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</tr>
<tr>
<td>(1)</td>
<td>fusiform gyrus, inferior/middle temporal gyrus, lingual gyrus, parahippocampal gyrus</td>
<td>L</td>
<td>19/20/34/35/37/38</td>
<td>-29</td>
<td>-71</td>
<td>-5</td>
<td>3.29</td>
<td>3.5 x 10^-6</td>
</tr>
<tr>
<td>(2)</td>
<td>fusiform gyrus, inferior temporal gyrus, lateral occipital cortex, lingual gyrus, parahippocampal gyrus</td>
<td>R</td>
<td>18/19/34/35/37</td>
<td>26</td>
<td>-27</td>
<td>-18</td>
<td>3.42</td>
<td>5.49 x 10^-4</td>
</tr>
<tr>
<td>(3)</td>
<td>superior temporal sulcus, middle temporal gyrus</td>
<td>R</td>
<td>21/22</td>
<td>60</td>
<td>-33</td>
<td>-6</td>
<td>3.51</td>
<td>2.13 x 10^-2</td>
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</tbody>
</table>

**Contrast: ASD males < TD males**

<table>
<thead>
<tr>
<th>ID</th>
<th>Regional Labels</th>
<th>Side</th>
<th>BA</th>
<th>$x$</th>
<th>$y$</th>
<th>$z$</th>
<th>$t_{max}$</th>
<th>$p_{correct}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1)</td>
<td>fusiform gyrus, inferior/middle temporal gyrus</td>
<td>L</td>
<td>18/19/20/37</td>
<td>-52</td>
<td>27</td>
<td>-19</td>
<td>2.61</td>
<td>1.04 x 10^-2</td>
</tr>
<tr>
<td>(2)</td>
<td>fusiform gyrus, lateral occipital cortex, lingual gyrus, parahippocampal gyrus</td>
<td>R</td>
<td>18/19/37</td>
<td>29</td>
<td>-60</td>
<td>-8</td>
<td>2.71</td>
<td>2.90 x 10^-2</td>
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</tbody>
</table>

**Contrast: ASD females < TD females**

<table>
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<th>Regional Labels</th>
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<th>$y$</th>
<th>$z$</th>
<th>$t_{max}$</th>
<th>$p_{correct}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1)</td>
<td>entorhinal cortex, fusiform gyrus, inferior/middle temporal gyrus, lingual gyrus, parahippocampal gyrus</td>
<td>L</td>
<td>18/19/20/34/35/37/38</td>
<td>-30</td>
<td>-69</td>
<td>-6</td>
<td>3.90</td>
<td>2.23 x 10^-6</td>
</tr>
<tr>
<td>(2)</td>
<td>entorhinal cortex, fusiform gyrus, inferior/middle temporal gyrus, lateral occipital cortex, lingual gyrus, parahippocampal gyrus</td>
<td>R</td>
<td>18/19/20/34/35/36/37</td>
<td>25</td>
<td>-26</td>
<td>-18</td>
<td>4.13</td>
<td>1.08 x 10^-6</td>
</tr>
<tr>
<td>(3)</td>
<td>superior temporal sulcus gyrus, inferior parietal cortex, middle/superior temporal gyrus</td>
<td>R</td>
<td>21/22/39</td>
<td>62</td>
<td>-33</td>
<td>-6</td>
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<td>1.08 x 10^-7</td>
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<td>Area</td>
<td>Side</td>
<td>MA</td>
<td>x</td>
<td>y</td>
<td>z</td>
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<td>--------</td>
</tr>
<tr>
<td>4</td>
<td>insula, pre-/post-central</td>
<td>L</td>
<td>40/41</td>
<td>-44</td>
<td>-18</td>
<td>16</td>
<td>-3.96</td>
<td>1.23 \times 10^{-2}</td>
</tr>
<tr>
<td></td>
<td>gyrus, supramarginal gyrus</td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>5</td>
<td>insula, pre-/post-central</td>
<td>R</td>
<td>40/41</td>
<td>46</td>
<td>-14</td>
<td>20</td>
<td>-3.67</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>gyrus, supramarginal gyrus</td>
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<td></td>
</tr>
</tbody>
</table>

Note. ID, cluster number; BA, approximate Brodmann Area; \( t_{max} \), maximal \( t \) test statistic within cluster; \( x,y,z \), Talairach coordinates at \( t_{max} \); \( p_{cluster} \), cluster probability; R, right; L, left.
**eTable 5. Differences in Demographic Measures Between Diagnostic Groups and Biological Sexes**

<table>
<thead>
<tr>
<th></th>
<th>Groups</th>
<th>ASD</th>
<th>Control</th>
<th>F(df=1)</th>
<th>p</th>
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<tr>
<td></td>
<td>(n=98)</td>
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<td>(n=98)</td>
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<tr>
<td>Age [years]</td>
<td>26.88 ± 7.18</td>
<td>27.39 ± 6.44</td>
<td>0.099</td>
<td>0.585</td>
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</tr>
<tr>
<td>Full-scale IQ (WASI)</td>
<td>113.74 ± 12.31</td>
<td>116.44 ± 9.48</td>
<td>3.142</td>
<td>0.078</td>
<td></td>
</tr>
<tr>
<td>Verbal IQ</td>
<td>113.52 ± 12.89</td>
<td>113.68 ± 10.75</td>
<td>0.027</td>
<td>0.871</td>
<td></td>
</tr>
<tr>
<td>Performance IQ</td>
<td>111.28 ± 14.02</td>
<td>115.61 ± 9.75</td>
<td>6.179</td>
<td>0.014*</td>
<td></td>
</tr>
<tr>
<td>Mean Cortical Thickness [mm]</td>
<td>2.41 ± 0.01</td>
<td>2.34 ± 0.09</td>
<td>0.010</td>
<td>0.921</td>
<td></td>
</tr>
<tr>
<td>Total Grey Matter Volume [litre]</td>
<td>0.72 ± 0.08</td>
<td>0.72 ± 0.07</td>
<td>0.003</td>
<td>0.955</td>
<td></td>
</tr>
<tr>
<td>Total Intracranial Volume [litre]</td>
<td>1.45 ± 0.24</td>
<td>1.45 ± 0.21</td>
<td>0.000</td>
<td>0.994</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Biological Sex</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Males (n=100)</td>
<td>Females (n=96)</td>
</tr>
<tr>
<td>Age [years]</td>
<td>26.70 ± 6.41</td>
<td>27.60 ± 7.21</td>
</tr>
<tr>
<td>Full-scale IQ (WASI)</td>
<td>113.66 ± 11.57</td>
<td>116.59 ± 10.32</td>
</tr>
<tr>
<td>Verbal IQ</td>
<td>110.18 ± 11.98</td>
<td>117.21 ± 10.59</td>
</tr>
<tr>
<td>Performance IQ</td>
<td>114.42 ± 12.49</td>
<td>112.45 ± 11.92</td>
</tr>
<tr>
<td>Mean Cortical Thickness [mm]</td>
<td>2.34 ± 0.085</td>
<td>2.33 ± 0.11</td>
</tr>
<tr>
<td>Total Grey Matter Volume [litre]</td>
<td>0.763 ± 0.06</td>
<td>0.677 ± 0.10</td>
</tr>
<tr>
<td>Total Intracranial Volume [litre]</td>
<td>1.58 ± 0.19</td>
<td>1.32 ± 0.17</td>
</tr>
</tbody>
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Note. Data expressed as mean ± standard deviation, F, p resulting from a general linear model (GLM) including a main effect of diagnostic group and biological sex. * statistically significant based on p < 0.05 (two-sided)
eMethods. Supplementary Methods

