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Abstract

Autism Spectrum Disorder (ASD) is a construct used to describe individuals with a specific combination
of impairments in social communication and repetitive behaviours, highly restricted interests and/or
sensory behaviours beginning early in life. The worldwide prevalence of ASD is just under 1%, but
estimates are higher in high-resource countries. Although gross brain pathology is not characteristic of
ASD, subtle anatomical and functional differences have been observed in postmortem, neuroimaging
and electrophysiological studies. Initially it was hoped that accurate measurement of behavioural
phenotypes would lead to specific genetic subtypes, but genetic findings have mainly applied to
heterogeneous groups that are not specific to ASD. Psychosocial interventions in children can improve
specific behaviours, such as joint attention, language and social engagement that may affect further
development and could reduce symptom severity. However, further research is necessary to identify the
long-term needs and treatments and the mechanisms behind them that could result in improved
independence and quality of life over time. Families are often the major source of support for people
with ASD throughout much of life and need to be considered, along with the perspectives of autistic persons, in both research and practice.

[H1] Introduction

Autism spectrum disorder (ASD) is a common, highly heritable and heterogeneous neurodevelopmental disorder that has underlying cognitive features and commonly co-occurs with other conditions. The behaviours, strengths and challenges of people with autism, or ASD, have attracted the attention of scientists and clinicians for at least 500 years (Fig. 1). ASD is a heterogeneous disorder and, reflecting this heterogeneity, the term autism has been used in various ways to describe both a broader presentation, and then a specific diagnosis when it was considered to be one subgroup within the general diagnostic category of ‘pervasive developmental disorders’ (PDDs), a group of disorders that was introduced in Diagnostic and Statistical Manual of Mental Disorders, Third Edition (DSM III) in 1980 to convey the idea of a broader spectrum of social communication deficits. Owing to of lack of clear borders between the PDDs and difficulties in reliably distinguishing them, the current diagnostic systems, the International Classification of Diseases 11th Revision (ICD-11) and the DSM-5 use the umbrella term ‘ASD’, and differentiate individuals using additional clinical specifiers and modifiers.

Manifestations of ASD include impairments in social communication and interaction, sensory anomalies, repetitive behaviours and varying levels of intellectual disability (Table 1). Together with these core symptoms, co-occurring psychiatric or neurological disorders are common in people with ASD, of which, hyperactivity and attention disorders (such as attention-deficit/hyperactivity disorder (ADHD)), anxiety, depression and epilepsy are fairly prevalent. A diagnosis of ASD is reached after obtaining a detailed developmental history, often from the parents, and observation of the individual interacting with parents or other individuals 1,2. Early intervention for children with ASD is key owing to common difficulties in communication. The types of interventions used change throughout life and include parent-mediated interventions and/or therapist-delivered interventions in childhood, school-based strategies and techniques to promote independence in adulthood. Pharmacological therapies can be used to treat some of the associated symptoms of ASD, such as irritability, and comorbidities, such as anxiety.

This Primer discusses the epidemiology and mechanisms of ASD, together with the diagnosis and treatment of people with this condition. Three themes are addressed: mechanisms of causality and change over time, heterogeneity within and between individuals with ASD, and outcomes across the lifespan.

[H1] Epidemiology

[H2] Prevalence [Au: subheading introduced for flow, OK?YES]

Epidemiological administrative and community-based [Au:OK?YES] studies have suggested that ASD is more common in males than in females, with reported ratios [Au:OK to add ‘with reported ratios’ here?] ranging from 2.1–5.1, with an estimate of 4.1 in the 2010 [Au:OK?] Global Burden of Disease study3,4. The sex ratio is slightly lower in studies that use population-wide testing [Au: I’ve edited ‘screening’ to ‘population-wide testing’ here based on this comment. Do we need to edit the text in the Diagnosis, screening and prevention section too?] to find community cases within a population compared with the
more common passive case-finding studies that review administrative data (for example, medical or special educational records), and that may result in less plausible associations and, therefore, artificially increase prevalence estimates.\(^5\) Active case-finding that does not rely on administrative records has demonstrated an equivalent community rate of ASD in men and women with moderate to profound intellectual disability\(^4\). Thus, even the most widely accepted tenet of our understanding of factors associated with ASD is far from straightforward.

Estimates of the prevalence of ASD in various populations and settings differ according to the method of ascertainment used in the study, including definition, sampling and the extent of independent population case assessment. Of note, the Global Burden of Disease study uses all known data from administrative and community survey sources on a disease or disorder to model associations (particularly with time) to examine trends. In the 2010 GBD study [Au:OK?], an estimated 52 million people had ASD globally, equating to a prevalence of 1 in 132 individuals\(^6\). Worldwide, little interpretable variation in the prevalence of ASD between regions or ethnicities has been reported [Peer reviewer comment: It would be useful to add a few comments about prevalence estimates around the world and how they differ based upon resources of the countries (i.e., are likely not due to regional variations in prevalence or differing environmental conditions, but instead reflect access to services). This could enhance the paper’s global perspective. [Au: could we quote additional data from the GBD study here? If these data are lacking I would mention this here so it’s clear to readers].] Indeed, one systematic review did not find a strong effect of ethnic, cultural or socioeconomic factors on the prevalence of autism [Au:OK?]. However, statistical power to detect any effects was limited in the available data sets, particularly in low-income countries. An increased prevalence of ASD has been reported in migrant groups in some studies\(^8\) with few clear factors that might contribute to a greater prevalence in an Afro-Caribbean population in higher income countries\(^9,10,11\) in the absence of any evidence of geographical variation.\(^7\) However, a survey of adults in the general population has shown that rates of ASD in black and minority ethnic groups may be lower than in the rest of the population\(^12\); data from indigenous and Aboriginal cultures are very limited.

Many individuals and groups presume that autism rates are increasing over time, but this supposition is based on data from administrative records rather community-based studies. Indeed, after accounting for methodological variations between studies, there was no clear evidence of a change in the prevalence of ASD in the community between 1990 and 2010\(^13\). In addition, general population and systematic case-finding community-based [Au:OK?] surveys (including testing of representative populations) have also confirm the lack of significant change in prevalence rates in childhood\(^14\) and adulthood\(^15\) over time. No significant evidence is available supporting that ASD is rarer in older people, which provides further evidence against the suggestion that ASD is increasing in prevalence over time\(^4\). Even in high-income countries with strong ASD public health policies, there is evidence that ASD in adults goes largely unrecognized, whereas administratively recorded diagnoses in children increase year by year\(^16\). This finding highlights the importance of obtaining information on ASD rates in settings where professionals may be able to improve its recognition. The prevalence of ASD in mental health inpatient settings is estimated to be far higher than in the general population, ranging from 4–9.9\(^%\)\(^17\).

[H2] Environmental factors
One review of systematic reviews and meta-analyses of environmental risk factors for ASD included a comprehensive coverage of the literature, a discussion of the limitations of research and the need for long-term prospective cohort-based studies to begin to address these limitations\(^{18}\) (Fig. 2). This and other studies identified environmental risk factors for ASD as advanced parental age\(^{19}\) and birth trauma, particularly if due to proxies of hypoxia\(^{18}\). Moreover, maternal obesity, a short interval between pregnancies, gestational diabetes mellitus and valproate use during pregnancy have all been associated with increased risk of ASD [Au:OK?] (Fig. 2). However, it should be noted that these factors cannot be considered causal, but could be reactive, independent or contributory for ASD. Studies evaluating risk factors for ASD that have reported an absence of association are equally, if not more important, to note, including clear evidence that ASD is not associated with vaccination\(^{20}\). Other negative associations include prolonged labour, delivery by caesarian section or assisted vaginal delivery, premature rupture of membranes and the use of assisted reproductive technologies, among other factors (Fig. 2). Environmental risk factors could underlie risk of ASD through several complex underlying mechanisms, such as [Au: edits to improve narrative flow ok?] genetic and epigenetic related effects (see Mechanisms/pathophysiology, below), inflammation and oxidative stress, hypoxic and ischemic damage\(^{18}\).

**[H1] Mechanisms/pathophysiology**

Many cognitive theories have been suggested to underlie the behavioural and developmental manifestations of ASD, although the prominence and the consensus on the potential explanatory value of these theories have declined in the past decade. These theories range from ‘social first’ theories, such as the theory of mind (or mentalizing) and social motivational deficit theories, to global processing deficit theories including attentional control, executive dysfunction and weak central coherence or enhanced perceptual processing theories\(^{21,22}\). Although many of these theories had a useful descriptive role and provide potential insights into differences in how autistic individuals might process and experience the world around them, the theories pertain to neurodevelopmental disorders in general and lack specificity for ASD), largely non-developmental, applying only to a single point in time, and lack evidence as explanatory models. Nevertheless, they have been useful in clinical practice and underlie some recently proposed interventions, such as CBT-oriented treatments for anxiety\(^{23}\).

