Improving treatment of neurodevelopmental disorders:
recommendations based on preclinical studies

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Abstract

Introduction: Neurodevelopmental disorders (NDDs) are common and severely debilitating. Their chronic nature and reliance on both genetic and environmental factors makes studying NDDs and their treatment a challenging task. Here, we discuss neurobiological mechanisms of NDDs, and present recommendations on their translational research and therapy, outlined by the International Stress and Behavior Society.

Areas covered: Common NDDs are a heterogeneous group which includes autism spectrum disorder, intellectual disability, communication/speech disorders, motor/tic disorders and attention deficit hyperactivity disorder. Various drugs currently prescribed to treat NDDs also represent a highly diverse group. Acting on various neurotransmitter and physiological systems, these drugs often lack specificity of action, and are commonly used to treat multiple other psychiatric conditions. There has also been relatively little progress in the development of novel medications to treat NDDs. Based on clinical, preclinical and translational models of NDDs, our recommendations cover a wide range of methodological approaches and conceptual strategies.

Expert opinion: To improve pharmacotherapy and drug discovery for NDDs, we need a stronger emphasis on targeting multiple endophenotypes, a better dissection of genetic/epigenetic factors or ‘hidden heritability’, and a careful consideration of potential developmental/trophic roles of brain neurotransmitters. The validity of animal NDD models can be improved through discovery of novel (behavioral, physiological and neuroimaging) biomarkers, applying proper environmental enrichment, widening the spectrum of model organisms, targeting developmental trajectories of NDD-related behaviors and comorbid conditions beyond traditional NDDs. While these recommendations cannot be addressed all in once, our increased understanding of NDD pathobiology may trigger innovative cross-disciplinary research expanding beyond traditional methods and concepts.

Keywords: neurodevelopmental disorders, animal models, autism, ADHD, translational research
Article Highlights:

- Neurodevelopmental disorders (NDDs) are common and widespread psychiatric illnesses caused by aberrant brain development.
- The chronic nature of NDDs and their strong reliance on both genetic and environmental factors complicates their treatment and prevention.
- Drugs currently used to treat these disorders lack specificity and are often ineffective.
- Here, we discuss neurobiology of NDDs and their available preclinical (experimental) animal models.
- Based on these analyses, we formulate recommendations to improve NDD drug discovery and pharmacotherapy.
- Improved understanding of NDD pathobiology is critical for fostering further innovative cross-disciplinary research in the field of translational neuroscience and biological psychiatry.
List of abbreviations:

ADHD - Attention Deficit/Hyperactivity Disorder
ASD - Autism Spectrum Disorder
CAM – complementary alternative medicine
CNS – central nervous system
CD – Communication Disorder
DSM – Diagnostic Statistical Manual
FDA – Federal Drug Administration
fMRI – functional magnetic resonance imaging
GxE – gene-environment (interaction)
G-E – gene-environment (correlation)
GWAS - genome-wide association study
ID - Intellectual Disability
ncRNA- non-coding RNAs
NDD – neurodevelopmental disorder
PPI - pre-pulse inhibition
PTSD – post-traumatic stress disorder
RDOCs - Research Domain Criteria
SERT – serotonin transporter
SHR – spontaneously hypertensive rat
SLD - Specific Learning Disorder
SNP - single nucleotide polymorphism
SSRI – selective serotonin reuptake inhibitor
WGS – whole-genome sequencing
WES - whole-exome sequencing
1. Introduction

Neurodevelopmental disorders (NDDs) are caused by aberrant brain development resulting in cognitive, motor, language and affective deficits (Table 1). Common NDDs include autism spectrum disorder (ASD), social communication disorders, intellectual disability (ID), attention deficit hyperactivity disorder (ADHD), motor and tic disorders (Fig. 1a). Aberrant neural development, usually beginning during early embryogenesis, causes a lasting dysregulation of a wide range of brain processes, including neurogenesis, glia/neuronal proliferation, cell migration, synapse formation and myelination (Fig. 1b). As genetic factors play a key role in NDDs (showing medium-to-high heritability estimates; Fig. 1b), they can also be evoked by environmental factors, such as maternal influences, nutrition or early exposure to immune challenges, toxicants, psychoactive drugs and stress. In addition to clinical studies, various animal models of experimental NDDs are widely used in translational research in this field (Tables 2-4).