Gaussian Process Classification of biological sex based on normative variability in CT

Gaussian process models were used for the probabilistic prediction of biological sex based on patterns of neuroanatomical variability in CT. Gaussian process classification (GPC) has previously been applied to MRI data, and a detailed description can be found in. Briefly, we begin with a set of training data \( X_{train} \) consisting of input vectors \( x \) (i.e. training samples with \( d \) features each), and \( y \) is a column vector of target variables or class labels where \( y \in \{-1, +1\} \). Training samples are indexed by \( i = 1, \ldots, m \). The training data are used to estimate a probability distribution that allows us to predict a target \( y^* \) for a new data sample \( x^* \) as accurately as possible. For binary GPC, predictions take the form of predictive or ‘class’ probabilities, where the probability for a class results from \( P(y^* = +1|x^*,X_{train}) \). These predictions are computed by first estimating the posterior distribution of a set of vertex weights using the rules of probability, then passing the output through a sigmoid function to constrain the output to the unit interval, ensuring a valid probabilistic interpretation. The mode of the distribution of weights, \( w \), can be understood as providing a spatial representation of the decision boundary. If GPC is applied to neuroimaging data, \( w \) has the same number of elements as the feature vector \( x \), and can subsequently be used to create a map of the most important brain regions underlying the prediction (analogous to Support Vector Machine (SVM) discrimination maps).

Here, we initially employed GPC to establish (i.e. train) a predictive model of biological sex based on normative phenotypic variability in CT. Thus, the input matrix \( X \) was an \( m \times d \) matrix of CT measures where \( m \) was defined as the number of typically developing (TD) controls (i.e. 51 males and 47 females), and \( d \) was defined as the number of vertices across the cortical surface. As target variable \( y \), we used the class labels of -1 and +1 for neurotypical females and males respectively. This allowed us to...
make probabilistic predictions for individuals based on the binary categories dictated by biological sex, utilizing their neuroanatomical brain phenotype. Class probabilities ranged from 0 to 1, wherein those being most confidently classified as phenotypically female (i.e. class probability=0) or male (i.e. class probability=1) representing the most prototypical (i.e. characteristic) male or female neurophenotype respectively. The GP classifier was trained by maximizing the logarithm of the marginal likelihood, which measures the total probability of the data given the Gaussian process prior and the model’s hyperparameters. The performance of the normative model was subsequently validated using cross validation, which provides robust parameter estimates of generalisation ability. Here, the classifier is trained repeatedly using all but one participant from each class label. This procedure was repeated \(n\) times, where \(n\) is the number of participants per class label. The accuracy of the classifier was measured by the proportion of observations that were correctly classified, e.g. the number of biological males (or females) correctly classified as male (or female) based on their brain phenotype. Permutation testing was used to assess the statistical significance of the model performance relative to chance level. To achieve this, we permuted the class (i.e. sex) labels 1000 times without replacement. Each time, the model was retrained and the accuracy was computed. All Gaussian process models were estimated using the GPML toolbox (www.gaussianprocess.org/gpml/code).

The normative predictive model of biological sex was subsequently used to predict biological sex for the independent sample of ASD individuals. To visualize the spatially-distributed patterns of neuroanatomical variability in CT carrying high and low risk of ASD in males and females, we employed a recently proposed approach for mapping the contribution of each vertex to the classifier decision. This approach has two advantages over the more conventional approach of mapping the weight vector: (1) it indicates the total contribution of each vertex to the prediction for each individual, i.e. not just the contribution from the training set; and (2) it can be used to map the neural constituents
for the predictions based on a second sample, independent from the training set (i.e. the clinical and normative samples respectively). To achieve this, we calculated the element-wise product of \( w \) and each individual's CT measures (i.e. \( x^\text{CT} \)) for (1) all males (or all females) classified as phenotypically male (i.e. class probability > 0.5), (2) all males (or all females) classified as phenotypically female (i.e. class probability < 0.5), and (3) across biological males and females with a class probability < 0.5 (or a class probability > 0.5). Within each of these groups and at each cerebral vertex, the resulting maps were then summarized by the mean value across individuals.

**Estimation of ASD probability based on normative sex-related phenotypic variability of the brain**

To determine whether normative sex-related variability in CT measures modulates the probability of ASD, we initially examined the (absolute) probability of ASD as a function of predictive probabilities for the TD male or female brain phenotype within our sample. The sample probability of ASD for females (or males) was determined as the ratio of females (or males) with ASD divided by the total number of females (or males) falling within eight bins (i.e. from 0 to 1 in steps of 0.125) along the axis of predictive class probabilities for the male neuroanatomical phenotype (\( M \)). Let \( D \) and \( S \) denote binary variables representing a diagnosis of ASD and biological sex respectively. This equals the posterior probability of having ASD (i.e. \( D=1 \)) under the condition of being biologically female or male (i.e. \( S=0 \) or \( S=1 \), respectively), and given a neuroanatomical phenotype \( M \) quantified by class probabilities, so that \( P(D=1|M, S=s) \), for every \( s \in \{0,1\} \).

Furthermore, this data allowed us to estimate the population risk of having a diagnosis of ASD, given that an individual exhibits a male or female neuroanatomical phenotype. In our notation, this may be expressed as \( P(D=1|M=m, S=s) \), for every \( m \in \{0,1\} \) and \( s \in \{0,1\} \) where 0 denotes female and 1 denotes male. Using Bayes theorem, such a conditional probability can be expressed as
\[ P(D = 1|M = m, S = s) = \frac{P(M = m|D = 1, S = s)P(D = 1|S = s)}{P(M = m|S = s)}, \quad (1) \]

in which the normalizing constant is obtained by integrating out \( D \), so that \( P(M|D, S) = P(M|D = 1, S)P(D = 1|S) + P(M|D = 0, S)P(D = 0|S) \) for every realization of \( M \) and \( S \). The probabilities in the numerator of the right-hand side of equation (1) are readily obtained from our sample, and from previous epidemiological studies (e.g. ⁴). Indeed, the term \( P(M = m|D = 1, S = s) \) corresponds to the number of biological males or females with a diagnosis of ASD, who have been identified as having a male or female neuroanatomical phenotype by our GPC model.

Moreover, the prevalence rates of ASD for males and females in the general population (i.e. \( P(D = 1, S = s) \)) are known from epidemiological priors (i.e. 1:42 for males, 1:189 for females ⁴). Based on these previous priors, and the sample estimates resulting from our study, we were therefore able to estimate the population risk of ASD given a male/female neuroanatomical phenotype for biological males and females, i.e. \( P(D|M, S) \). Note that the estimation of \( P(D = 1|M, S) \) presented here is analogous to the correction usually applied to a sensitivity test in order to compute the probability of having a disease, given a positive test result. In our context, the probability of a particular neuroanatomical phenotype can thus be regarded as a 'test' for disease status.