Following cohorts of infants from gestation or birth to 2 or 3 years of age (that is, when a diagnosis of ASD can be established) enables the study of the brain and behavioural manifestations of ASD as they emerge\(^{24}\). Indeed, prospective studies of infants with a relative with ASD have yielded a number of insights into the mechanisms of this disorder. For example, infants who develop ASD later in childhood have substantially typical profiles of interest in faces\(^{25}\) and eyes\(^{26}\) at 6 months of age, which have cast doubt on social orienting theories in which ASD originates from a primary deficit in innate patterns of subcortically-mediated social orienting\(^{27}\). In addition, subtle but diffuse differences in encephalography (EEG) and in other measures of brain function have been demonstrated in autistic people (see ‘Findings from electrophysiological studies’, below), which could represent alternative pathways to a common end-state phenotype or to whole-brain alterations in synaptic signalling pathways that have effects on development [Au: edits for brevity ok?]\(^{28}\). Such considerations highlight the limitations of deterministic models of ASD, in which a genetic change leads to a synaptic change that relates to a canonical symptom\(^{29}\). Rather, there
is likely a complex set of developmental interactions, in which the child’s emerging brain activity and behaviour have bidirectional relationships to synaptic signalling and gene expression.\(^{30}\)

[**H2**] Genetics

Twin and family studies consistently demonstrate that ASD has a particularly large genetic contribution, with estimated heritability ranging from \(~40\) to \(90\%\).\(^{21,32}\) In addition, one analysis demonstrated that ASD is among the most heritable common medical conditions.\(^{33}\) More than 100 genes and genomic regions have now been confidently associated with ASD\(^{34,35}\), mostly based on the study of heterozygous, germ-line, \textit{de novo} mutations. These genetic changes range in size from a single base (or nucleotide)\(^{36-38}\) to submicroscopic segments of DNA of thousands to millions of bases (also known as copy number variations (CNVs))\(^{39,40}\). Whether these genetic changes lead to alterations in the sequence of DNA or the structure of the chromosome, changes that have a functional effect on protein-coding regions of the genome have the strongest and most reliable association with ASD risk. Collectively, these \textit{de novo} heterozygous mutations are rare and confer relatively large risks of ASD.\(^{41}\) With genetic studies now including cohorts of up to tens of thousands of individuals and the associated increase in statistical power, common, transmitted alleles of modest effect size, mostly corresponding to the non-coding regions of the genome, have begun to be identified.\(^{42}\)

Studies of the genetics of ASD contrast broadly with studies of adult-onset psychiatric disorders, in which most successful gene discovery has emerged from genome-wide association studies (GWAS), which assess common alleles of small effect size. Indeed, the earliest successes in ASD presaged a more general finding that the contribution of rare, \textit{de novo} mutations in coding regions of the genome is relatively greater among a range of early-onset disorders\(^{43-45}\) than for typically later-onset common conditions such as schizophrenia and bipolar disorder, although there is also a surprising degree of overlap in genetic risk for overtly disparate neuropsychiatric phenotypes that remains to be further elucidated.\(^{31}\)

The extent to which rare, high effect size mutations account for ASD risk raises some important definitional issues. Considering the overall population, the contribution of \textit{de novo} mutations to ASD risk is quite small (\(~3\%)\(^{32}\). Indeed, the vast majority of individuals who harbour genetic risk for a common condition, particularly those with variants of small effect size, will never develop symptoms or need clinical attention. By contrast, there is a marked enrichment of individuals with rare and \textit{de novo} mutations in the clinical ASD population. Conservative estimates are that 10–20\% of people with ASD harbour a \textit{de novo} rare point mutation or CNV contributing to their presentation (34, 49,50). If the clinical population is constrained to those with ASD who are female or who have intellectual disability, multiple unaffected siblings or seizures, \(~20-30\%) have a \textit{de novo} mutation [\textbf{Au: edits for clarity based on Matt’s comment, ok?}]\(^{34}\). For example, the yield of \textit{de novo} structural and sequence variations contributing to ASD is nearly double in girls than in boys (34).

However, irrespective of the precise proportion of risk conveyed by these mutations, their most substantial contribution to the understanding of ASD is likely to be in elaborating the mechanisms of this disorder.\(^{48,49}\). In ASD, a single \textit{de novo} germ-line heterozygous loss-of-function point mutation can convey more risk than the cumulative effect of the top decile of polygenic risk for schizophrenia.\(^{47,50}\). Unfortunately, although manifestly more tractable than modelling hundreds of alleles simultaneously,
addressing a single ASD mutation at a time is not synonymous with an easy avenue to clinical care of most people with ASD.

**[H3] Molecular pathophysiology.** Over the past decade there have been many studies using model systems to recapitulate so-called single gene (or monogenic) versions of ASD, such as fragile X syndrome and tuberous sclerosis complex – which cumulatively are estimated to account for <10% of clinical cases of ASD\(^51\). In addition, more recent studies have modelled the effects of rare and *de novo* mutations identified in idiopathic ASD. This literature is far too vast to review comprehensively here \(^52,53\). Although the study of ASD risk genes in model systems has revealed a great deal about general biology, how these findings relate to the pathophysiology of ASD is less clear\(^48,49\). In general, ASD risk genes tend to have a role in multiple functions in many brain regions that unfold in a spatiotemporally defined manner across development. Consequently, although manipulation of a single risk gene in a model system may lead to interesting phenotypes—including social-behavioural phenotypes in evolutionarily distant organisms—it does not necessarily illuminate its contribution to human social disability. Moreover, although a single mutation can confer a several fold increase in the risk of ASD, these variants do not demonstrate the type of causal clarity that is associated with classic monogenic neurodevelopmental disorders, such as fragile X syndrome, Angelman syndrome, Rett syndrome or tuberous sclerosis complex. In addition, the well-established sexual dimorphism of social disability adds yet another dimension to the expansive search space that exists between risk gene and human behaviour\(^48\). The challenges of disentangling the spatiotemporal dynamics of risk gene expression and protein function are made even more difficult by the reality that these may play out differently in males versus females.

Owing to these challenges, multiple approaches have emerged focusing on convergence\(^38,40,54–57\), that is, searching for points of commonality across different ASD risk genes, with the reasoning that this approach could identify shared pathological mechanisms. In fact, the earliest successes in gene discovery quickly revealed important general properties that have held up well over time, including that, *prima facie*, most proteins encoded by ASD risk genes are involved either in synaptic structure and function or chromatin modification and regulation of gene expression\(^38,46,47,58\) (Fig. 3). More recently, there has been an additional focus on spatiotemporal convergence and several studies have supported a nexus in mid-fetal, glutamatergic neurons during cortical development, with modestly divergent findings regarding deep \(^56\) versus superficial\(^54\) cortical layers. With improvements in technology, additional regions, including striatum, have also begun to emerge as points of potential risk convergence for ASD\(^59\).

The ability to constrain future experiments to examine mutations in specific risk-associated regional, cellular and developmental contexts should allow the narrowing in on relevant mechanisms. Of note, one study used single cell technologies to examine specific cell types and developmental stages using brain tissue from people with ASD \(^60\), and demonstrated changes in transcription in multiple cell types including upper-layer cortical neurons. These types of post-mortem studies ask important but somewhat broader questions from the approaches described above, such as underlying pathology and how the brain changes and responds to pathology over time. In these studies, similar to any cross-sectional study, it can be challenging to differentiate cause from effect. Consequently, the pursuit and intersection of studies that seek to define convergence early in development and those that examine subsequent molecular, cellular and circuit level changes will be critical to illuminating pathological mechanisms.
[H2] Neurobiology

[H3] Findings from MRI Studies. This section does not summarize the structural or functional literature and focuses predominantly on prospective study designs. Broader coverage of the neurobiology of ASD is needed] [Au: I suggest adding a sentence or two at the end of this paragraph summarizing Rebecca’s comment here, so it’s clear why this section of the paper focuses on prospective studies and those in infants] MRI can facilitate understanding how the brain structurally and functionally develops differently in people with ASD, although to date, MRI results in ASD are not definitive. Although neuroimaging is typically more expensive than EEG and studies are limited by issues of replication, sometimes that is related to head motion that occurred during the scan which can erode signal\(^{61}\), structural studies including those using diffusion tensor imaging (DTI)\(^{62}\) and functional MRI (fMRI)\(^{63}\) have accelerated our understanding of how altered neural circuits relate to clinical symptoms of ASD\(^{64,65}\).

Studying circuitry in childhood that is specifically associated with the social brain (a network of brain areas involved with processing social information), including visual areas, areas of the prefrontal cortex, subcortex and areas integrating information (such as temporal parietal function and superior temporal sulcus), could also offer insight into the neural mechanisms of ASD\(^ {66}\). In addition, MRI may facilitate understanding the heterogeneity of ASD demonstrating subgroups of individuals with specific neurobiological alterations that could account for their symptomology.

The first MRI studies of ASD focused on cerebral and cerebellar grey matter and white matter volumes in young children\(^{67,68}\), although these studies were limited by studying toddlers and children \(\geq 18\) months, missing the opportunity to detect biomarkers of ASD in the first year of life. More recently, longitudinal studies have obtained multiple brain MRIs of infants at high risk of developing ASD (that is, those with a sibling with ASD; known as baby sibling studies) during their first 2 years of life, and assessed these children for ASD at this age. In these studies, detectable differences in brain structure were observed at 6 months of age in the fractional anisotropy trajectories for 12 of 15 neural fibre tracts in the brain in children diagnosed with ASD at 2 years of age compared to children not diagnosed\(^ {69}\). Furthermore, abnormal growth in the cortical surface between 6 and 12 months of age and greater brain volume between 12 and 24 months of age was seen in children who were later diagnosed with ASD, compared with those not diagnosed with ASD\(^ {70}\) (Fig. 4). In addition, white matter integrity in the genu pathway at 6 months of age predicted the presence of restricted and repetitive behaviours at 2 years of age\(^ {71}\) and computational work demonstrated that whole brain functional connectivity at 6 months of age predicted a diagnosis of ASD at 2 years of age\(^ {72}\). Collectively, these studies suggest the presence of disrupted neural pathways before the emergence of behavioural symptoms in children with ASD, and might provide clues about the underlying neural mechanisms of ASD. Although data from MRI studies has revealed differences in neurobiology between young children diagnosed with ASD and those without\(^ {73}\), given that replication has been particularly difficult in these studies, more work is required before MRI can be used as a reliable biomarker of ASD\(^ {74}\).

Task-based fMRI studies investigate circuits that are responsible for core challenges in ASD (such as language production and comprehension\(^ {75}\)), and have demonstrated hyper-activation of the superior temporal gyrus and inferior frontal gyrus and hypoactivation of the bilateral middle temporal gyrus\(^ {72}\). In addition, these studies have demonstrated challenges in processing emotions in faces and the “social brain”\(^ {74}\), and deficits in attention\(^ {75}\). Studies have also shown greater sensitivity to sensory information, showing increased connectivity between the anterior insula and sensorimotor areas, and the anterior
insula and amygdala, together was associated with greater sensitivity to slightly aversive sounds and
tactile information. Although this area of research has revealed similarities or differences in people with
ASD compared with comparison groups, it has been limited by averaging data across many individuals,
which can mask heterogeneity and differences across age groups. In addition, the work has been limited
by small sample sizes and problems with replication that is likely caused by the many challenges with MRI
data collection in people with ASD, such as differences in data processing, inter-subject variability and
data quality. Longitudinal imaging, as well as associating neuroimaging data with longitudinal
behavioural outcomes can address some of these limitations characterizing differences within
participants.