The growing socio-economic impact and high prevalence of NDDs require urgent attention to improving their therapy. The treatment and prevention of NDDs are complicated by their chronic nature and strong dependence on both genetic and environmental factors. With little progress in the development of novel medications for NDDs, the existing drugs often lack specificity of action, and are mainly used to treat other psychiatric conditions beyond NDDs (Table 3). Recognizing these important challenges, the International Stress and Behavior Society (ISBS) has established the Strategic Task Force on NDDs. Comprising international experts from different fields of biological psychiatry, the Panel comprehensively evaluated the neurobiological mechanisms, genetics, psychopharmacology and in-vivo animal models of NDDs. Based on mounting translational and preclinical evidence, the ISBS Task Force has made critical recommendations for improving pharmacotherapy of NDDs, which are summarized here.

2. The complex genetics of neurodevelopmental disorders

Described as a “ballet choreographed over time between the action of multiple genes, environmental and epigenetic factors”8, brain disorders often display complex, polygenic non-Mendelian
genetics\textsuperscript{9}. Used to identify disease-causing variants, genome-wide association studies (GWAS) utilize multiple single nucleotide polymorphisms (SNPs) maintained in linkage disequilibrium\textsuperscript{10}. However, the contribution of GWAS to our understanding of brain disorders, and NDDs in particular, has been limited so far\textsuperscript{9}, partially due to the small sample sizes, the difficulty of ascertaining how SNPs alter gene function, the variability of clinical NDD symptoms and the poor operationalization of these symptoms into measurable entities. The common variants uncovered by GWAS only account for a small percentage of heritability estimates, with surprisingly poor correspondence of data across studies\textsuperscript{10, 11}. Yet, in some cases, genome-wide approaches followed by fine-mapping may provide useful insights into the genetics of NDDs. For example, Latrophilin 3 (LPHN3) was identified as a human ADHD ‘susceptibility’ gene by linkage analysis of affected families\textsuperscript{12} and is now being confirmed in preclinical models of this disorder\textsuperscript{13}.

Increasing evidence indicates that some NDDs (e.g., ASD and ADHD) can also be triggered by mutations in single genes\textsuperscript{14-17}. These genes often contain rare mutations with <1\% of the minor allele frequency (the rate at which the less common allele occurs within a given population)\textsuperscript{10}. Rare mutations include both SNPs and copy-number variants\textsuperscript{18} - deletions or duplications of chromosome segments of variable size that can affect single or multiple genes at the same time. Rare mutations can impact brain and its development in several ways. In addition to reducing gene function, disease-causing mutations may constitutively activate proteins, show dominant negative activity, or create new or abnormal biochemical functions\textsuperscript{19, 20}. Furthermore, a single disease gene can be mutated in multiple positions, each of which may give rise to a very different phenotype. For example, a defect in a single gene, Disrupted in Schizophrenia 1 (DISC1), can lead to different disorders, including schizophrenia (psychoses), bipolar disorder, major depression or ASD, even when patients carry the same mutation\textsuperscript{21}. Therefore, for NDDs with an unknown genetic basis, it may be critical to identify causative mutations in both known and novel genes.

The non-Mendelian inheritance of some single disease-causing mutations can be explained by incomplete penetrance of the disease-causing alleles, by protective variants in the genetic background or
by the acquisition of novel mutations. Single mutations provide an excellent entry point for studying NDDs, as they are easier to model in animals, provide construct validity for the disease and offer opportunities to uncover the underlying neural circuits\textsuperscript{11}. Furthermore, comparison of the effect of multiple mutations may also highlight the general changes in brain function that lead to pathogenesis\textsuperscript{22}: just as “all roads lead to Rome”, alteration of the function of many genes may ultimately affect only a few common disease-causing pathways\textsuperscript{23} (see further). However, the relative contribution of rare or common mutations to disease susceptibility is currently unclear. Rather than being caused by a single type of genetic lesion, NDDs could be caused by a combination of many. For example, single mutations can predispose patients to ADHD, while other SNP polymorphisms in the genetic background (or mutations in a second critical gene) may alter the penetrance of the disease\textsuperscript{11}. Thus, a single mutation may be necessary, but not sufficient, to trigger the disorder. Such a combined model may explain the large heterogeneity of NDD symptoms and their low penetrance often observed in patients and in preclinical models\textsuperscript{24}.

Since most of the rare mutations linked to NDDs affect genes active during embryonic development, even subtle changes can have far-reaching consequences for neural development and permanently alter the mature brain function\textsuperscript{25}. In some cases, genetic mutations cause specific cellular phenotypes linked to neurological disorders, impairing symmetric cell division, differentiation, survival, neurite outgrowth, axon pathfinding and dendritic architecture\textsuperscript{26, 27}. Alternatively, mutations may cause diffuse and variable changes to brain function by influencing the outcome of random variation at a number of developmental choice points, thereby affecting the global developmental trajectory of an organism\textsuperscript{28}. Such broad phenotypic changes may cause miswiring of neural circuits, leading to altered brain function and pathophysiology\textsuperscript{29}. Overall, diffuse changes to brain function seem likely when considering that genes do not directly control behavior, but rather act via the formation, connection and function of neural circuits in the brain. Thus, it is perhaps predictable that many disease-causing mutations are active during embryonic development, a time when embryos may be both more plastic and more susceptible to alteration.
3. **Recommendations of the ISBS Strategic Task Force on neurodevelopmental disorders**

Outlined here are several strategies suggested by the ISBS Panel to foster translational research into the pathogenesis and treatment of NDDs (also see Table 5). Covering different aspects of translational cross-species and multi-phenotype modeling, these recommendations are expected to improve current and future drug discovery and pharmacotherapy of NDDs.