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**eFigure 1.** Sex-Related Pattern of Neuroanatomical Variability in Cortical Thickness

(A) Spatial distribution of the weight vector $w$ resulting from the Gaussian Process Classification of biological sex in TD controls. (B) Un-thresholded statistical difference map representing t-test statistics resulting from the GLM-comparison between TD males and females using CT measures. Positive $t$-values (yellow to red colormap) indicate increased CT in TD females relative to males, while negative $t$-values (cyan to blue colormap) indicate decreased CT in females. Clusters with significant differences in CT following statistical thresholding are shown in eFigure 2 in the Supplement. (C) Patterns of normative neuroanatomical variability in CT associated with a phenotypic shift of the brain from a characteristic female to male presentation. These patterns were derived using a predictive mapping approach, which quantifies the extent to which the individual's neuroanatomy interacts with the weight vector $w$. At each vertex, the color-scale therefore represents the product of $w$ and CT, averaged across all individuals falling within four bins along the normative axis of predictive probabilities. (D) Patterns of sex-related neuroanatomical CT variability associated with high and low risk for ASD across sex categories. The risk of ASD was determined as the number of males and females with ASD relative to the total number of individuals falling within four intervals of predictive class probabilities. At each vertex, the color-scale indicates the product of $w$ and CT, averaged across all individuals within the four risk groups. Brackets indicate the 95% confidence interval associated with the risk for ASD.
A) Clusters with significant between-group differences using random-field theory (RFT)-based clustering with a cluster threshold of 0.5 (two-tailed). The orange to yellow colormap indicates brain regions where CT is significantly increased in females relative to males. (B) Clusters with significant between-group differences using a non-parametric clustering approach with $n=10,000$ permutations at the same cluster-threshold and colormap.
**eFigure 3.** Sample Probability of ASD for Females (A) and Males (B) as a Function of Normative Sex-Related Phenotypic Variability in CT

The sample probability of ASD was determined as the ratio of females (or males) with ASD relative to the total number of females (or males) falling within eight bins (i.e., from 0 to 1 in steps of 0.125) along the normative axis of predictive probabilities. The colored area around the regression line indicates the 95% confidence interval (CI) resulting from a binomial test, based on the hypothesis that the probability of having ASD is larger (or smaller) than the probability of not having ASD.

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eFigure 4. Clusters With Significant Differences in the Neuroanatomical Pattern of CT Associated With High Probability of ASD Between Males and Females (i.e., Product of w and CT, See Figure 2 Right Panel)

The red colormap indicates regions where females had a significant increased w x CT as compared to males (RFT-based cluster-corrected, p<0.05). There were no significant between-sex differences in the low ASD probability maps.

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**Figure 5.** Clusters With Significant Group-by-Sex Interactions Following a Non-parametric Clustering Approach With \( n = 10,000 \) Permutations of Group (i.e. Sex) Labels (Permutation-Based Cluster-Corrected, \( p < 0.05 \)) (A). (B) Clusters where CT was significantly reduced in females with ASD relative to female controls (permutation-based cluster-corrected, \( p < 0.05 \)). No significant clusters remained when comparing ASD males with male TD controls using a permutation-based cluster correction.
**eFigure 6.** Group-by-Sex Interaction Graph Based on Mean CT in the Left-Hemisphere Cluster That Extended Into the Hippocampal/Entorhinal Cortex, the Fusiform and Lingual Gyrus, and the Inferior/Middle Temporal Lobe (see cluster ID 1 in eTable 4)
eFigure 7. Differences in Full-Scale, Verbal and Non-verbal (i.e. Performance) IQ Between Individual Subgroups of ASD vs. TD Individuals, and Males vs. Females, Examined Using t Tests for Independent Samples

Note. ** indicates $p<0.01$, * indicates $p<0.05$. 

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**eFigure 8.** Spatially Distributed Pattern of Negative (Blue) and Positive (Red) Correlations Between Verbal IQ Measures and CT Assessed Across Male and Female Neurotypical Controls

The colormap indicates the t test statistics for vertex-wise correlation coefficients.
eAppendix. Supplementary Appendix

Relationship between sex-related neuroanatomical differences and inter-individual variability in clinical and/or neurocognitive measures

To determine how the observed sex-related differences in brain structure may be related to variability in clinical or neurocognitive measures, we also examined if differences in verbal and non-verbal (i.e. performance) IQ between diagnostic groups and biological sexes could explain the phenotypic shift of the brain in females. For example, neurotypical females tend to have a significantly higher verbal IQ than neurotypical males, which has previously been suggested to be a ‘protective’ factor for ASD (e.g. 5). It is therefore possible that females with ASD display a neurocognitive profile that is more similar to the neurotypical male rather than female behavioral phenotype, which in turn could account for a more male-typical pattern of brain anatomy.

However, when examining similarities and differences in IQ measures between females with ASD, and male and female controls, we found that ASD females differed significantly from neurotypical males in both verbal (t(97)=-2.993, p<0.01) and performance IQ (t(97)=2.406, p<0.05). Moreover, ASD females were not statistically different from female controls (p>0.05, two-tailed) (see eFigure 7 in the Supplement). Based on IQ profiles, the ASD females assessed in our sample therefore seem functionally closer to female rather than male TD controls. Thus, based on IQ measures exclusively, our finding of increased predictive probabilities for the male neuroanatomical brain phenotype in ASD females seems to reflect a sex- rather than function-related difference in brain anatomy. Moreover, there was no significantly correlation between predictive probabilities for a male neuroanatomical brain phenotype and variability in verbal or performance IQ in females (p>0.05, two-tailed), as would be expected based on the absence of a significant difference between females with ASD and female controls. Given that females tend to have a significantly increased verbal IQ compared to males, we also compared the

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normative patterns of neuroanatomical variability indicative of biological sex (i.e. the weight vector \( w \), eFigure 1a) with the spatially distributed pattern of correlations between CT measures and verbal IQ (see eFigure 8 in the Supplement). Overall, however, the distributed patterns indicative of biological sex, and the patterns associated with high and low ASD probability, are quite different from the spatial distribution of high/low correlations between CT and verbal IQ. Based on these findings, it seems unlikely that the results of our multivariate prediction of biological sex are driven by sex-dependent differences in verbal IQ, rather than by biological sex itself.

Last, we also explored the relationship between autistic symptoms and CT variability in the two main clusters with significant sex-by-group interactions (see Figure 3a) within the ASD group. Here, we utilized a GLM predicting CT variability using separate effects of biological sex and measures of symptom severity to (1) examine whether CT in these clusters is significantly correlated with measures of symptom severity within the ASD group, and to (2) disentangle the effect of biological sex from the effect of inter-individual variability in symptom severity. We examined these relationships in two separate statistical models, one including the three ADI-R subdomains, and one including the ADOS total and repetitive domain. Overall, we found that CT measures in the right-hemisphere cluster correlated with ADOS total scores exclusively (\( r(97) = 0.283, p < 0.05 \), two-tailed). No significant correlations were observed in the left-hemisphere cluster. However, when predicting CT by biological sex and ADOS total scores, both effects remained statistically significant in the right hemisphere (\( p < 0.05 \), two-tailed). In the left hemisphere, only the effect of biological sex exceeded the level of statistical significance (\( p < 0.05 \), two-tailed). Thus, while measures of symptom severity may explain some CT variability in addition biological sex, it seems that both effects are statistically separable. Although a more thorough investigation of this issue is required in the future, these preliminary results
indicate that our findings reflect neuroanatomical differences related to biological sex, and having a
diagnosis of ASD, rather than functional differences in specific clinical or neurocognitive features.
eReferences. Supplementary References