Resting state functional connectivity MRI studies that require participants to look at a blank screen with
no task demands have been used to study intrinsic connections in the human brain. Large datasets, such
as the Autism Brain Imaging Date Exchange (ABIDE), have enabled researchers to pool data to allow
more highly powered studies to address known limitations of small sample sizes and many dataset have
relied on resting state studies to study neural connectivity in ASD. In these studies, evidence has emerged
of both hyper-connectivity and hypo-connectivity in short-range and long-range connections throughout
the brain. Differences in results between studies could be due to the age of the participants, sex
differences, heterogeneity, methodological concerns or that both connectivity states exist in ASD.

In future, MRI could be well suited to categorize subgroups of ASD, as well as parsing out commonalities
and distinctions among other developmental disorders. Using MRI to better understand differences
between boys and girls on the spectrum, such as differences in whole brain connectivity or the social
brain, a field in its infancy, or as a marker of biological change due to treatment has growing interest.

**[H3] Findings from electrophysiological studies.** EEG has been historically used for the
diagnosis of comorbid epilepsy in people with ASD, although it can also be used to study the mechanisms
of ASD. Compared with MRI, EEG is more economical, easier to use and less invasive—which is particularly
important for paediatric populations—whilst granting access to brain dynamics at millisecond timescales.
Magnetoencephalography (MEG), although more expensive, provides higher spatial resolution than EEG.

Since the early recordings, the first focus of quantitative EEG was to study people with ASD in task-free
conditions. Pioneering studies have revealed alterations in oscillatory activity during the resting state in
people with ASD, with more slow waves and less alpha waves, as well as less intra-hemispheric and inter-
hemispheric asymmetry compared to people without ASD. More recent work has demonstrated the
presence of developmental trajectories as revealed through increasingly sophisticated spatio-spectral
analyses, and has revealed how differences in the trajectories of EEG power in high-risk infants may
represent an endophenotypes of ASD.

In terms of mechanisms, other studies have started to focus on task-based modulation of cognitive
function, such as low-level perceptual anomalies and action observation that relate to the ASD phenotype.
One theory proposing a specific failure in ASD of the ability of the brain to ‘mirror’ observed actions of
another person (thereby named the ‘broken mirror’ theory) was based on altered μ-wave suppression in
ASD, but was later questioned both theoretically and empirically, pointing toward a more complex
picture of dysfunctional executive functions and visual attention. Other studies, particularly those
assessing event-related potentials (ERP), have demonstrated the modulation of sensory processing in
people with ASD, with observed changes in sensitivities and latency. Differences in auditory and visual
processing could have a role in the development of core features of ASD, such as language delay and
difficulty in emotion recognition although this hypothesis requires further study. Although perceptual
processes appear different in people with ASD, the electrophysiological underpinning is still far from clear
regarding the main ERPs like the MisMatch Negativity (MMN)\textsuperscript{100} or the N170\textsuperscript{101}. Although data from
metanalyses have suggested smaller MMN amplitudes and delayed N170 latencies on average in people
with ASD compared to typically developing controls, additional studies are required that account for the
large heterogeneity of this disorder, by moving away from averaging the data to focus either on specific
subgroups\textsuperscript{102} or refined modelling strategies that can capture individual differences in developmental
trajectories\textsuperscript{91}. Although this avenue of research has not yet been fully explored, interactive tasks that
encompass real-time social interaction could allow the study of brain activity in experimental contexts
that are more relevant for core ASD symptoms, rather than the more passive tasks that are used in most
functional imaging studies\textsuperscript{103}. Experiments focusing on human-human interaction\textsuperscript{104} and human-machine
interaction\textsuperscript{105} have been undertaken but, so far, no study has ever made explicit use of such methods to
study the electrophysiology of ASD.

In a further search for mechanisms of ASD, prospective baby siblings studies have suggested that the
gradual emergence of behavioural symptoms of ASD is preceded by earlier subtle alterations in the activity
of regions and networks of the social brain\textsuperscript{24}. For example, early work on a small group of 5–6-month-old
infants who later developed ASD observed faster but less prolonged neural activation and delayed
sensitization responses to faces compared with infants who did not develop ASD\textsuperscript{106}, and one report
demonstrated that newborns with an increased familial likelihood of ASD showed higher signal
homogeneity within core social brain networks (right fusiform and left parietal cortex\textsuperscript{107}). By comparison,
reduced frontal power, particularly in the high-alpha band, during quiet play at 3 months of age\textsuperscript{108} and
cortical hyperexcitability in the right tempo-parietal region during auditory repetition of pure tones at 9–
10 months of age have been found in babies at familial risk for ASD\textsuperscript{109}, suggesting that atypical patterns
occur in brain regions other than those involved in social processing. Such alterations could have a
cascading effect on social learning and contribute to the later emergence of behavioural symptoms of
ASD, although a causal link remains to be demonstrated. Replications across different research centres
are needed because many of these studies had small sample sizes, different definitions of groups and
varied measures and time points.

Interestingly, results from MEG and EEG studies jointly point toward two physiological mechanisms of
ASD: excitation/inhibition (E/I) imbalance and alteration of large-scale functional interactions of brain
systems as quantified through connectivity analysis\textsuperscript{110}. An E/I imbalance is supported by results from
computational modelling of how reductions in the amount of inhibition can account for the previously
observed perceptual consequences of ASD\textsuperscript{111} and transcranial magnetic stimulation (TMS) studies
demonstrating a neurophysiological deficit in \(\gamma\)-aminobutyric acid (GABA) receptor-mediated function in
people with ASD\textsuperscript{112}. In parallel, decreased long-range functional connectivity has also crystalized as a
consistent mechanism\textsuperscript{113}. MEG studies have especially suggested a complex functional connectivity
pattern in the somatosensory cortex with reductions in the feedback (top-down) direction, but increased
in the feed-forward (bottom-up) direction\textsuperscript{114}. Clarifying the extent to which this pattern is a
methodological artifact that could result from the predominant average-brain approach, as suggested by
fMRI studies, is critical\textsuperscript{115}.

Beyond use to understand the pathophysiology of ASD, the scalability and accessibility of EEG suggest that
this technique could be an ideal candidate for use as a brain-based biomarker. Measures from information
theory have already provided promising case-control classification\textsuperscript{116}, but developing generalizable biomarkers may require a combination of multiple EEG measures supported by robust machine learning methods\textsuperscript{117}. Against the background of the current reproducibility crisis that characterizes many studies\textsuperscript{118}, as well as the defining heterogeneity of ASD, the next breakthrough will certainly demand large-scale collaboration between researchers and clinicians.

[H1] Diagnosis, screening and prevention

Diagnosis of ASD is made on the basis of behavioural presentation. Although substantial heterogeneity exists between and within individuals across development, a set of core diagnostic features of ASD (covering social interaction, communication and flexible or sensory behaviour) can be reliably identified by trained clinicians\textsuperscript{119,120}.

[H2] Diagnostic criteria

The re-formulation of the diagnostic criteria for ASD in the DSM-5 (Table 1)\textsuperscript{121}, which is similar to the criteria in ICD-11\textsuperscript{122}, contains several changes from previous editions that were based on good empirical and clinical evidence\textsuperscript{123}. First, the sub-classification of ‘Asperger’s disorder’ was subsumed under the unitary term ASD as the diagnosis was inconsistently applied even by expert groups\textsuperscript{124}. This change is controversial, but the evidence supporting the inclusion of Asperger’s disorder as a separate condition is very weak\textsuperscript{125}. The important questions are how better to consider the factors that characterize differences among autistic individuals and ensuring that these differences are measured and addressed using neurobiological and clinical research, rather than contained within very poorly defined categories of Asperger’s and PDD Not Otherwise Specified (NOS) as defined in DSM-IV. In addition, some individuals with social communication problems but not restricted and repetitive behaviours who would previously have fallen into the now-removed subcategory of PDD-NOS now receive a different diagnosis of Social communication disorder, which is not yet well-validated. Although these changes have led to concerns that the DSM-5 ASD criteria are more restrictive than those in DSM-IV, many clinicians feel that the changes better reflect clinical consensus and practice. Second, the social and communication domains of the diagnostic criteria were unified to reflect the factor structure of symptomatology. Third, sensory anomalies (hypersensory and hyposensory responsiveness and sensation-seeking) in DSM-5 were included under the ‘restricted, repetitive behaviours and interests’ domain to reflect their pervasiveness\textsuperscript{126}. Fourth, the DSM-IV criteria required symptoms to be present in the first 3 years of life, but criteria in DSM-5 recognise symptom onset occurring in the early developmental period with the caveat that symptoms might not fully manifest until social demands exceed limited capacities. This change recognizes the developmental nature of ASD, wherein for some individuals, clear manifestation of ASD might not be apparent until mid-childhood, adolescence or even adulthood. In addition, late diagnosis (that is, diagnosis beyond early childhood) can occur even in those who received intensive early monitoring\textsuperscript{127}. In addition, the DSM-5 criteria supports the use of specifiers that can denote those with a dual diagnoses, such as individuals with ASD and ADHD or other psychiatric disorders, as well genetic conditions such as fragile X syndrome or down syndrome. Beyond the clinic, these changes have
implications for large-scale data pooling efforts; for considering domains of behaviour to be modelled; and for identifying shared and distinct developmental pathways to conditions like ASD and ADHD.

[H2] Diagnosis and screening in children

The two core elements of the diagnostic process of ASD in children are a detailed developmental history that is usually obtained from parents, covering first concerns and early history to the present day, and an observation of the child’s interactions with their parents and with unfamiliar adults during a combination of structured and unstructured assessments. Ideally, observations of the young person in peer-group settings such as school or nursery would also form part of the diagnostic process. Of note, in one population-based study in the UK, girls with similar levels of symptom expression to boys were less likely to receive a diagnosis of ASD from clinical services. This finding might reflect socio-cultural factors in the application of the diagnostic criteria, greater resilience or protective factors in girls that reduce the need for clinical services at a given symptom level, or the need for the revision of instruments used to identify symptoms to more fully cover female autistic traits.