**4.1. Focus on multiple (vs. single) endophenotypes**

While prenatal neural development is associated with cell differentiation, neuronal outgrowth and network formation (Fig. 1b), after birth, these networks are refined by the experiences of the newborn that impinge on the ongoing developmental processes. Therefore, it is insufficient to elucidate neurodevelopmental mechanisms by studying individuals with full-blown disordered phenotypes. Indeed, it has been demonstrated that NDDs are associated with susceptibility genes that are relatively upregulated in the prefrontal cortex during fetal life\(^3^0\). For example, genes associated with syndromic NDDs (e.g., ID and ASD), are relatively enriched in prenatal transcript abundance compared with the overall transcriptome, whereas genes associated with neurodegenerative disorders are significantly under-expressed during fetal life\(^3^0\). In addition, during normal prefrontal cortex development, gene splicing changes are most frequent early in development. More than 60% of all splicing changes represented a single splicing pattern reflecting preferential inclusion of gene segments potentially targeting transcripts for nonsense-mediated decay in infants\(^3^1\). Risk for NDDs is increased when nonsense-mediated mRNA decay becomes aberrant\(^3^2\). Thus, it is essential to follow developmental processes before phenotypes become overt. Some of these processes may remain sub-threshold and not present recognizable features. Even in this case, a feature appearing will not lead to a diagnosis. As a consequence, a continuum in developmental processes and associated phenotypes is expected. Longitudinal and prospective cohort studies providing information about maternal and paternal genotype and personality, prenatal factors, and genotype, brain morphology and function, life events and developmental milestones can provide important insights into this continuum of developmental processes and phenotypes that predispose to the full-blown diagnosed disorder. However, such studies require a
large number of subjects and more budget, given clinical heterogeneity of NDDs (Fig. 1a-c). Traditionally, experimental modeling of NDDs and other brain disorders is performed by mimicking their behavioral (affective, cognitive, social, motor or reward) endophenotypes. Current preclinical studies of NDDs may therefore benefit from modeling the dynamic interplay of multiple pathogenic endophenotypes (and their sensitivity to pharmacotherapy) rather than focusing on individual phenotypes or their screens. Note, however, that this approach can be complicated in longitudinal studies (revealing phenotypes and mechanisms occurring before the onset of the full-blown disorder). Thus, development of novel ‘early’ biomarkers or endophenotypes of NDDs, and performing both short-term and ‘clinic-like’ longitudinal experiments in animal models, is recommended.

4.2. Dissect genetic and epigenetic influences

Modern genetic sequencing tools have revealed multiple previously unknown genes linked to NDDs. Many of the sequencing studies performed to date have utilized whole-exome sequencing (WES), which focuses only on the ~2% of the human genome that encodes proteins. Recently, the Encyclopedia of DNA Elements (ENCODE) project found that the non-coding regions of the genome not sequenced in typical WES play crucial regulatory functions in cellular dynamics and pathways. In contrast, whole-genome sequencing (WGS) attempts to scan the majority of the genome for disease-linked genetic variants, and is successful in small cohorts of ASD patients. While WGS poses financial challenges, this Panel recommends an increased focus on WGS, as this may lead to further discoveries of rare variants in non-coding RNAs (ncRNA) and other regulatory regions contributing to NDDs. The examination of non-coding genomic regions and ncRNA also highlights the need for detailed epigenetic analyses of NDDs. Indeed, epigenetics play a critical role in the pathogenesis of Angelman syndrome (loss of imprinting at UBE3A), Fragile X syndrome (hypermethylation of FMR1 promoter caused by trinucleotide expansion), and Rett syndrome (mutation in the epigenetic enzyme MECP2). Mutations in genes coding for chromatin-modifying enzymes were also recently identified in ASD. Therefore, we call for the detailed investigation of both epigenetic markers (e.g., histone acetylation/methylation and DNA methylation) in NDDs, as well as deep sequencing analysis of genes encoding epigenetic enzymes (such
as MeCP2, DNA methyltransferases and histone deacetylases/acetyltransferases), since mutations in these enzymes may play a role in stochastic epigenetic variation and transgenerational epigenetic effects (e.g., 79).