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Appendix IV: Chapter 6 Supplementary Materials

Supplementary Figure 6. Morphological Differences Between Typically Developing (TD) Females and Males as Identified Using a General Linear Model: Sex differences between TD males and females for (A) cortical thickness, (B) surface area, (C) the local gyrification index, and (D) sulcal depth. Uncorrected t-maps are shown in the left panel, random field theory cluster corrected ($p<0.5$) maps are shown in the right panel.
Supplementary Figure 3. Probability for ASD as a Function of Normative Sex-Related Phenotypic Variability in Brain Morphology: Probability estimates for ASD across eight discrete bins along the axis of predictive class probabilities for biological sex are plotted for (A) cortical thickness, (B) surface area, (C) the local gyrification index, and (D) sulcal depth for biological females and males (left and right panel respectively). For each model (excluding local gyrification index measures in biological males) probability for ASD was seen to increase along with increasingly male-typical class predictions. This relationship was particularly apparent in females due to the statistically significant shift in females with ASD towards male-typical neuroanatomical phenotypes.
Supplementary Figure 2, Group x Sex Interaction Effects Identified Using a General Linear Model: Uncorrected $t$ maps of group x sex interaction effects are shown in the left panel for (A) cortical thickness, (B) surface area, (C) the local gyrification index, and (D) sulcal depth. Significant clusters after random field theory (RFT) correction (p<0.05) are shown in the middle panel. Plots of the mean values from cortical thickness cluster C1, surface area cluster C2, and local gyrification index cluster C3 are shown in the right panel.
## Supplementary Table 1

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**Supplementary Table 1**, Clusters of Significant Differences in Cortical Morphometry Utilizing a General Linear Model (TD Females – TD Males): left (L), right (R), Vertices indicates the number of vertices within the cluster, t<sub>max</sub> represents the maximum t-statistic within the cluster located at the x y z Talairach coordinates listed, p<sub>cluster</sub> is the cluster corrected p value.
### Supplementary Table 2

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**Supplementary Table 2, Population Prevalence of ASD for Biological Males and Females with Male- or Female-typical Neuroanatomical Brain Phenotypes:** Population estimates are given based on the predictive class probabilities for each morphological feature (i.e. cortical thickness [CT], surface area [SA], the local gyrification index [lGI], and sulcal depth [SULC]) as well as the average of these class probabilities across features (mean). D: diagnosis of ASD (0=no, 1=yes), S: biological sex (0=female, 1=male), N: neuroanatomical brain phenotype (0=female-typical, 1=male-typical).
## Supplementary Table 3

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<td>fusiform gyrus, inferior temporal gyrus, lateral occipital cortex, lingual gyrus, parahippocampal gyrus</td>
<td>R</td>
<td>6835</td>
<td>26</td>
<td>-27</td>
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<tr>
<td></td>
<td>3</td>
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<td>R</td>
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<td>-33</td>
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<tr>
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<td>R</td>
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<td>R</td>
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<tr>
<td></td>
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<td>L</td>
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**Supplementary Table 3, Clusters of Significant Group-by-Biological Sex Interaction Effects:** left (L), right (R), *Vertices* indicates the number of vertices within the cluster, $t_{\text{max}}$ represents the maximum $t$-statistic within the cluster located at the x y z Talairach coordinates listed, $p_{\text{cluster}}$ is the cluster corrected $p$ value.
Appendix V: Chapter 6 Manuscript Proof

Normative Sex-Related Phenotypic Diversity Across Multiple Measures of Brain Anatomy is Associated with Probability for Autism Spectrum Disorder

Journal: The American Journal of Psychiatry

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Normative Sex-Related Phenotypic Diversity Across Multiple Measures of Brain Anatomy is Associated with Probability for Autism Spectrum Disorder

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KEY WORDS: ASD, FreeSurfer, Sex, Gender, MRI
Abstract

Objective: Males are 2 to 5 times more likely to be diagnosed with Autism spectrum disorder (ASD) than females. The reasons for this bias are not fully understood. However, normative diversity in brain structure associated with biological sex is one possible mediator for ASD risk. The authors therefore examined the probability of ASD as a function of sex-related variability in multiple measures of brain anatomy.

Method: 98 adults with ASD (49 males, 49 females) and 98 typically developing controls (51 males, 47 females) underwent structural MRI scanning. Measures of cortical morphometry (i.e. cortical thickness, surface area, the local gyriﬁcation index, and sulcal depth) were used to establish normative models of biological sex. These models were subsequently applied to males and females with ASD. Sample probabilities for ASD were calculated across features according to sex-based neuroanatomical phenotypes and extrapolated to estimate ASD risk within the general population.

Results: Across morphometric features, females with ASD displayed a statistically significant neuroanatomical shift towards more male-typical presentations \( (p<0.001) \). Moreover, within both biological males and females sample probabilities for ASD increased with predicted probabilities for male-typical brain phenotypes. For biological females, this translated to an estimated variability in population prevalence ranging from 0.01 to 2.6% depending on having either a female- or male-typical neurophenotype.

Conclusions: Male-typical neurophenotypes are associated with higher probability for having an ASD diagnosis than female-typical neurophenotypes across a range of neuroanatomical features. In addition to genetic and environmental factors, normative sex-related phenotypic diversity in brain structure should be considered when investigating risk and resilience for ASD.
Introduction

Autism spectrum disorder (ASD) is a complex neurodevelopmental condition estimated to be 2-5 times more common in males than females (1, 2). While the preponderance of males with ASD has partially been attributed to sex differences in clinical phenotypes (3-7), etiological models propose that factors inherent to normative sex differences may modulate ASD liability (for review see (8)). However, it remains unknown what "male risk" and/or "female protective" factors may contribute to the male preponderance of ASD, and how these factors may affect the biological underpinnings of the condition.

To identify potential sex-related risk (and protective) factors, we recently examined the probability of ASD as a function of normative sex-related phenotypic diversity in brain structure (9). To do so, we initially developed a multivariate predictive model of biological sex based on differential patterns of brain structure in a sample of typically developing (TD) males and females. This model of normative sex differences was subsequently applied to males and females with ASD, which allowed us to (i) examine the probability of ASD along a phenotypic axis, ranging from the characteristic female to male brain phenotype, and (ii) identify patterns of sex-related neuroanatomical variability associated with low and high probability of ASD. In this previous study we reported that normative sex-related phenotypic diversity in brain structure significantly affects the risk of ASD in addition to biological sex alone, with male-atypical neuroanatomical characteristics carrying a higher intrinsic risk for ASD than female-atypical characteristics. This increased risk was predominantly driven by a phenotypic shift of the brain in females with ASD for a male-atypical rather than female-atypical presentation. Moreover, using a mapping approach that enabled us to estimate patterns of atypically at the level of each individual subject (10), we demonstrated that high and low risk of ASD is associated with particular patterns of neuroanatomical variability that included many of the brain regions where males typically differ from females on average (e.g. (11, 12)). Taken together, these findings highlight the need for considering normative sex-related
phenotypic diversity in addition to genetic and environmental factors when estimating an individual's risk for ASD.