A number of structured diagnostic interviews and observational assessments for ASD exist, but only a limited number have been rigorously tested for diagnostic accuracy relative to the gold-standard of expert clinician judgement. Although these interviews and assessments have reasonably robust sensitivity, specificity and reliability (see for a review) and are widely used in some services in communities, there are also challenges to the widespread adoption of the best validated instruments: the Autism Diagnostic Interview—Revised (ADI-R) and the Autism Diagnostic Observation Schedule—2nd Edition (ADOS-2). These challenges include the cost of the instruments and training, the time required to complete them and the need for substantial training to use them reliably. Although expert clinical judgement was previously believed to be more reliable than reliance on instrument scores alone for the diagnosis of ASD, more recent evidence suggests this may not be true at least in toddlers and preschool children. The need to take a global perspective on ASD is driving attempts to develop more scalable tools, but this work is currently in its infancy.

The stability of a diagnosis of ASD from the preschool years to mid-childhood is relatively high. However, although diagnostic systems currently presuppose that ASD is a lifelong condition, there is a growing recognition that ASD has a heterogeneous developmental time course. Indeed, sub-groups of individuals with ASD and improving or worsening symptoms over time can be identified. Such developmental trajectories might be a more meaningful phenotype on which to map aetiological mechanisms than a static case-control dichotomy. Some individuals diagnosed as children have no clinically meaningful (or even detectable) impairment later in life (so-called ‘optimal outcome’); one critical question in identifying mechanisms is whether this profile is associated with successful effects of early intervention or is an aetio logically distinct subtype of ASD.

[ Au: please see the comment in the Epidemiology section (line 121) regarding the use of the term ‘screening’ - do we need to introduce some edits here? I’d be grateful if you could take a look through this and amend as needed. ] [H3] Screening and early identification. The potential for early testing to prospectively identify children with ASD at a young age has considerable interest, and several studies have evaluated the performance of parent-report instruments between 14 and 24 months of age, such as the Modified Checklist for Autism in Toddlers (M-CHAT) and the Early Screening of Autistic Traits (ESAT).
Examples added based on Tony’s comment YES fine. However, there are contrasting views on the strength of the evidence for universal population-wide testing. Of note, research is lacking on the effectiveness of therapeutic interventions in those identified with ASD through universal screening. In addition, although it is possible to identify some children with ASD before parents or professionals have identified concerns, diagnosis is missed in many children, and most tested cohorts have not been systematically followed up to identify later-onset ASD in children who initially tested negatively. Screening also often identifies children with broader developmental difficulties as well as those with ASD. In general, such instruments could be more useful for identifying possible signs and symptoms of ASD in high-risk populations, for example in young children with older siblings with ASD, or in those referred for speech or other developmental concerns to community paediatric services. In addition, population-wide testing may also play a part in improving awareness and recognition of the early signs and symptoms of ASD in both professionals and the general public, which alongside ongoing developmental surveillance pathways in community services, could help to bring down the age of recognition and diagnosis. These principles also apply in low-income and middle-income countries in which testing for ASD and other neurodevelopmental disabilities has only just begun to be developed. Very little research has been devoted to cultural and ethnic differences in either child early presentation and parents’ understanding or the experience of autism, which may in fact affect how screening instruments work and thus impact on parents and families as much as autistic individuals.

[H3] Early developmental profiles. Understanding of onset patterns of ASD has dramatically expanded over the past 10 years, through work on infants with a first degree relative with ASD, who due to the high heritability of the condition have a 20% chance of developing ASD themselves. Symptoms of ASD have a gradual developmental onset. Indeed, although the average age of ASD diagnosis remains 4–5 years of age, parents typically report first concerns to health professionals at ~2 years of age. In many individuals, symptoms emerge during the second and third year of life (although, as per the DSM-5 onset criteria above, in others, onset might not be noticed until the child reaches school-age or later) whereas in others, symptoms become apparent after a seeming period of typical development, including a period of regression or stasis. To this end, conceptualization of what has been called ‘regression’ prior to 2 years of age has been reconsidered. Over the first two years of life, a substantial proportion of infants who later receive ASD diagnoses show gradually accumulating delays across social, communication and language domains, suggesting that ‘regression’ represents a spectrum ranging from frank loss of acquired skills, to a gradual erosion (or ‘plateauing’) of developmental potential to individuals in whom these skills never emerge.

[H2] Diagnosis and screening in adults

Information on diagnostic methods to identify ASD in adulthood is in its infancy, with little methodologically acceptable evaluation of interview methods or screening questionnaires (including self-completion questionnaires). Clinical approaches rely heavily on extending methods developed for use in childhood to adulthood. These methods tend to rely on childhood developmental data, although validation research in adult general population-wide testing suggests good specificity and sensitivity for the observationally based ADOS Module 4. However, typically, much research has depended on the
judgment of expert clinicians and of standardized data collection on early child development that is unlikely to be obtainable for many older adults. Given that (undiagnosed) autistic adults presenting for an ASD assessment are also more likely to have co-occurring adult mental health disorders, any method of assessment must be capable of differentiating such abnormalities in symptoms and behaviour from abnormalities due to ASD. This point has led to the suggestion that clinical examination methods to identify adult psychopathology could be extended to include ASD in addition to depression, anxiety and psychosis, among other disorders. Semi-structured adult psychopathology interviewing has been fruitful in the assessment of closely related neurodevelopmental disorders in adults, most notably ADHD. Given that most people in the world who are autistic are adults, and as many of these individuals have not received a diagnosis of autism, the development and evaluation of such adult assessment approaches is an urgent research priority.

[H2] Co-occurring disorders

In addition to the core features of ASD, co-occurring difficulties or disorders are much more widely recognized in research, although they are not necessarily adequately addressed in clinical practice. For preschool children with ASD, language delays, motor problems, epilepsy, difficulties with sleep and eating, and high levels of activity are most commonly observed. By comparison, ADHD, anxiety, obsessive-compulsive disorder (OCD), intellectual disability, academic challenges, irritability and disruptive behaviours become more apparent in school-aged children. The proportion of individuals with depressive symptoms becomes higher in adolescents and adults, whereas other issues often remain. Moreover, growing evidence (although it is reliant on administrative case-finding data) suggests that people with ASD have premature mortality and increased risk of self-harm and possibly suicide, although the mechanisms involved have yet to be elucidated. Studies using electronic health records have demonstrated that adults with ASD are more likely to be diagnosed with many physical health conditions such as immune conditions, sleep disorders and obesity, compared with adults in the general population.

Collectively, these difficulties and disorders contribute to ASD severity and independence and well-being at each age. However, it is important to note, in the context of heterogeneity, that the prevalence of each of these co-occurring conditions varies considerably with the context of the sample (such as from psychiatry referrals, neurological referrals, or schools) and the methodology used (administrative, self-report or assessed), as well as with age, level of cognitive function and perhaps region. As many of these conditions are treatable, they are very important as clinical considerations but are also more complex than sometimes conveyed.

[H1] Management

[H2] Early intervention
Early intervention is seen as a priority because many young children with ASD struggle to communicate and interact with others, restricting their opportunities to learn and affecting their parents who can find their child’s behaviour perplexing and challenging to manage. Thus, outcomes of such interventions include changes in the individual’s availability for learning and increased parent understanding. Intervention delivered in the preschool years at an age when there is increased brain plasticity might lead to additional benefit, although this theory has not yet been empirically supported.

The primary models of psychological intervention for preschool children with ASD are developmental and behavioural. Although some consensus has been reached on the interventions that have more supporting evidence (termed ‘naturalistic developmental behavioural interventions’), there is some uncertainty and disagreement about the strength of evidence for different approaches, with almost no direct comparisons of treatments or studies to assess which child should receive what treatment or treatment intensity. Indeed, clinical trials in ASD are limited by cost, time, placebo effects and limited outcome measures, and are far behind much of the other research. This gap leaves parents and practitioners at the mercy of what is available and sometimes marketed in their region. Indeed, access to early intervention services is variable in most communities, including in high-income countries, and is mostly carried out by non-specialists supervised by specially trained professionals. In low-income and middle-income countries, most children and young people with ASD — similar to those with intellectual and developmental disabilities — will not receive specialized services, although a number of groups have begun to test community delivery of early intervention in such settings.

Many current interventions build on the original ‘Applied Behaviour Therapy’ (ABA) and have shifted to more natural, child-initiated developmentally appropriate strategies and tasks instead of dependence on repeated ‘discrete trials’ (known as discrete trial training, or DTT). In addition, considerable variation exists between different intervention models in terms of mode of delivery (for example, parent-mediated versus therapist-implemented), length (12-week versus 2-year programs), intensity (from a few hours a week to ~15 hours per week) and the balance between the developmental or dyadic versus behavioural components.

Lower-intensity approaches include parent-mediated interventions whereby parents are coached to become more attuned to their child’s communication signals and style (which are considered an intermediate child outcome) and to facilitate more joint engagement in play and everyday activities, designed to increase social and communication skills in the child. Some studies have demonstrated enhanced joint engagement and joint attention (which are considered important intermediate child outcomes), with these lower-intensity approaches in preschool children compared to a control group, such as the 12-week Joint Attention Symbolic Play Engagement and Regulation (JASPER) program, both when delivered by parents in the home and by teaching assistants in school. However, other lower-intensity, time-limited parent-mediated interventions such as Focus Playtime Intervention (FPI) have not improved child outcomes (such as social orienting and joint attention), although some interventions have increased parental responsiveness. A longer program (Preschool Autism Communication Trial (PACT)), which consists of fortnightly parent-therapist sessions for 6 months, then monthly sessions for another 6 months, demonstrated improvements in parent and child dyadic behaviours such as parental synchrony and child initiations when interacting with each other (those close to the intervention target) but not symptom reduction at immediate follow-up. A subsequent 6-year follow-up to mid-childhood at age 7 to 11 years identified modest reductions in overall ASD symptoms using the ADOS over the whole course of the study that were not detectable at the immediate endpoint, suggesting that a longer-term perspective is critical in considering outcomes.
A higher intensity, more comprehensive approach is the Early Start Denver Model (ESDM), which combines behavioural and developmental or dyadic approaches. The ESDM is delivered by therapists for ~15 hours per week, and as part of this programme, parents are trained to improve social communication and interaction with their child. A small-scale trial demonstrated improvements in child developmental and adaptive outcomes, primarily in the language and communication domains, following 2 years of ESDM compared with treatment as usual\textsuperscript{182}. One larger multi-site trial found attenuated benefits with improvement in language outcomes at two of the three trial sites, but no differences between the treatment as usual and ESDM groups in overall developmental ability, adaptive behaviour or ASD severity\textsuperscript{179,183}.