4.3. Focus on symptoms that bridge across several neurodevelopmental disorders

The symptoms of NDDs frequently overlap (Fig. 1b), providing an opportunity to uncover common down-stream pathways that affect multiple diseases. For example, aggression is a common comorbid symptom of ASD, ADHD, schizophrenia, conduct disorder and some types of depression34. Although heightened aggression levels can prevent treatment of other symptoms of a disease, the neural circuits that control this behavior are not well understood. Additionally, there are currently few medications that specifically target this behavior and, therefore, an urgent need for effective pharmacotherapy. Therefore, studying the genetic and neurological basis of behaviors that are present across neurodevelopmental disorders may provide insight into each separate disease. This recommendation is generally consistent with the recently proposed Research Domain Criteria (RDOCs) of the US National Institutes of Health (NIH) to address mechanisms and phenotypic dimensions, rather than symptomatic categories, of psychiatric diagnoses80. By focusing on discrete phenotypes, rather than on whole, criteria-based NDDs, new insights may be gained that would have otherwise remained obscured.

4.4. Address the problem of ‘hidden heritability’

The problem of ‘hidden’ (missing) heritability35 is highly relevant to pathobiology of NDDs, since impaired neural development often causes global behavioral and cognitive deficits. This, in turn, may mask disorder-specific phenotypes, which may then become undetectable by conventional (e.g., GWAS) analyses. Recognizing its potential importance, we call for more thorough analyses of ‘hidden heritability’ of NDDs in both clinical and preclinical studies. Additionally, given the importance of epigenetic factors in CNS modulation, detailed analyses of epigenetic variation81 (both inherited and random) may further contribute to the "missing heritability" of complex brain disorders, such as NDDs. For example, genetic risk for a specific NDD may be passed through the germline by either “true”
epigenetic inheritance (i.e., an epigenetic mark that is present in the germline and inherited across multiple generations) or by mutations in epigenetic enzymes which may bias the chromatin confirmation of a patient\textsuperscript{36,37}, thus increasing the risk for diagnosis of a particular NDD.

4.5. Examine potential developmental role of neuromediators

Serotonergic drugs, such as SSRIs, are commonly used to treat NDD symptoms (Table 3). Mounting evidence suggests that some neurotransmitters, such as serotonin, may play a dual role in the brain, acting as both classical neurotransmitters and developmental ‘neurotrophic’ hormones\textsuperscript{38}. For example, although the genetic ablation (knockout) of serotonin transporter (SERT) in mice shares some similarities with the effects of postnatal SSRI exposure, some differences exist, including anxiety and elevated extracellular serotonin levels in SERT knockouts vs. reduced serotonin levels and anxiety in adult mice following early postnatal SSRI (escitalopram)\textsuperscript{39}. Therefore, it remains unclear to what extent serotonin-mediated neurodevelopmental changes contribute to serotonin-related traits, and whether such traits are solely dependent on serotonin levels \textit{per se}. The developmental phase during which serotonergic perturbations take place can play a crucial role, as during the late prenatal phase in rodents serotonergic neurons grow and migrate, whereas at the early postnatal phase brain circuits are shaped and refined\textsuperscript{40}. Furthermore, it remains unclear whether and which serotonin-mediated structural neurodevelopmental changes contribute to behavioral phenotypes of NDDs. Therefore, approaches like in-utero electroporation and manipulating Sert expression during embryonic development may help clarify the postnatal structural and functional neurodevelopmental role of serotonin. Focusing on developmental processes during the postnatal period, the use of conditional knockout animals may further reveal how changes in neural development contribute to NDD-like behavioral phenotypes. For example, SSRIs used during pregnancy cause subsequent developmental delays, increased social behavioral disturbances and increased risk for ASD\textsuperscript{41,42} in children. In addition, SSRIs induce chromatin alterations, suppressing protein kinase calmodulin-dependent protein kinase II (CaMKII) transcription in the rat nucleus accumbens\textsuperscript{36}. Collectively, this suggests a larger (than traditionally recognized) role for central serotonin, as it can modulate NDDs \textit{directly} as neurotransmitter, and \textit{indirectly} as a neurotrophic developmental
hormone. Investigating this emerging complexity, and its implications for pharmacotherapy of NDDs, are recognized by this Panel as an important strategic direction of research.