However, in this prior 'proof-of-concept' study, our analysis was based exclusively on measures of cortical thickness (CT) as previous neuroimaging studies have shown CT to be highly variable between TD males and females (11, 12), and significantly altered in individuals with ASD (13, 14). Thus, as has been raised by others (15), it is unknown whether or how our initial findings generalize to other morphometric features that have been reported to be atypical in ASD. These include vertex-wise estimates of surface area (SA; (14)), the local gyriﬁcation index (LGI; (16, 17)), and sulcal depth (SULC; (18)). Here, we extended these findings to consider multiple aspects of brain anatomy including both volumetric and geometric cortical features in order to (i) provide a more comprehensive characterization of normative sex-related differences in brain structure, (ii) determine how diversity in sex-based neurophenotypes are associated with ASD probability, and (iii) estimate how much ASD prevalence in the general population might vary as a function of normative sex-related phenotypic diversity. We expect that our initial finding of a phenotypic shift of the brain in ASD females towards a more male-typical neurophenotype generalizes to other morphometric features, and for sex differences in cortical morphometry to be associated with population risk for ASD.

**Methods**

**Participants**

98 right-handed adults with ASD (49 males and 49 females) and 98 age and IQ matched TD controls (51 males and 47 females) aged 18-42 years were recruited by advertisement and assessed at the Institute of Psychiatry, Psychology and Neuroscience (IoPPN), London, and the Autism Research Centre, Cambridge. Approximately equal ratios of cases to controls, and males to females, were recruited within sites (Table 1). Exclusion criteria included a history
of major psychiatric disorder (e.g., psychosis), head injury, genetic disorder associated with ASD (e.g., fragile-X syndrome, tuberous sclerosis), or any other medical condition affecting brain function (e.g., epilepsy). We also excluded participants on antipsychotic medication, mood stabilizers or benzodiazepines.

All participants with ASD were diagnosed according to International Statistical Classification of Diseases, 10th Revision (ICD-10) research criteria. Clinical diagnoses were confirmed using the Autism Diagnostic Interview Revised (ADI-R: (19)) or the Autism Diagnostic Observation Schedule (ADOS: (20)). All ADI-R diagnosed participants reached algorithm cut-offs in the three domains of the ADI-R, although failure to reach cut-off in one domain by maximally two points was permitted. Due to reliable informants being unavailable, we were unable to obtain ADI-R diagnostic data from five females with ASD, for whom diagnostic status was confirmed using ADOS-G cut-offs for ASD. In all other participants, ADOS-G scores were used to assess the severity of current symptoms and were not used as an inclusion criterion. One female with ASD fell short one point on the ADI-R communication and repetitive domain but met ADOS-G criteria. Overall intellectual ability was assessed using the Wechsler Abbreviated Scale of Intelligence (21). All participants had a full-scale IQ greater than 80 and gave informed written consent in accordance with ethics approval by the National Research Ethics Committee, Suffolk, UK.

Structural MRI Data Acquisition

Scanning took place at the IoPPN, London, and the Addenbrooke's Hospital, Cambridge, using matched 3T GE Signa systems (General-Electric, Milwaukee, USA). A quantitative T1-mapping sequence was used to ensure standardization of structural MRI scans across scanner platforms (22, 23), which resulted in high-resolution structural T1-weighted inversion-recovery images, with 1x1x1mm resolution, a 256x256x176 matrix, TR=1800ms, TI=50ms, FA=20°, and FOV=25cm.
Cortical Reconstruction using FreeSurfer

FreeSurfer v5.3.0 software (http://surfer.nmr.mgh.harvard.edu) was used to derive tessellated models of the cortical surface for each T1-weighted image. These well-validated and fully automated procedures have been extensively described elsewhere (e.g. (24-28)). Resulting surface models were visually inspected for reconstruction errors. Surface reconstructions with visible inaccuracies were further excluded from the statistical analysis and are not described in this study. Dropout rates were equal across groups and were <10% of the total sample.

From each individual’s surface reconstructions four separate vertex based morphometric cortical features were calculated. (i) Cortical thickness (CT), i.e. the closest distance between corresponding vertices between the grey-white matter and gray matter-cerebrospinal fluid boundaries (28); (ii) vertex-wise estimates of pial (i.e. grey matter) surface area (SA) (29). And measures of cortical geometry including (iii) the local gyrification index (GII) (30) and (iv) sulcal depth (SULC, i.e. average convexity, the depth/height of each vertex above the average cortical surface). Prior to further analysis all features (excluding GII) were smoothed using a 15mm surface-based FWHM Gaussian kernel. As GII utilizes SA values sampled within spherical regions of interest with a 15mm diameter, no additional smoothing was performed for this measure.

Gaussian Process Classification

Gaussian Process Classification (GPC) (31, 32) models were employed to discriminate biological sex among TD participants on the basis of each of the four cortical features (i.e. CT, SA, GII, SULC) separately. These classifiers provide a predictive class probability ($\hat{p}_Y$) for each individual ranging from 0 to 1 for the characteristic female to male brain phenotype, with $\hat{p}_Y=0.5$ serving as the binary cut-off between classes. The overall classification accuracy of each model was estimated using leave-one-out cross validation and tested for significance by repeating the cross-validation procedure $n=1,000$ times after randomly permuting the class labels. Subsequently, we applied these four classifiers for biological sex in TD participants to
predict the sex of males and females with ASD. The resulting classification accuracies for the ASD cases were tested for statistical significance using permutation testing as outlined above.

To visualize the patterns of brain regions representative of normative sex differences, we present spatial representations of the maximum a-posteriori estimate of each model's weight vector ($w$), which is indicative of the regional contribution to the overall prediction of biological sex (32). For each classifier, $w$ was derived by training the GPC on all TD individuals. As individuals with ASD were not included in the initial training of the classifiers they do not contribute to $w$, thus we used a “predictive mapping” approach (10) to determine which cortical features are driving sex related differences in ASD among groups of males and females with high and low probability for ASD. Here, groups of high and low ASD probability were determined according to predicted class probabilities, with $y_x>0.5$ (i.e. male-typical phenotype) representing high and $y_x<0.5$ (i.e. female-typical phenotype) representing low probability. For each test example, in both male (and female) high (and low) probability groups, we calculated the element-wise (i.e. Hadamard) product of each individual's data ($x_i$) and the corresponding weights ($w$), so that $(x_i) \cdot w$ (10). For each group (i.e. male and female, high and low ASD probability) brain regions contributing the most to the classification were summarized by taking the mean of the $(x_i) \cdot w$ values at each vertex.

**Estimating ASD Probability**

To examine how normative sex based phenotypic diversity in brain structure is associated with probability for ASD we first calculated the mean of each individual's predictive class probabilities across the four classification models. This provided a single class probability for each individual that takes into account normative sex variability across all four cortical features. We then converted the continuous axis of class probabilities into a set of discrete bins (e.g., from 0 to 1 in steps of 0.125). Within each bin, the sample probability of ASD was calculated as the ratio of the number of individuals with ASD relative to the total number of individuals within the bin. Confidence intervals for these sample probabilities were determined using an exact binomial test implemented in R Project for Statistical Computing.
Furthermore, these data allowed us to estimate the population prevalence of ASD given a male- or female-typical neuroanatomical phenotype in addition to being biologically male or female. Details of these methods are provided in the supplementary materials. In brief, using Bayes theorem, we combined our sample prevalence estimates with previously published prevalence rates of ASD for biological male and female individuals in the general population (i.e., 1:42 for male and 1:189 for female individuals; (33)). In this way, we were able to estimate the population prevalence of ASD diagnosis (D = 1) given an individual’s predicted neuroanatomical sex phenotype (M) and biological sex (S), that is P(D = 1 | M, S).