Many of these early intervention approaches are based on models of typical development. Increasingly, studies are using a combination of methods to define treatment outcomes and to better understand the mechanisms and models of change of interventions. These methods include analysis of the degree to which changes in the direct target of the intervention (for example, parent behaviour) mediate later changes in child behaviour\textsuperscript{182}, and the use of experimental methods such as EEG to examine whether there are accompanying changes in relevant brain networks\textsuperscript{184}. Many parents seek complementary medical approaches, which to date have not been supported and sometimes are dangerous\textsuperscript{185}. A note of general caution is that even in the context of significant treatment differences between groups, individual outcomes are very variable, and some children do not improve, although reliable predictors of response to treatment have not been demonstrated in rigorous, randomized controlled trials. As ASD is a heterogeneous developmental condition, different interventions may be required at different stages throughout life and different individuals might benefit from different interventions [Au: edits for brevity ok? Yes]. One area which many consider to hold much promise, that of neurobiologically or biomarker ‘informed’ psychological intervention, is on the horizon but such targeted therapies have not yet been developed.

**[H2] School age children and adolescents**

Many children and young people with ASD can also benefit from interventions at later ages. A number of programs and approaches are available that focus on the core social communication difficulties of ASD; for example, social skills training programs for which moderate evidence of benefit exists\textsuperscript{186,187}. In addition, non-verbal young people with ASD can benefit from use of augmentative communication systems, such as the Picture Exchange Communication System (PECS) that use picture symbols and behavioural training methods to allow children to request and make choices\textsuperscript{188} or other technology-based augmentative communication systems. Increasingly, more generic interventions that target co-occurring emotional and behavioural problems are being adapted for youths with ASD, and initial studies suggest moderate benefits\textsuperscript{189}. These interventions include modified cognitive behavioural therapy (CBT) for anxiety (modified, for example, to include parents, increase the duration of sessions, use more visual materials and specific work on understanding one’s own emotion states)\textsuperscript{190} and parent-mediated interventions for disruptive behaviour and ADHD\textsuperscript{191}. More recently, there have been efforts to develop and test interventions that target aspects of parental wellbeing, such as parental stress and self-efficacy\textsuperscript{192}. Increasingly, interventions for school-age children and young people with ASD are being delivered within the school environment, rather than the clinic, which has natural advantages for programmes that consist of groups or peer-to-peer interactions and an emphasis on social skills. Indeed, it is hoped that this approach may facilitate generalization of the skills learned\textsuperscript{193,194}.

**[H2] Adult services**
As individuals with ASD progress into and through adulthood, the focus of management shifts from treating the core symptoms of ASD to addressing associated symptoms or behaviours and promoting independence. However, there are few intervention studies to guide treatment options in adulthood. Indeed, a 2012 systematic review identified only 32 studies published between 1980 and 2010 that evaluated treatment studies for adolescents and young adults with ASD. A more recent review identified 41 studies of interventions targeting social functioning in adults over a 37-year period.

Despite the low number of treatment studies, there is some evidence supporting treatment efficacy for a limited number of symptoms, behaviors, and functional outcomes such as employment, social skills, and anxiety; however, in general, the evidence-base is weak. For example, only three randomized controlled trials (all of which included small cohort sizes) that tested job interviewing skills curricula have been published. Social skills interventions have a somewhat more robust literature base, but most of these studies had very small sample sizes and were not well controlled. In addition, it is unclear whether social skills interventions can be generalized to other social settings and situations, that is, whether skills learned in the treatment context are used by the participants in other settings, such as with peers or at work. There is some evidence for the use of cognitive-behavioural therapy (CBT) for effectively treating anxiety in people with ASD who do not have cognitive delays or language problems.

Formal service systems and social care can help fill in the treatment gaps. Indeed, although many adults with ASD do not receive adequate services and support, their receipt can improve outcomes across a number of domains. For example, transportation services can allow adults with ASD to engage in employment and access therapies and programs in the community. In addition, comprehensive job support services can promote finding and maintaining employment, particularly for adults with more severe impairments. Public health insurance can increase access to psychiatric care for those with co-occurring mental health problems, and income supports can reduce dependence on families.

[H2] Medications

All medications that have evidence of benefit for ASD treat the associated symptoms or co-occurring diagnoses, rather than the symptoms of ASD directly (including social communication or repetitive behaviours). As mentioned earlier, ASD is an extremely heterogeneous disorder, and individuals with ASD can have a number of common co-occurring disorders that can also vary in severity.

Risperidone and aripiprazole (both of which are atypical antipsychotics) are approved in the USA to treat irritability and agitation — including aggression, self-injury and tantrums — in children and adolescents with ASD. However, both treatments are associated with adverse events, including sedation, risk of movement disorders and weight gain, which limit their use to people with severe irritability with agitation. The anti-diabetes drug metformin has been shown to limit weight gain from these medications, possibly broadening their safe use.
As mentioned previously (see Co-occurring disorders, above), co-occurring mental health conditions are common in people with ASD. Methylphenidate, atomoxetine and guanfacine are beneficial for ADHD symptoms in ASD ([Au: given that these medications target different symptoms of ADHD are they ever prescribed in combination? Or will a child be given one medication only?]). Although serotonin reuptake inhibitors (SRIs), such as fluoxetine and citalopram, are used for the treatment of depression, anxiety and OCD in the general population, they have differing efficacy in people with ASD ([Au:OK?]). Indeed, although fluoxetine improves symptoms of OCD in adults with ASD, citalopram has demonstrated poor tolerability and no benefit for repetitive behaviour in children with ASD. Medications for depression or anxiety have not systematically been tested in people with ASD.

Some excitement has accompanied the recent studies of medications targeting the neurohormonal oxytocin or vasopressin systems, both of which modulates social behaviour across species. Underpowered studies of intranasal oxytocin have demonstrated mixed results that are overall not supportive of a large effect size, with results pending from adequately powered studies ([Au: please provide the NCT number of this ongoing study here so readers can follow up in due course]). In addition, a pilot study of intranasal vasopressin suggested possible benefit in people with ASD, although this study was underpowered. A large trial of balovaptan, a vasopressin AVPR1A antagonist in adults with ASD showed negative results on its primary outcome (a general rating of ASD symptoms), with suggestive results on a key secondary parent report measure of adaptive behaviour, including social and communication behaviour. A few studies have also focused on the hypothesis that, at the level of neural circuits, ASD may result from excessive excitation or insufficient inhibition, with some promising but inconclusive results for medicines that target the GABAergic system. Medications targeting genetic syndromes that can cause ASD have not yet yielded consistent improvement, but there is much hope for a precision medicine approach that links genetic subgroups with neurobiology-based treatments.

[H1] Quality of life

[H2] Objective and subjective measures

Several aspects of intervention research speak straight to the heart of current debates within the clinical field and broader ASD community, including how a good outcome is classified for an individual with ASD, as well as who should decide what outcomes are used ([Au: edits for style ok?]) in intervention studies. This point is aligned both with the debates about medical versus social models of disability but also with a more general shift in medicine away from focusing on symptom reduction to improving the wellbeing and quality of life (QOL) of patients. QOL research in adults with ASD has focused on two aspects: objective and subjective QOL. Objective QOL encompasses social achievements such as employment, adequate living conditions, supportive relationships, and good physical and mental health, whereas subjective QOL focuses on individuals' perceptions and subjective assessments of their own lives. Both subjective and objective QOL are often related, but not synonymous, and both are important to take into account when considering outcomes for individuals with ASD (Table 3).
**[H3] Objective QOL.** Adults with ASD tend to have poor objective QOL. Unemployment is high in this population, and even among those employed, individuals are often working below their skills and abilities.\textsuperscript{20,221} Moreover, independent living can be \textsuperscript{[Au:OK? As presumably some individuals may be able to live independently with relative ease?] a challenge, and adults often lack meaningful relationships with peers.\textsuperscript{222} When aggregating across these domains of life, many adults with ASD have ‘poor’ or ‘very poor’ outcomes.\textsuperscript{223,224}

ASD is a highly heterogeneous condition and several factors have been associated with higher versus lower objective QOL. Most of the studied factors associated with higher objective QOL have been characteristics of the individuals (versus families, service system or communities), and consistent predictors of higher objective QOL include better early language development,\textsuperscript{[Au:OK?] higher IQ and adaptive behaviour scores, less severe ASD symptoms, and fewer challenging behaviours.\textsuperscript{225} In addition, more recent research suggests that women with ASD may have a more difficult time maintaining employment positions\textsuperscript{226} and are more likely to ‘camouflage’ their ASD symptoms than men, which can lead to mental health challenges.\textsuperscript{227}

**[H3] Subjective QOL.** Meta-analysis have suggested that across the lifespan, subjective QOL tends to be lower among individuals with ASD compared to typically-developing peers,\textsuperscript{228} but is often more positive than indicators of objective QOL.\textsuperscript{223,229} Predictors of subjective QOL tend to be inconsistent across studies, except for perceived stress and supports, the latter of which encompasses services, family and social support.\textsuperscript{230–232}

**[H2] Self-advocate perspective**

[Peer reviewer comment: The self-advocate perspective is important. It was quite noticeable to use identity-first language (“autistic people”) only in this section and some explicit reference to this choice might be made, to help readers understand the politics and differing viewpoints.] It is clear that ASD has heterogeneous outcomes and biological underpinnings; what is less clear-cut are the differing and nuanced views of autistic people regarding how ASD should be approached and researched. (Box 2, Autistica,\textsuperscript{233} see also Ontario Brain Institute.\textsuperscript{234}) Indeed, some people with a diagnosis see ASD as being a fundamental part of their identity whereas other people do not. In addition, many people feel that social change is required, whereas other individuals want therapies to meet a range of their needs. The key is respect for a variety of views and ultimately respect for autistic people. Researchers can demonstrate respect by considering how ASD as a topic is distinct from, for example, cancer. To this end, terms like ‘disease’ are inappropriate and are scientifically inaccurate when referring to ASD. Ultimately, active participation in the design, implementation and interpretation of research studies,\textsuperscript{[Au:OK?] clear consideration of research ethics and the consequences of research involvement, and broad consultation of autistic people in research is key to authentically addressing the substantial inequalities autistic people face as a group and ensuring they live long, healthy, happy lives.