4.6. Improve the validity of animal models and biomarkers

Given their immense translational importance, we recommend the continued use of genetically-modified animal models to investigate the neurobiology of NDDs, with a particular focus on genes involved in synaptic function and epigenetic modification\textsuperscript{2,60}. However, efforts should also focus on the development of environmental epigenetic models that recapitulate early life factors and may be involved in these disorders. The combination of these preclinical models will be crucial in identifying gene x environment interactions important for the progression of NDDs. Notably, basic research continues to over-utilize male animals and cells\textsuperscript{43}. As various NDDs demonstrate robust sex differences (Fig. 1), both sexes must be included in a balanced manner in preclinical studies to improve drug discovery. Likewise, with the growing availability of rodent or fish strains for NDD research, a few selected inbred strains are typically used for drug screening assays, to ensure a better genetic control of the experiments\textsuperscript{44}. As the global human population is highly heterogeneous genetically, a more balanced use of both inbred and outbred animal strains in CNS drug discovery may lead to more valid treatment and side-effects data, reflecting ‘demographic’ aspects of drug action and providing important clinical insights. Finally, although biological markers are not yet approved as part of the diagnostic criteria for NDDs, they may help predict the potential disease trajectory, and support decisions for specific early therapeutic and prophylactic measures. Other well-validated physiological biomarkers relevant to NDDs include fMRI imaging and, more broadly, other electrophysiological markers (e.g., mismatch negativity) which have shown high sensitivity to NDDs\textsuperscript{80, 81}. We strongly emphasize the need to validate and discover new predictive markers of neurodevelopmental markers, in order to improve both drug discovery and pharmacotherapy.

4.7. Widen the spectrum of model organisms

Preclinical animal models are a critical tool in biological psychiatry\textsuperscript{45}, including translational modeling of NDD-related phenotypes and pathogenesis. For example, rodent models of social
interaction, communication deficits and repetitive behaviors are relevant to studying ASD, whereas rodent hyperactivity and/or inattention can parallel clinical ADHD. Although rodents are currently the most commonly used animal model in the pre-clinical research (Fig. 2), they are limited by lower throughput and costs. Therefore, increasing the spectrum of model organisms is recognized as an important strategy in biological psychiatry, and is fully endorsed by the ISBS. This endorsement is even more based on the conceptual value of identifying core, evolutionarily conserved pathogenic mechanisms of brain disorders – which can only be achieved by modeling brain disorders across taxa. For example, zebrafish (Danio rerio) are a time-efficient and low-cost model with high genetic and physiological homology to humans (Fig. 2). They also show rapid development, which beneficial for NDD models larval zebrafish display robust motor and affective behaviors (which can be then monitored throughout the 4-year lifespan). In adult zebrafish, group behavior (shoaling) can be a useful model of ASD, whereas hyperlocomotion, impulsive swimming and inattention in both larval and adult zebrafish can be relevant to ADHD. Chicks (Gallus gallus) also display robust social and locomotor phenotypes, recapitulating several NDD symptoms, including social deficits, accompanied by aberrant brain growth and affective behaviors. Likewise, non-human primates, such as common marmosets (Callithrix jacchus), represent another useful model organism with robust social and affective behaviors highly sensitive to environmental manipulations that cause NDD-like phenotypes (e.g., ASD-like social deficits or ADHD-like pathological hyperactivity). Finally, complementing vertebrate models, invertebrate models (e.g., fruit flies, Drosophila melanogaster) are also demonstrating potential for modeling some aspects of NDDs. Although these models have been comprehensively evaluated in the literature (see above), and will not be discussed here in detail, increasing the spectrum of model organisms is recognized by this Panel as one of the top priorities in translational NDD research.

4.8. Assess the role of environmental enrichment

There is currently a growing interest in the impact of early life adversity on neural development. Less attention, however, has been paid to the effects of environmental enrichment. Rearing laboratory animals in a complex, enriched environment can stimulate the development of the CNS, aid in
recovery of brain function following acquired injury, and improve performance on many cognitive and behavioral tasks, affording it special interest in the study of NDDs\textsuperscript{37, 60, 61}. The potential for modulation of NDD phenotypes by enriched environments has received some attention (e.g., in particular, with respect to refinement of animal models), and evidence is emerging that the effects of some teratogen-linked NDDs can be improved by environmental enrichment\textsuperscript{62}. For example, environmental enrichment reverses pre-pulse inhibition (PPI) deficits, alterations in pain sensitivity, stereotypic behavior, and social behavioral deficits in a mouse model of ASD induced by pre-natal valproate exposure\textsuperscript{62}. Furthermore, environmental enrichment can significantly improve both the physical and behavioral response to prenatal alcohol exposure in rodents\textsuperscript{63-65} and enhance cognitive performance in spontaneously hyperactive rats (SHR), a genetic model of ADHD\textsuperscript{66}. Although the mechanisms of action on NDD-related phenotypes are not particularly well understood, they may include the modulating effects of environmental enrichment on brain neuroplasticity and the efficacy of dopamine receptors in the pre-frontal cortex\textsuperscript{67, 68}. Therefore, gaining a more thorough understanding of the role of environmental enrichment on the modulation and mediation of NDDs will be of critical importance moving forward. Additionally, the Panel emphasizes the interaction of genetic factors and environmental enrichment on the affectation of NDDs as a potential priority for further research in this field.