**Vertex-wise between-group comparison of cortical measures**

Vertex-wise statistical analyses of each cortical feature were conducted using the SurfStat toolbox (www.math.mcgill.ca/keith/surfstat/) for MATLAB (R2014a, The Mathworks, Massachusetts). For the comparisons between TD males and females, parameter estimates were obtained by regression of a GLM at each vertex i with biological sex and center as categorical fixed-effects factors, and total grey matter (GM) volume as a continuous covariate. To examine whether the neuroanatomy of ASD is significantly modulated by biological sex, we also included a main effect of diagnostic group and a group-by-sex interaction, i.e.

\[ Y_i = \beta_0 + \beta_1 \text{Group} + \beta_2 \text{Sex} + \beta_3 (\text{Group} \times \text{Sex}) + \beta_4 \text{GM}_i + \beta_5 \text{Center} + \epsilon_i \]

where \( \epsilon_i \) is the residual error. Additionally, total SA was used as a continuous covariate for analyses of SA. Effects of interest were estimated from the coefficients \( \beta_{1,3} \), normalized by the corresponding standard error. Corrections for multiple comparisons were performed using a "random field theory" (RFT)-based cluster analysis for non-isotropic images at a cluster-threshold of \( p<0.05 \) (two-tailed) (34).
Results

Participant Demographics and Global Brain Measures

There were no significant differences between all individuals with ASD and TD controls in age (t(194) = -0.53, p = 0.598), full-scale IQ (t(194) = 1.72, p = 0.086), mean CT (t(194) = 0.08, p = 0.936), mean SA (t(194) = -0.03, p = 0.976), mean AGI (t(194) = 0.24, p = 0.810), mean SULC (t(194) = -0.67, p = 0.499), and total GM volume (t(194) = 0.20, p = 0.839). There were also no significant differences between males and females in age (t(194) = -0.93, p = 0.356), full-scale IQ (t(194) = 1.87, p = 0.063), mean CT (t(194) = -0.95, p = 0.344), or mean SULC (t(194) = -2.80, p = 0.006) across diagnostic groups. As expected, total grey matter volume was significantly larger in males than in females (t(194) = 9.11, p < 0.001). Furthermore, across the entire sample males showed significantly greater mean SA (t(194) = 9.43, p < 0.001) and mean AGI (t(194) = 5.56, p < 0.001) than females. There were no significant differences in any of these measures between TD and ASD males, or between TD and ASD females (p=0.05, 2-tailed).

Prediction of Biological Sex Based on Normative Variability in Cortical Morphometry

Across all four vertex based cortical features GPC was able to separate male from female TD controls at cross validated accuracies significantly above what would be expected by chance (SA 78%, p = 0.001; CT 71%, p = 0.001; SULC 74%, p = 0.001; AGL 77%, p = 0.001). Using each classifier’s predictive class probabilities for individuals we were able to represent normative sex based variability in brain phenotypes along a singular phenotypic axis, ranging from the female- to male-typical neurophenotype (Figure 1). For each morphological measure, the GPC model weights (w) followed a spatial pattern similar to the distribution of group differences observed between TD males and females as determined via a conventional GLM (Figure 1, Supplementary Figure 1, Supplementary Table 1).
Prediction of Biological Sex Among Individuals with ASD

When predicting biological sex for individuals with ASD we found that females with ASD were more frequently allocated to the male rather than the female class in all four models (SA 71%, \( p=0.002 \); CT 80%, \( p=0.002 \); SULC 65%, \( p=0.009 \); JGI 78%, \( p=0.001 \)). Females with ASD were significantly allocated to the male class more frequently than would be expected based on the typical rates of misclassification for female controls (SA \( \chi^2=26.41, p<0.001 \); CT \( \chi^2=22.15, p<0.001 \); SULC \( \chi^2=13.65, p<0.001 \); JGI \( \chi^2=26.01, p<0.001 \)). Furthermore, the mean across the predicted class probabilities for ASD females was significantly increased in females with ASD relative to female controls, indicating a shift towards male-typical morphological patterns in the brain (SA \( r=0.158, p=0.001 \); CT \( r=0.13, p=0.001 \); SULC \( r=0.44, p=0.001 \); JGI \( r=0.85, p=0.001 \)). Across all four models there were no significant differences (\( p<0.05 \)) between the predicted class probabilities for TD and ASD males, or between males and females with ASD.

ASD Probability as a Function of Normative Sex Variability in Cortical Morphometry:

Within our sample we found that the number of individuals with ASD significantly increased along the female to male phenotypic axis (i.e. from female to male) as defined by predicted class probabilities. This effect was predominantly driven by the shift of females with ASD towards a more male-typical neueroanatomical brain phenotype (See Figure 2, Supplementary Figure 2). When these sample probabilities were combined with previously reported ASD prevalence rates in the general population among males (1/43, i.e. 2.3%) and females (1/190, i.e. 0.5%) (33), we found that biological females with male-typical brain morphology (0.1% estimated prevalence) were twenty six times more likely to have ASD compared to biological females with female-typical neueroanatomical phenotypes (2.6% estimated prevalence). Biological males with male-typical brain morphometry showed an estimated population prevalence of 2.4% in line with previously reported rates. However, males with female-
typical phenotypes showed a reduced prevalence estimate for ASD (1.9%) indicative of a ‘female protective effect’ in relation to brain anatomy (see Supplementary Table 2).

Predictive maps identifying patterns of neuroanatomical variability associated with high and low probability of ASD across sexes are shown in Figure 3. While a similar set of brain regions conferred high (and low) probability for ASD in men and women within features, certain brain regions were unique to or differed in sign between particular risk groups and sexes. For example, negative predictive mapping values for CT in the left parahippocampal gyrus and SU/LC in the superior temporal gyrus were observed across all males and high-probability females, while low probability females showed positive values in these regions. For predictive mapping measures of SA and iGl, regions of the inferior and superior temporal gyri appear to show a sex-specific relationship with high probability males showing negative values compared to positive values amongst females and low probability males. This indicates that morphometric measures in particular brain regions that are important for classifying sex differences differ between individuals with high and low risk for ASD and may be sex-specific.

Cortical Morphometry in ASD is Modulated by Biological Sex

Lastly, we used a conventional GLM to test for significant between-group differences and diagnosis-by-group interactions in the four examined morphometric features. Overall, we found that measures of CT, SA, and iGl in ASD were significantly modulated by biological sex (Figure 4, also Supplementary Figure 3). Such group-by-sex interactions therefore need to be taken into account when interpreting main effects of group. Significant group-by-sex interactions in measures of CT were observed across the temporal lobe, including the fusiform and parahippocampal gyri. Across this cluster the degree of CT abnormality in females with ASD exceeded those of males, despite no differences in measures of ASD.
symptoms severity (9). Additionally, group-by-sex interactions were observed for measures of SA and jGII in the medial orbital frontal cortex showing decreases in females with ASD and increases in males with ASD. Statistical details of these clusters are provided in Supplementary Table 3.