**[H2] Family perspectives**

Families of people with ASD are also heterogeneous, yet, as a group, they experience lower QOL than families with a member with other neurodevelopmental conditions, even before receiving the formal diagnosis.\textsuperscript{237} For this reason, it is essential that parents, other family members, clinicians, educators, and
the entire external support system coalesce around common goals for outcomes whilst accessing and maximizing resources for the betterment of the child and family. Parents are typically at the centre of this support network and carry much of the responsibility of direct care, coordination and advocacy, over and above typical parental responsibilities. The exact parental roles are dependent on the child’s strengths and challenges, and frequently shift over time (Fig. 6). During this process, it is important that parents maintain motivation by setting realistic goals and tracking progress to experience the many achievements that their loved one with ASD can attain.

Effective parents often work closely with experienced providers who can track development of the child with ASD and can provide guidance on next actions. Early in childhood, this role includes identifying and engaging with early and school-based interventions. It is never too early for parents to begin planning for the adult transition process, including (dependent on the person with ASD’s capacity) promoting self-advocacy, preparation for life after secondary education, vocational training and employment supports, living needs, community participation, and long-term financial considerations. During adulthood, for cognitively-able adults, parental roles might shift to more traditional relationships, whereas for those with cognitive disability, parental caregiving often continues and culminates in planning for late life needs. Although the journey can be challenging, for many parents, it can be incredibly rewarding and a source of life meaning.

Many parents recognize the need to give back to the community through research. Accordingly, it is crucial that researchers foster this desire carefully, communicating with parents to ensure that any potential immediate or future risks or benefits are clear. Even if the study period is brief, in many cases, the goal should be to develop a positive longer-term relationship, as this can lead to parents and people with ASD continually re-engaging in and developing positive feelings about the research process.

[H1] Outlook

ASD research has substantially expanded in the past 50 years, particularly the past 20 years, as reflected in the websites listed in Box 3. Although it seems unlikely that the incidence of ASD is truly rising at the rate suggested in prevalence studies, these data have increased awareness and the numbers of diagnosed children in schools and clinics, although adult services and recognition run far behind. The lives of people with ASD diagnoses have improved at least in some high-income countries, with a greater proportion of children using some language, more adults with educational qualifications and less institutionalization, although the changing nature of diagnoses has to be considered when interpreting historical trends. Some risk factors for ASD have been identified (such as increased parental age, birth trauma and a positive family history) which has implications at least for more careful follow-up. In addition, the genetics of ASD has yielded surprising discoveries with substantial implications for heritable neurodevelopmental disorders, such as ADHD, language delay and named syndromes associated with profound intellectual disability. The perceived value of routine genetic screening for ASD diagnosis is disputed, with American medical academies strongly in favor whereas those in other countries much more selective. Studies of brain structure and function have added similarly intriguing findings that are just beginning to be integrated into both developmental and more mechanistic models of behaviour with possible targets or markers for change. Despite the intellectual contribution of these studies to research, at this point, neither EEG nor imaging are recommended as part of standard practice for diagnosis of ASD but can be used for other neurological indicators (such as if there are concerns beyond ASD symptoms that merit an EEG or imaging). In this field, replication of findings across
sites and even within individuals, as well as larger samples through collaboration are the promise of the future.

One way of bringing the three themes of mechanisms, heterogeneity and outcomes of ASD together is to consider the trajectories of this disorder over time (Fig. 7), and how knowledge of these trajectories can contribute to investigations of the biological and cognitive underpinnings of ASD, and how treatments and supports could make the lives of children and adults with ASD more positive.

In terms of mechanisms, despite earlier hopes for simple genetic explanations of ASD, instead, we have identified many single gene germline loss of function point mutations yielding some initial models of disruption in very basic molecular patterns, as well common genes with small effects that are just beginning to emerge. Attempts to study genes-first have shown heterogeneity even within highly specific CNVs, with a few exceptions. In addition, hope exists that genetically based interventions for ASD may be possible, although this will likely involve much further research.

One way of bringing the three themes of mechanisms, heterogeneity and outcomes of ASD together is to consider the trajectories of this disorder over time (Fig. 7), and how knowledge of these trajectories can contribute to investigations of the biological and cognitive underpinnings of ASD, and how treatments and supports could make the lives of children and adults with ASD more positive.

Data from genetic approaches that might yield targeted genetic interventions may be most relevant to rare, severe neurodevelopmental difficulties in general rather than ASD as a specific entity. With more information about the differing developmental trajectories of ASD, more continuous measures of language and intellectual function, behavioural phenotypes and changes over time can be quantified across different neurobiologically defined subgroups. This approach could potentially identify different ‘routes’ to different outcomes, whether ASD or not, and could have a practical benefit in terms of selecting and monitoring appropriate treatments. In addition, with the heterogeneity of ASD, our growing understanding of mechanisms, be they causal or mechanisms for change, needs to be linked to trajectories in development and not be considered as static. Researchers modelling ASD in other species might find the incorporation of early developmental manifestations, such as regressions or motor delays, more tractable than the current focus on ASD-related social communication symptoms seen in humans. With collaborations and studies of sufficient sample sizes, investigators have begun to focus on findings within different developmental periods that could provide insight into trajectories and targets for intervention. Thus, more study of the development of ASD both in studies of human behaviours and in animal models might have an effect on the identification and treatment of ASD as a neurodevelopmental disorder. Prospective studies, including epidemiological and direct behavioural work across developmental periods, moving beyond very young children to later childhood, adolescence, and adulthood are needed.
Similarly, limited findings about adult development and patterns that lead into ASD (Figs 5 and 7), call for measurement of different outcomes that respect individual differences in autistic people and in families (Box 2). By young adulthood, available supports for places to live, employment and mental health services are needed for individuals who have a range of skill levels, with supports not always well matched to the needs of individuals; however, comparisons of treatments or treatment intensities have not historically been made, even though they are continually called for. The types and specific goals of treatments differ greatly for autistic people who are verbally fluent versus those who have difficulty speaking for themselves, such that alternative systems need to be in place that take into account co-occurring conditions, strengths, preferences and challenges. More studies of well-defined, more homogeneous subgroups of autistic children and adults over time would provide different and more useful information about real-life issues, as in Table 3 than large-scale surveys of very heterogeneous samples.

Progress in the biology of more generally defined neurodevelopmental disorders may have the greatest yield for children with ASD in their early years [Au: compared with what? In their later years?]. Clinical trials that compare known treatments (both psychosocial and biological), with new ones and treatment as usual would allow us to build on previous findings in a more meaningful way and begin to address the priorities listed in Box 2, which strikingly, are seldom priorities in autism research. To move from science to practice including evaluation and treatment, ASD researchers need to find a way to select and fund studies of more mundane, but critical evidence gaps in understanding heterogeneity, mechanisms of change and outcome that affect practice in any circumstance, not just internationally, within academic systems that reward creativity and novelty. Unique methodologies, including the baby sibling studies, accumulation of large data sets (such as ABIDE, and the Simons Simplex Collection (SSC)), prospective epidemiological studies and mechanistic studies of intermediate biomarkers may begin to bring together information from molecular to pathophysiological to cognitive and behavioural levels. However, for now, as for other neurodevelopmental and psychiatric disorders including schizophrenia, the distance between science and practice remains great, and the amount of research that attempts to address solvable problems for autistic people alive today and their families remains modest.
Box 1. Global challenges in autism research

Recently, there have been calls for more attention to global issues in autism research\(^\text{249}\) (Global Research on Developmental Disabilities Collaboration – Lancet Global Health, 2016), including a number of related issues with somewhat different potential solutions. For example [Au: edits to improve the narrative flow ok?], broader populations should be included in autism research, including individuals from Lower Resource and Middle Income countries (LMICs), but also inclusive representation of the ethnic, linguistic and socio-economic diversity of many High Resource countries and people whose autism is unrecognised. Moreover, there should be the creation of opportunities to carry out research in LMICs\(^\text{251}\) (Patel et al, 2015). Open source and shared databanks, including autism-specific resources such as the Simons Simplex Collection and Autism Brain Imaging Data Exchange [Au:OK?] (ABIDE), as well as broader collaborations such as PsychENCODE [Au:OK?] could assist in promoting international research [Au: edits for brevity ok?]. In addition, the science of autism should be disseminated in ways that are useful for practice in all countries, [Au: edits ok?] but with particular attention to the needs of communities and families with fewer resources\(^\text{252}\)\(^\text{253}\). [Au: sentence deleted as suggested] More immediately, searches for scalable methods of identification and perhaps intervention with children and adults [Au:OK? Yes] with autism\(^\text{135,254}\) have begun [Au: globally or in LMICs?]. However, the need to develop scalable global practices highlights how little is known about when we need population-wide testing for autism versus broader neurodevelopmental disorders, the minimal intensity and duration of effective interventions, behavioural mechanisms behind changes in behaviour and which treatments work with which children and adults and families, all of which have a bearing on interventions locally and globally. In addition, global issues of stigma, governance and paucity of resources also have to be taken into account\(^\text{251}\). 