4.9. Address Gene-Environment interactions vs. Gene-Environment correlations

The Gene x Environment \textit{interactions} (GxE) characterize how genotypes modify the sensitivity to environmental factors, and are widely analyzed in neuropsychiatric research. The gene-environment (G-E) \textit{correlations} represent another important source of variation for complex behavioral traits, reflecting the individuals’ exposure to the environment as a function of their genotype\textsuperscript{69}. The understanding that environment is not a static uniform factor, but can partly depend on individual’s own behavior, is an important conceptual development in modern biological psychiatry\textsuperscript{69}. This concept has direct implications for NDDs and their association with environmental factors, especially those which are behaviorally modifiable (e.g., social environment). Consider, for example, parent-
offspring and early peer-peer interactions, which represent ‘early’ reciprocal social interactions during neural development\textsuperscript{69}. While most animal studies view the ‘environment’ as a static factor equally imposed on all animals in the laboratory, our ‘real life’ experience depends on subjects’ behavior with parent, relatives or peers. For children with genetically caused poorer social skills (e.g., ASD), their peculiar mode of interactions with others can make social environment more adverse, compared to ‘smoother’ healthy subjects. On the one hand, such behaviorally-mediated interactions with the environment may affect neural development in a correlated (or even synergistic) manner. On the other hand, breaking this vicious pathogenetic cycle may reduce the negative impact of environmental adversity, therefore contributing to a better therapeutic outcome. The Panel recognizes the growing importance of G-E correlations in NDDs, and recommends more animal and clinical studies of reciprocal influences between environment and neural development.

4.10. Use imaging phenotypes in the field of neurodevelopmental disorders

Modern imaging technologies enable \textit{in vivo} monitoring of NDDs and \textit{in vivo} assessment of the efficacy of novel pharmacological treatments. Magnetic resonance imaging and diffusion weighted imaging are established longitudinal methodologies for identifying neuroanatomical alterations. Three-dimensional brain atlases exist for a range of species and enable automatic segmentation of brain regions to identify differences in volume or white matter microstructure\textsuperscript{70-75}. These imaging modalities have been widely used to study various NDDs, including ASD, ADHD and ID\textsuperscript{76}. More recent functional imaging techniques have begun to identify the neural circuitry linked to the behavioral and social phenotypes relevant to NDDs. For example, manganese-enhanced magnetic resonance imaging\textsuperscript{77-79} and functional magnetic resonance imaging (fMRI)\textsuperscript{80, 81} can identify abnormal circuitry, while optogenetics permits identification and precise manipulation of the circuitry related to neurological disorders\textsuperscript{82-84}. \textit{In vivo} imaging also plays an important role in drug discovery – e.g., as it helps to establish vital biomarkers that are necessary to determine if therapeutic candidates have elicited their targeted biological effects\textsuperscript{85}. Following identification of a disease biomarker, imaging can then be used to profile drug mechanisms, efficacy, and safety. A wide range of such techniques include ultrasound, magnetic resonance
spectroscopy, positron emission tomography, single-photon emission computed tomography, and optical imaging methods (e.g., near-infrared fluorescence microscopy). Method to generate transparent brains are now also emerging, which jointly with novel microscopic techniques enable 3D brain imaging. Each technique has different spatial resolutions and time scales, resulting in unique advantages and limitations. As a result, studies should carefully consider which technique(s) are best suited to produce desired conclusions. Overall, the Panel recognizes the value of the diversity of neuroimaging tools, as well as simultaneously analyzing physiological biomarkers and genomic/epigenetic mechanisms. We strongly believe that increased understanding of NDDs must be paralleled by advances in imaging. Imaging can play a large role in many suggestions of the Panel listed here, such as identifying similarities and differences between sexes or strains, developing canonical template spaces for new animal models, investigating the effects of environmental enrichment, and longitudinal investigations of brain development. Ultimately, by combining with existing behavioral and social paradigms, imaging methodologies will further foster translational research of NDDs.