Discussion

The present study builds upon a previous 'proof-of-concept' suggesting that probability of ASD increases significantly from a characteristic female to male neuroanatomical brain phenotype (9). In this prior study, which examined measures of cortical thickness (CT) only, the increase in ASD probability was predominantly driven by a phenotypic shift of the brain in females with ASD towards a male-typical neurophenotype (9). In the current study, we show that this phenotypic shift is not restricted to measures of CT, but generalizes to other cortical features including vertex-wise estimates of surface area (SA), the local gyrification index (jGII), and sulcal depth (SULC). The findings of the present study therefore not only support our original report in suggesting that male-typical neuroanatomical characteristics carry a higher probability for ASD than female-typical characteristics, but further show this effect to be broadly and consistently related across multiple measures of cortical morphometry.

The extension of our approach to include multiple morphometric features is of crucial importance as there is evidence to suggest that different cortical features are etiologically and phenotypically distinct. For example, it has been shown that measures of CT and SA (i) are mediated by different sets of genes (35), (ii) have different developmental trajectories (36), and (iii) represent distinct aspects of the cortical architecture (37). Therefore, the observation that the phenotypic shift of the brain in ASD females is consistent across morphometric features may indicate the existence of a common underlying mechanism, such as atypical
cortical neurogenesis and migration (38), that steers multiple aspects of brain anatomy towards a similar phenotypic endpoint overall. Yet, the neuroanatomical patterns associated with the prediction of biological sex, and the patterns carrying high/low ASD probability in males and females, differed in sign and regional distribution between the cortical features.

Thus, while there may be a common etiology for the phenotypic differences in brain anatomy in ASD within the sexes, our findings are consistent with the notion that sex differentiation mechanisms in the brain have region specific effects (for review see (39, 40)). Therefore, differences in these neuroanatomical patterns associated with high probability for ASD may be beneficial in identifying brain regions where sexual differentiation may contribute to male preponderance in ASD.

Moreover, we established that the neuroanatomy of ASD in some regions of the brain is modulated by biological sex by examining sex-by-diagnosis interactions for multiple cortical features. This suggests that in these brain regions (i) there are not only quantitative differences in the degree of neuroanatomical abnormality between males and females with ASD and their respective normative population, but also that (ii) ASD potentially interacts with sexually-differential neurobiological phenotypes to give rise to qualitative differences in brain anatomy between males and females with the condition. This is also relevant in view of a recent genetic study, suggesting that while ASD risk genes are expressed at similar levels in males and females, the impact of these genes is likely modulated by their interactions with sex-differential processes (e.g. in cortical microglia and/or astrocytes) that may play a critical role in setting males at a greater risk for ASD (41). An individual’s risk or probability of ASD is therefore not only dependent on genetic load itself, but also on naturally occurring sex-differential biological pathways.

With respect a biological organ as complex as the brain it has been suggested that on a phenotypic level, the brain should be considered a ‘mosaic’ of regions that differ between
degrees of relative maleness or femaleness, or not at all (i.e. unisexual), resulting in a potentially infinite variety of brain phenotypes both within and between sexes (39, 40). In order to identify an individual’s position on such a multi-dimensional “spectrum” of sex-related phenotypic variability, it is important to develop analytical frameworks that can accommodate inter-individual phenotypic diversity within and across the binary categories dictated by biological sex, such as the probabilistic multivariate pattern classification approach utilized in the present study (see also (9)). In this way we were able to estimate an individual’s risk of ASD based on phenotypic, rather than biological sex and in doing so demonstrated that an individual’s probability for developing ASD may vary from as much as 0.1 to 2.6% in the general population depending on having a female- or male-typical neuroanatomical phenotype, in addition to being biologically male or female. Our approach to modeling normative sex-related phenotypic diversity may therefore be extended to various additional phenotypic features in the future, and to contribute to the development of multivariate disease liability models for complex neurobiological phenotypes.

Despite the advantages of the current approach over the original proof-of-concept study, a number of limitations remain. Most importantly, it is crucial to highlight that while we were able to show a statistical association between normative sex-related phenotypic diversity in brain structure and ASD probability, this relationship should not be interpreted as being ‘causal’. Future studies will be needed to uncover the genetic and/or neurobiological mechanisms underpinning this relationship (e.g. (41-43)). Second, due to the limited range of clinical/behavioral measures acquired in the current study, we were unable to determine the functional relevance of our findings. In the future it will be crucial to identify how diversity in normative sex-related morphometric phenotypes relates to sex-differences in general cognitive and/or behavioral profiles (44, 45), or to different clinical ASD phenotypes (3, 4, 6, 7). Last, our study was based on a sample of adults with ASD without learning disability using a cross-sectional study design. The fact that our sample is not representative of the
entire ASD population limits the generalizability of our sample probability estimates for ASD to the general population. Furthermore, our population estimations are based on previously reports of male preponderance in ASD being roughly a 4:1 male to female ratio (33), however other studies have shown this ratio may actually to be closer to 2:1 (8). The nature of our sample means further studies will thus be required to (i) replicate our findings in other subgroups on the autism spectrum (e.g. in children and adolescents, or individuals with a learning disability), and (ii) to identify how stable (or variable) our neuroanatomical indicators of biological sex are across different stages of development.

Conclusions

Taken together, our findings suggest that the male-typical neurobiological brain phenotype carries a higher probability for ASD than the female-typical phenotype across the binary categories dictated by biological sex, and across a range of neuroanatomical features that include both volumetric as well as geometric measures. In addition to genetic and environmental factors, multiple morphometric features characteristic of normative phenotypic diversity in brain structure should therefore be considered when investigating risk and resilience for ASD.
### Table 1. Participant Demographics and Global Brain Measures