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Box 2. Top ten questions for ASD research proposed by autistic people, family members and professionals. (Reference: Autistica\textsuperscript{233})

1. Which interventions improve mental health or reduce mental health problems in people with autism spectrum disorder (ASD)? How should mental health interventions be adapted for the needs of people with ASD?

2. Which interventions are effective in the development of communication and language skills in ASD?

3. What are the most effective ways to support or provide social care for autistic adults?

4. Which interventions reduce anxiety in autistic people?

5. Which environment supports are most appropriate in terms of achieving the best education, life and social skills outcomes in autistic people?

6. How can parents and family members be supported and/or educated to care for and better understand an autistic relative?

7. How can ASD diagnostic criteria be made more relevant for the adult population? And how do we ensure that autistic adults are appropriately diagnosed?

8. How can we encourage employers to apply person centred interventions and support to help autistic people maximize their potential and performance in the workplace?

9. How can sensory processing in ASD be better understood?

10. How should service delivery for autistic people be improved and adapted in order to meet their needs?
Box 3. Examples of ASD websites [Au: based on Tony’s comment I’ve introduced some broader headings here rather than adding a description under each individual website - what do you think? Please check carefully and let me know if you prefer the original version (we can revert the changes, no problem!)]

Sites for health care professionals or research scientists:
[H1] American Academy of Pediatrics

[H1] International Society for Autism Research
ASD:
https://www.autism-insar.org

[H1] National Autistic Society
https://www.autism.org.uk

[H1] Royal College of General Practitioners

Information about treatment, research and advocacy for people with ASD and their families:
[H1] Autism Canada
https://autismcanada.org

[H1] Research Autism
http://www.researchautism.net/

[H1] Autism Europe
https://www.autismeurope.org/

[H1] Autism India
http://www.autism-india.org

[H1] WHO
https://www.who.int/news-room/fact-sheets/detail/autism-spectrum-disorders

[H1] Autism Spain
http://www.autismo.org.es

[H1] Autism Speaks
www.autismspeaks.org

[H1] Autismo Deutschland
https://www.autismus.de

[H1] Autistica
https://www.autistica.org.uk
Information about research funding, and up-to-date information for people with ASD and families:

[H1] Simons Foundation
https://www.sfari.org

[H1] US NIH

[H1] Autism Science Foundation
https://autismsciencefoundation.org

Epidemiological information [Au:OK?]

[H1] US CDC
https://www.cdc.gov/ncbddd/autism/index.html

Figure 1. Theories and findings regarding ASD mechanisms, outcomes and heterogeneity.

Original descriptions of the cardinal features of autism spectrum disorder (ASD) were attributed to a range of causes including being raised by wolves (the Wild Boy of Aveyron), inborn limitations in affective contact and unfeeling parenting (such as 'refrigerator mothers') and holy people (such as fools for Christ))

Conceptualizations of ASD as a common highly heritable neurodevelopmental disorder with underlying cognitive features began with the recognition of differences in brain function and cognition in the 1960s and the first twin study in the 1970s. Other proposed mechanisms include maturational lags in neurophysiology and cognitive mechanisms such as joint engagement. With the search for pathways to and sometimes out of ASD on many levels, conceptualization of positive outcomes has been more recent, but has also varied markedly. In the 1970s, autism societies and collaborative clinical programs focused on community integration and de-institutionalization (such as National Autistic Society (NAS) and National Society for Autistic Children (NSAC)) Priorities shifted in the 1980s and 1990s, with still unreplicated claims of ‘recovery’ in children who participated in intensive behavioural interventions, new advocacy groups focusing on biomedical discoveries to yield potential biological treatments and even ‘cures’ (such as National Alliance for Autism Research (NAAR) and Cure Autism Now) which rejected ‘cures’ and called for adaptation of environments to support autistic people, using terminology preferred by self-advocates and community participation. Recognition of the marked heterogeneity within ASD began in the 1970’s with the triad of impairments in language, play and social interaction characterizing many children with intellectual disabilities (ID) or those with classical autism. The first twin study demonstrated that monozygotic twin pairs, though concordant for difficulties associated with ASD, differed in specific characteristics and co-occurring conditions including ID. More recently, phenotypic heterogeneity has been the rule in most, though not all, gene-first phenotypic studies. Thus, developmental aspects of differences in strengths, difficulties and trajectories, as well as biological factors, require highly personalized conceptualizations of the needs of autistic individuals and their families. ABA, applied behaviour analysis; AGRE, Autism Genetic Resource Exchange; CDC, US Centers for Disease Control and Prevention; DSM, Diagnostic and Statistical Manual of Mental Disorders; EEG, electroencephalography; GRASP, Global and Regional Asperger
Syndrome Partnership; IDEA, Individuals with Disabilities Education Act; MCEP, the gene associated with Rett Syndrome; PACT, Preschool Autism Communication Trial; PDD, pervasive developmental disorder; SNAP, Special Needs and Autism Project; SPARK, Simons Foundation Powering Autism Research for Knowledge, TEACCH, Treatment and Education of Autistic and Communication related handicapped Children.

Figure 2. Environmental risk factors for ASD.

Data from studies aiming to identify risk factors for autism spectrum disorder (ASD) can be broadly split into three categories, those with evidence supporting an association (panel a), those with inconclusive evidence (panel b) and importantly, those with no supporting evidence (panel c). Bars represent ranges. *Represents recurrence risk. Figure adapted from 18 with added findings from select reviews and empirical papers: neonatal hypoxia estimate 267, childhood vaccines 20, valporate use during pregnancy 268, parent age estimates 269, preterm birth estimate 270,271, maternal obesity estimate 272, folic acid intake estimate 273, siblings estimate 274,275, interpregnancy interval estimate 276, assisted reproductive technologies estimate 277,278, pesticide and air pollution estimate 279, caesarian section estimate 280.

Figure 3. Encoded proteins associated with ASD risk. [Au: please ensure this figure is added to the third party rights table and we will obtain permission to reuse it on your behalf. Note that I’ve added the protein names in after the ASD syndromic genes, below]

Simplified schematic of the major cellular components of a neural circuit in the cerebral cortex, with a focus on pyramid-shaped glutamatergic excitatory projection neurons. Proteins encoded by selected high-confidence (FDR < 0.1) autism spectrum disorder (ASD) risk genes 34 and proteins encoded by selected syndromic ASD genes have a role in these neurons during development. These proteins have a diverse intracellular distribution; those at the synapse, have roles in cell adhesion, scaffolding and signalling. In addition, some of these proteins are localized to the nucleus and have been shown, broadly, to mediate chromatin modification and transcriptional control. Syndromic ASD genes include FMR1 (encoding fragile X mental retardation protein; fragile X syndrome), UBE3A (encoding Ubiquitin-protein ligase E3A; Angelman syndrome), TSC1 and TSC2 (encoding hamartin and tuberin; tuberous sclerosis complex), PTEN (encoding Phosphatase and tensin homolog) and MECP2 (encoding methyl-CpG-binding protein 2; Rett syndrome). Adapted from 48.

Figure 4. Longitudinal trajectories of total brain volume, surface area and cortical thickness in ASD.

[Au: please ensure this figure is added to the third party rights table and we will obtain permission to reuse it on your behalf] Brain trajectories from 6–24 months of age for total brain volume (TBV, panel a), surface area (SA, panel b) and cortical thickness (CT, panel c). [Au: These data are quite complex and may be difficult for non-experts to understand; accordingly, please briefly describe the key trends in this figure in 2-3 sentences, what are the take-home messages from the figure?] Corrected age [Au: edited Length-age to ‘corrected age’ as per the original figure, ok?] refers to the age corrected by length (body size). From Hazlett et al. 2017.
Figure 5. Co-occurring disorders [Au: this title was too long to adhere to our production guidelines so I've edited it down, ok? I've tried to incorporate the original title into the first line of the legend, is this ok?].

Primary and secondary disorders and disadvantage can accumulate through development in people with autism spectrum disorder (ASD). These disorders can form additional targets for treatment and policy. Prevalence [Au:OK?] estimates from QUEST$^{281}$ SNAP$^{133,282}$ and EDX$^{142}$ cohorts [Au: please advise on how to proceed with Terry’s comment here].

Figure 6. Major parental milestones in advocating and supporting their child with ASD.

Families of children and adults with autism spectrum disorder (ASD) have many decisions and expectations across the lifespan of their children, from seeking initial diagnostic evaluation and intervention to preparing for aging-related services. These decisions [Au:OK?] vary across different cultures, regions and countries and depend on many factors, including the resources and services available. However, several decisions are common across all regions [Au:OK?], including LMR [Au: please define LMR - do you mean LMICs?], such as choices about who will care for their child if the parents are temporarily unable [Au: edits for brevity ok?], the amount of time parents and other family members can spend with the child with ASD versus meeting other needs, ways to modify their home environment to ensure the safety and independence of the individual with ASD and the kinds of behavioural expectations that are most helpful for their child or adult. Of note, for many families, these choices and responsibilities are lifelong and are relevant, for children, adolescents [Au:OK?], adults and elders [Au:OK?] with ASD.

Figure 7. Changes in daily living scores as predicted by IQ scores and autistic symptoms.