4.11. Increase focus on neurodevelopmental trajectories

Many NDDs are triggered during embryonic development and their symptoms predominantly affect children or young adults. However, the majority of studies in animal models have focused on the behavior and neurobiology of fully mature adults. We recommend that research be extended to more actively include juvenile animals as well. The adolescent brain undergoes widespread developmental changes that affect axon guidance and proliferation, circuit wiring, and synaptogenesis and pruning. These structural changes are associated with the modulation of genetic and epigenetic factors. As NDDs likely affect these processes, a critical step will be to compare neuronal position, connectivity and the function of neural circuits in brains that are not yet fully mature, as well as their underlying molecular mechanisms. For example, ADHD has been linked to a delay in brain maturation and aberrant myelination, manifested as a reduced volume and aberrant microstructure of the white matter. Furthermore, the effect of pharmacotherapy on brain development trajectories should also be investigated further (Fig. 3).
4.12.  Widen the spectrum of disorders relevant to NDDs

NDDs frequently overlap with other neuropsychiatric disorders, including anxiety, depression, schizophrenia and post-traumatic stress disorder (PTSD, Fig. 1). Therefore, expanding the spectrum of NDD-related disorders is an important strategy in translational NDD research, and can include modeling brain illnesses that are not conventional NDDs. Consider, for example, the potential link between NDDs and PTSD, as neurodevelopmental deficits can later make an adult individual more vulnerable to stress, thereby triggering or exacerbating developing PTSD after a traumatic event.90, 91 Alternatively, depending on age, early traumas (e.g., child abuse) could potentiate neurodevelopmental deficits, which will parallel the developing PTSD symptoms. Thus, vigilant pharmacotherapy of PTSD may represent an important adjunct therapy to prevent or reduce NDDs (Fig. 1). The finding that some NDD symptoms (Table 3) and some forms of PTSD exhibit improvement following treatment with SSRIs suggests a role of the serotonergic system in these mechanisms.92 Two SSRIs, paroxetine and sertraline, are currently the only FDA-approved pharmacological treatments for PTSD. However, response rates to SSRIs in PTSD patients rarely exceed 60%, whereas full remission following SSRI treatment is achieved in only 20-30% cases.93 An indication of the accentuated sympathetic activity in PTSD patients is the hyperresponsivity to the administration of yohimbine, an α2-adrenergic receptor antagonist that inhibits noradrenergic autoreceptors and increases central norepinephrine activity.94 Along with a generally greater baseline norepinephrine levels in PTSD patients, this indicates a critical role of this neurotransmitter in the hyperarousal component of PTSD. Based on observed hyperactivity of the noradrenergic system in PTSD, the efficacy of anti-noradrenergic drugs in treating the disorder is supported by the use of propranolol, a β-adrenergic receptor antagonist, to reduce PTSD symptoms when administered after a traumatic event or with re-experiencing a traumatic memory.95 Moreover, in preclinical studies, stress consistently increases glutamate levels, inhibits glutamate uptake, increases the expression and binding of glutamate receptors and increases calcium currents. For example, the primary effect of tianeptine involves the stabilization of glutamatergic neurotransmission and the enhancement of synaptic plasticity, particularly under stress conditions – the two mechanisms that may bridge NDD-
and PTSD-related pathogenesis. Taken together, this evidence suggests that examining disorders not normally considered as NDDs, such as PTSD, may provide new important insights leading to novel treatments that may be beneficial for comorbidity with, and/or target some symptoms of, NDDs.

5. Conclusion

In recent years, research into NDDs has revealed important information about their pathogenesis and underlying biology. However, due to the challenging and complex nature of NDDs and their shared or unique etiologies, many of the neural mechanisms involved remain unknown, resulting in a lack of effective treatment options for patients. Increasing the throughput of the translational pipeline from in-vitro assays to clinical trials (Fig. 3), and using mechanistic insights generated from experimental observations, will lead to novel effective, target-specific therapies available to patients suffering from NDDs.

5. Expert Opinion

Translational in-vivo research aims to replace symptomatic drug therapies with those based on a principled understanding of the disease causes. In relation to NDDs, this process can be presented as a translational cycle with four multidisciplinary steps (Fig 2). The first step, animal modeling, uses animals to examine cognitive, motor and social behaviors, and their underlying genetics, molecular biology, neurophysiology and anatomy. Next, therapeutic strategies step aim to design interventions based on biological findings in animal models and optimize drug safety and efficacy. The drug development step then involves optimization of lead compounds to improve drug-target specificity, bioavailability and pharmacokinetics, as well as to determine the most efficacious dose, dosing strategy and route of administration. Finally, clinical trials examine the drug’s therapeutic effects as well as their behavioral and cognitive effects in humans (Fig. 2 and 3).