<table>
<thead>
<tr>
<th></th>
<th>ASD (n = 98)</th>
<th>TD Control (n = 98)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Male Participants (n = 100)</strong></td>
<td>n = 49</td>
<td>n = 51</td>
</tr>
<tr>
<td>Age [years]</td>
<td>26.16 ± 7.2 (18 - 41)</td>
<td>27.22 ± 5.58 (18 - 42)</td>
</tr>
<tr>
<td>Full-scale IQ (WASI)</td>
<td>112.61 ± 12.27 (89 - 135)</td>
<td>114.67 ± 10.88 (93 - 137)</td>
</tr>
<tr>
<td>Verbal IQ</td>
<td>110.51 ± 13.01 (83 - 137)</td>
<td>109.86 ± 11.03 (88 - 137)</td>
</tr>
<tr>
<td>Performance IQ</td>
<td>112.00 ± 13.65 (85 - 138)</td>
<td>116.74 ± 10.90 (93 - 133)</td>
</tr>
<tr>
<td>ADE-R Social</td>
<td>17.57 ± 5.50 (10 - 28)</td>
<td>-</td>
</tr>
<tr>
<td>ADE-R Communication</td>
<td>13.86 ± 4.16 (8 - 24)</td>
<td>-</td>
</tr>
<tr>
<td>ADE-R Repetitive Behavior</td>
<td>4.96 ± 2.37 (1 - 10)</td>
<td>-</td>
</tr>
<tr>
<td>ADOS Social + Communication</td>
<td>9.43 ± 4.39 (1 - 21)</td>
<td>-</td>
</tr>
<tr>
<td>ADOS Stereotypic Behaviour</td>
<td>1.27 ± 1.27 (0 - 5) *</td>
<td>-</td>
</tr>
<tr>
<td>Mean Cortical Thickness [mm]</td>
<td>2.35 ± 0.10</td>
<td>2.33 ± 0.10</td>
</tr>
<tr>
<td>Mean Surface Area</td>
<td>0.7680 ± 0.71</td>
<td>0.7683 ± 0.05</td>
</tr>
<tr>
<td>Mean local Gyrunification Index</td>
<td>3.04 ± 0.12</td>
<td>3.04 ± 0.10</td>
</tr>
<tr>
<td>Mean Sulcal Depth</td>
<td>0.0341 ± 0.007</td>
<td>0.0356 ± 0.006</td>
</tr>
<tr>
<td>Total Grey Matter Volume [litre]</td>
<td>0.76 ± 0.08</td>
<td>0.76 ± 0.05</td>
</tr>
<tr>
<td>Total Intracranial Volume [litre]</td>
<td>1.59 ± 0.21</td>
<td>1.58 ± 0.17</td>
</tr>
<tr>
<td></td>
<td>Female Participants ($n = 96$)</td>
<td>$n = 49$</td>
</tr>
<tr>
<td>----------------------</td>
<td>--------------------------------</td>
<td>----------</td>
</tr>
<tr>
<td>Age (years)</td>
<td>$27.60 \pm 7.12$ (18 - 48)</td>
<td>$27.60 \pm 7.31$ (19 - 52)</td>
</tr>
<tr>
<td>Verbal IQ*</td>
<td>$116.60 \pm 12.15$ (76 - 144)</td>
<td>$117.83 \pm 8.81$ (96 - 135)</td>
</tr>
<tr>
<td>Performance IQ*</td>
<td>$110.56 \pm 14.50$ (67 - 138)</td>
<td>$114.38 \pm 8.26$ (96 - 128)</td>
</tr>
<tr>
<td>Full-scale IQ (WASI)</td>
<td>$114.88 \pm 12.36$ (84 - 136)</td>
<td>$118.38 \pm 7.32$ (99 - 129)</td>
</tr>
<tr>
<td>ADOS-R Social</td>
<td>$16.32 \pm 4.24$ (10 - 26)</td>
<td>-</td>
</tr>
<tr>
<td>ADOS-R Communication</td>
<td>$12.43 \pm 3.96$ (7 - 22)</td>
<td>-</td>
</tr>
<tr>
<td>ADOS-R Repetitive Behavior</td>
<td>$4.36 \pm 1.98$ (1 - 9)</td>
<td>-</td>
</tr>
<tr>
<td>ADOS Social + Communication</td>
<td>$7.71 \pm 5.28$ (0 - 19)</td>
<td>-</td>
</tr>
<tr>
<td>ADOS Stereotypic Behaviour</td>
<td>$0.75 \pm 0.97$ (0 - 3) *</td>
<td>-</td>
</tr>
<tr>
<td>Mean Cortical Thickness (mm)</td>
<td>$2.32 \pm 0.11$</td>
<td>$2.34 \pm 0.10$</td>
</tr>
<tr>
<td>Mean Surface Area</td>
<td>$0.6895 \pm 0.05$</td>
<td>$0.6865 \pm 0.06104$</td>
</tr>
<tr>
<td>Mean local Gyrification Index</td>
<td>$2.95 \pm 0.09$</td>
<td>$2.96 \pm 0.10$</td>
</tr>
<tr>
<td>Mean Sulcal Depth</td>
<td>$0.0878 \pm 0.008$</td>
<td>$0.0880 \pm 0.007$</td>
</tr>
<tr>
<td>Total Grey Matter Volume [litre]</td>
<td>$0.67 \pm 0.06$</td>
<td>$0.68 \pm 0.06$</td>
</tr>
<tr>
<td>Total Intracranial Volume [litre]</td>
<td>$1.32 \pm 0.18$</td>
<td>$1.32 \pm 0.17$</td>
</tr>
</tbody>
</table>

Note: Data expressed as mean ± standard deviation (range). There were no significant between-group differences in age, FSIQ, mean cortical thickness, total grey matter volume, or
total intracranial volume ($p<0.05$, two-sided). ADI-R values based on 49 males and 44 females

with ASD. ADOS values based on 49 males and 48 females. * statistically significant between

males and females based on $p<0.05$, two-sided ($t(95)=2.23$, $p=0.028$). + verbal and performance

IQ data was not available for one ASD female
Figure 1. Gaussian Process Classification of Biological Sex: The results of Gaussian Process Classification (GPC) according to biological sex categories are shown for each of the four cortical features investigated: (A) cortical thickness (CT), (B) surface area (SA), (C) the local gyrification index (IGI), and (D) sulcal depth (SULC). Each model was able to separate TD males from females at accuracies significantly above what would be expected by chance (top left). When these models of normative sex differences in cortical morphology were applied to males and females with ASD, females with ASD were more frequently classified to the male class (bottom left) indicating a male-typical morphological presentation amongst females with ASD. Spatial representations of each discriminative model (i.e. weight vectors) are shown (right). These patterns largely corresponded with brain regions that differed between TD males and females as identified using a conventional general linear model (see supplementary Figure 1).

Figure 2. Probability for ASD as a Function of Normative Sex-Related Phenotypic Variability in Brain Morphology: Probability estimates for ASD across eight discrete bins along the axis of probabilistic class predictions for biological sex are plotted for (A) biological females and (B) males. Each individual’s probabilistic class prediction was calculated as the mean prediction across all four classification models. Probability for ASD was seen to increase along with increasingly male-typical class predictions. This relationship was particularly apparent in females due to the statistically significant shift in females with ASD towards the male-typical neuroanatomical phenotype.

Figure 3. Prediction Maps for High and Low ASD Probability Males and Females: Prediction maps highlighting brain regions most relevant to classifying biological sex among high and low risk males and females are shown for each cortical feature. (A) cortical thickness (CT), (B) surface area (SA), (C) local gyrification index (IGI), and (D) sulcal depth (SULC). For male and female individuals high and low probability groups were determined according to class probabilities with <0.5 (i.e. female-typical) indicating low and >0.5 (i.e.
male-typical) indicating high probability. For each subgroup, the product of the decision function for classifying biological sex (w) and each individual’s data (x_i) was calculated. Mean values at each vertex were then used to summarize maps within subgroups.

Figure 4. Group by Sex Interaction Effects in Cortical Morphometry: Clusters of significant (RFT p<0.05) group x sex interactions in (A) cortical thickness (CT), (B) surface area (SA), and (C) the local gyrification index (LGI). No significant group x sex interactions were observed in measures of sulcal depth (SULC). Statistical details of these clusters are provided in Supplementary Table 3.
References


38. Casanova MF. Autism as a sequence: from heterochronic germinal cell divisions to abnormalities of cell migration and cortical dysplasias. Medical hypotheses. 2014; 83(1):32-3


259x296mm (300 x 300 DPI)