Changes in independent daily living skills can be observed in people with ASD over time. This sample consists of ~100 young adults with a mean age of 26 years with autism spectrum disorder (ASD), who were evaluated at 2, 3 and 9 years of age and followed up to 26 years of age. [Au: I’ve incorporated the discussion of how these scores were obtained and what they mean below (highlighted in yellow) for flow, ok? Text discussing divergence and heterogeneity moved to later on in the legend for flow] Daily living scores [Au: instead of ‘outcomes’, ok?] are very diverse, ranging from age-appropriate levels of independence at adulthood (represented by a daily living score of 100, assessed using the Vineland II$^{283}$ [Au:OK?] ) to very limited skills (represented by a score of <30). [Au: green text moved to here from earlier on for flow] Increasing divergence shows where measurement after 2 years of age is additionally predictive, with the line thickness indicating the proportion of early referred children that followed each trajectory. Heterogeneity in intellectual functioning and severity of ASD symptoms (social communication and restricted, repetitive, sensory behaviors) can be observed. In addition, improvements and worsening of autistic symptoms and intellectual functioning can occur over time. A, B| Referred children had verbal IQs predominantly <50 (over 3 standard deviations below average) but could show improvement in daily living standard scores [Au:OK?] from 2 to 3 years of age that were indicative of eventual greater independence in adulthood. Relatively less early change in non-verbal IQ is seen but, like verbal IQ,
adulthood the association with eventual adult daily living skills is strong. C, D | Variation in ASD symptom severity in social-communication (CSS refers to The Autism Diagnostic Observation Schedule, Second Edition (ADOS-2) Comparison Scores) showed a stronger association with adult independence than restricted-repetitive behaviours and continued to change over the lifespan following more divergent pathways than intellectual functioning. Data from the EDX cohort compiled from ¹,¹⁴²,²⁸⁴.
Table 1: ASD as defined in DSM-5.

[Au: I do think this version is improved, however, we can no longer include bullet points within tables as this leads to problems at the layout stage - what about converting this to a text box (see example below)]

<table>
<thead>
<tr>
<th>Domains</th>
<th>Other criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>A. Social communication and social interaction</strong></td>
<td><strong>B. Restricted, repetitive behaviours and interests</strong></td>
</tr>
<tr>
<td>Must have evidence across multiple contexts of all 3 subdomains currently or by history</td>
<td>Must have evidence of 2 of 4 subdomains currently or by history</td>
</tr>
<tr>
<td><strong>Subdomains</strong></td>
<td><strong>Subdomains</strong></td>
</tr>
<tr>
<td>• Social reciprocity</td>
<td>• Stereotyped, repetitive behaviours</td>
</tr>
<tr>
<td>• Nonverbal communication</td>
<td>• Insistence on sameness</td>
</tr>
<tr>
<td>Developing, maintaining and understanding relationships</td>
<td>• Highly restricted, fixed interests</td>
</tr>
<tr>
<td></td>
<td>Hyper- or hyposensitivity or interest in sensory inputs</td>
</tr>
<tr>
<td><strong>Note:</strong> Previously established DSM-IV diagnoses of any pervasive developmental disorder, including Asperger’s disorder should be assumed to be equivalent to DSM-5 ASD</td>
<td><strong>Note:</strong> ASD may co-occur with many other disorders including ADHD, intellectual disability, language delay and genetic syndromes</td>
</tr>
</tbody>
</table>

ADHD, attention-deficit/hyperactivity disorder; ASD, autism spectrum disorder; DSM-5, Diagnostic and Statistical Manual of Mental disorders, Fifth Edition.

Box xx. ASD as defined in DSM-5.

The Diagnostic and Statistical Manual of Mental disorders, Fifth Edition (DSM-5) criteria for autism spectrum disorder (ASD) comprise 5 symptom clusters (A-E)

A. Social communication and social interaction.

- Must have evidence across multiple contexts of all of the following 3 subdomains currently or by history
  - Social reciprocity
  - Nonverbal communication
B. Restricted, repetitive behaviours and interests.

- Must have evidence of 2 of 4 of the following subdomains currently or by history
  - Stereotyped, repetitive behaviours
  - Insistence on sameness
  - Highly restricted, fixed interests
  - Hyper- or hyposensitivity or interest in sensory inputs

C. Symptoms must be present in early development but may not be fully manifest until later or may be masked later in life by learned strategies

D. Symptoms must cause clinically significant impairment in current functioning

E. Not better explained by intellectual disability or global developmental delay

Note: Previously established DSM-IV diagnoses of any pervasive developmental disorder, including Asperger’s disorder should be assumed to be equivalent to DSM-5 ASD. ASD may co-occur with many other disorders including ADHD, intellectual disability, language delay and genetic syndromes.
Table 2: Evidence-based medication in ASD (adapted from Lord et al\textsuperscript{385}) [Au: please ensure this is added to the third party rights table and we will obtain permission to reuse it on your behalf]

<table>
<thead>
<tr>
<th>Medication</th>
<th>FDA or EMA Indication and age</th>
<th>Effect Size (d)</th>
<th>Common Adverse Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Typically used for ADHD symptoms</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methylphenidate</td>
<td>FDA and EMA approval for ADHD (not specific for ASD) in those ≥ 6 years of age [Au: edits ok?]</td>
<td>d=0.78 (teacher rated)</td>
<td>Sleep disruption and decreased appetite</td>
</tr>
<tr>
<td>Atomoxetine</td>
<td>FDA and individual country approval for ADHD (not specific for ASD) in those ≥ 6 years of age [Au: edits ok?]</td>
<td>d=−0.68–0.84</td>
<td>Decreased appetite, nausea and irritability</td>
</tr>
<tr>
<td>Guanfacine</td>
<td>FDA and EMA approval for ADHD (not specific for ASD) in those 6–17 years of age [Au: edits ok?]</td>
<td>d=1.67</td>
<td>Fatigue, sedation and decreased pulse and blood pressure [Au:OK?]</td>
</tr>
<tr>
<td><strong>Typically used to treat agitation and irritability</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Risperidone</td>
<td>FDA approval for irritability associated with ASD and EMA approval only for other indications [Au: what indications?] in those 5–17 years of age [Au: edits ok?]</td>
<td>d=0.94</td>
<td>Increased appetite, sedation and weight gain</td>
</tr>
<tr>
<td>Aripiprazole</td>
<td>FDA approval for irritability associated with ASD and EMA</td>
<td>d=0.87</td>
<td>Nausea and weight gain</td>
</tr>
<tr>
<td>approval only for other indications [Au: what indications?] in those 6–17 years of age</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>---</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

ADHD, attention-deficit/hyperactivity disorder; ASD, autism spectrum disorder; EMA, European Medicines Agency.
**Table 3:** Factors that affect QOL. Cited within table: (Duncan & Bishop) [Au: please ensure this is added to the third party rights table and we will obtain permission to reuse it on your behalf]

<table>
<thead>
<tr>
<th>Type of QOL</th>
<th>Factor</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Objective QOL</td>
<td>Early language</td>
<td>Follow-up studies of adults with ASD who were diagnosed as children have examined the amount of spoken language during early childhood. Individuals with ASD who had fluent speech are more likely to have higher levels of objective QOL life in adulthood than those with phrased speech or those with no speech or who spoke in single words.</td>
</tr>
<tr>
<td></td>
<td>Indicators of intelligence</td>
<td>Studies examining IQ scores using standardized IQ tests administered both in early childhood and adulthood find that individuals with ASD and higher IQ scores have higher levels of objective QOL than those with lower IQ scores [Au: comparator ok?]. Other, less-standardized measures of intelligence (such as those used in large cohort studies [Au: what measures? Please provide another example here]) have similar findings.</td>
</tr>
<tr>
<td>Adaptive behaviour</td>
<td></td>
<td>Higher levels of adaptive behaviour – and particularly more activities of daily living – are associated with better objective QOL in people with ASD. Adaptive behaviour is a challenge for many individuals with ASD, who have scores below what would be expected based on IQ. Adaptive behaviour is changeable, making it a promising avenue for interventions to improve objective QOL.</td>
</tr>
<tr>
<td>Autism symptom severity</td>
<td></td>
<td>Individuals with more severe autism symptoms tend to have lower objective QOL in adulthood.</td>
</tr>
<tr>
<td>Challenging behaviours</td>
<td></td>
<td>Higher levels of challenging behaviours in people with ASD, which can include both internalizing problems and externalizing problems, are related to lower objective QOL.</td>
</tr>
<tr>
<td>Sex or gender</td>
<td></td>
<td>Sex or gender associations with objective QOL have been demonstrated in terms of employment or post-secondary education. Indeed, women with ASD obtain employment and post-secondary educational positions at the same rate as men with ASD but have a more difficult time maintaining those positions over time.</td>
</tr>
<tr>
<td>Subjective QOL</td>
<td>Perceived stress</td>
<td>Many adults with ASD perceive high levels of stress in their own lives. These perceptions are related to lower subjective QOL.</td>
</tr>
<tr>
<td></td>
<td>Supports</td>
<td>Several different types of supports have been related to subjective QOL, including formal services, support from family members (most often parents) and more general social support from others.</td>
</tr>
</tbody>
</table>

ASD, autism spectrum disorder; QOL, quality of life.
References

Highlighted references:

   This paper presents a different, broad overview of the changes in perspective about autism and ASD over the years.

   This paper uses active case-finding to provide representative estimates of the prevalence of ASD and demonstrated that rates of ASD in men and women were equivalent in adults with moderate to profound intellectual disability.

   This seminal paper, through careful recruitment and methodology was the first to show significant early differences that may contribute to our understanding of developmental features in neural structure and circuits.

   This was the first paper to show a de novo loss of function mutation in a synaptic gene associated with non-syndromic ASD and really was a harbinger for so many of the findings that came after.

   This paper was the first to focus explicitly on simplex autism and show the importance of de novo CNVs in simplex cases, versus familial cases, versus controls.

   Paper replicating the mu suppression deficits in autism during action observation but questioning through high-density spectral analyses and sources reconstruction its previously drawn relation to the MNS.

Review supporting that psychiatric disorders are more commonly characterized by impairments of social interaction rather than social observation, and advocating for an interactive turn in neuropsychiatry.


This is a position paper highlighting challenges to translating knowledge on better awareness, understanding, identification, diagnosis (and then treatments) from the past two decades of clinical research in HICs into LMICs


This paper clearly establishes that autism was a stable diagnosis (as a spectrum) beginning at least by age 2. The paper also established parent interview and clinician observation as predictive of autism at age 9. Finally, it was the first paper that showed that the specific DSM-IV-TR diagnoses were unstable across childhood but that the instability was almost all shifting across categories not outside the spectrum.


Despite its potential importance as biological marker and/or subgroup of ASD, developmental regression has remained very poorly understood. This paper outlines recent data and reconceptualization about patterns of onset (and loss) that chimes with a more contemporaneous understanding of ASD as a heterogeneous condition in terms of its manifestation both within and across individuals.


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