Careful study of brain pathology and behavior establishes how well an animal model represents human disease (Step 1). Therapeutic strategies are evaluated in animal models by measuring deficits in performance in various behavioral tasks (Step 2, Table 2). A particular therapeutic strategy may address only a subset of cognitive or motor functions, so multi-target therapies that correct distinct deficits and
brain areas may be desirable. Drug development optimizes a therapeutic compound to improve drug-target specificity, reduce or eliminate dangerous side effects, and determine dose and route of administration (Step 3). Finally, a lead compound enters clinical trials to test the safety and efficacy of therapeutic strategies discovered in animal models (Step 4). An approved pharmacotherapy for a NDD (see Table 3 for a partial listing of currently used and approved drugs) would reduce one or more cognitive deficits or maladaptive behaviors. Such a therapy completes the translational cycle by addressing the disease phenotype. However, animal models are often an imperfect representation of human disease or developmental disorders, and the differences between species may carry special significance for disease pathology\textsuperscript{97-99}. Thus, improved characterization of animal models of NDDs requires better behavioral assays and physiological measurements. In addition, a better dissection of genetic determinants of NDDs is necessary - especially since, as already mentioned, even a single mutation can result in different CNS diseases within family pedigrees\textsuperscript{100}.

New animal models developed in the future will hopefully improve the correspondence with human conditions we hope to achieve, especially since more specific and efficacious second-generation therapies will require improved descriptions of the mechanisms underlying successful pharmacotherapeutic intervention. Once new and effective therapies are developed for NDDs by preclinical studies, designing and executing their clinical trials will require coordinated effort and significant resources from academia, government, private foundations and pharmaceutical companies.

Discussing the neurobiological mechanisms of NDDs, the Special Task Force of the International Stress and Behavior Society (ISBS) has presented recommendations on improving drug discovery, pharmacotherapy and translational research of NDDs. Based on clinical, preclinical and translational models, these recommendations cover multiple areas ranging from methodological considerations to conceptual strategies of future research (Fig. 3). We realize that these recommendations cannot be addressed all in once, and that studies employing experimental animals will always remain an approximation of the human condition. Despite these limitations, increasing our awareness of factors
relevant for improved understanding of NDDs will advance this field and may trigger innovative cross-disciplinary research that goes beyond traditional methods and concepts.

**Declaration of Interest:**

This collaborative study was initiated and coordinated by the International Stress and Behavior Society (ISBS) Task Force on Neurodevelopmental Disorders. SN, MK and HY are supported by Grants-in-Aid for Scientific Research (KAKENHI) of the Japan Society for the Promotion of Science (JSPS. Grants 25282221, 21200017, 25119509 and 15K15404. This research is also supported by Guangdong Ocean University (AVK, CS), ZENEREI Research Center, St. Petersburg State University, the Government of the Russian Federation (Act 211, contract 02-A03.21.0006; AVK), and the Institute of Experimental Medicine RAS (VMK). AVK is the Director of ZENEREI Research Center and President of ISBS. The authors have no other relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript, apart from disclosed.
Figure 1. Neurodevelopmental disorders (NDDs) and their treatment. Panel A outlines therapeutic approaches to treating NDDs and associated other neuropsychiatric illnesses (see Table 1). ASD – autism spectrum disorder, ADHD – attention deficit hyperactivity disorder, OCD – obsessive compulsive disorder, PTSD – post-traumatic stress disorder. Panel B shows the ontogenetic timeline of key neuronal processes related to neural development, as well as the prevalence and heritability estimates (bottom left) and key behavioral symptoms (bottom right) of NDDs; ID – intellectual disabilities, CD – communication disorders. Panel C shows high comorbidity rates among NDDs and with selected other related disorders (ICD – impulse control disorder).
NDDs

- Intellectual Disabilities
- Communication Disorders
- ASD
- ADHD
- Specific Learning Disorder
- Motor Disorders
- Tic Disorders

Improved NDD outcome

NDD Pharmacotherapy

Adjunct therapy

Obesity
Depression
Anxiety
PTSD
OCD
Psychoses
Sleep disorders
Aggression
Seizures
Addiction
B

Prenatal

Postnatal

Proliferation

Migration

Synaptogenesis

Myelination

Neurogenesis

Prevalence

Medium Heritability estimates

High Heritability estimates

Key symptoms

Language

Social deficits

Repetitive behaviors

Cognitive deficiencies

Aggression

Hyperlocomotion

Cognitive deficiencies

Depression

OCD

Psychoses

ICD
Figure 2. Steps involved in *in-vivo* development of anti-NDD therapies. Animal size reflects the relative usage of respective species (rodents, chicks, non-human primates, zebrafish, fruit flies) in NDD research.
Figure 3. The proposed integrative approach to improving pharmacotherapies of clinical neurodevelopmental disorders (NDDs), based on the recommendations proposed by the ISBS Task Force on NDDs

- Use a wider spectrum of drugs with different mechanisms of action
- Examine a wider range of genetic, epigenetic and environmental factors (e.g., enrichment), gene x environment interactions and G-E correlations
- Using more behavioral measures
- Using more physiological biomarkers
- Examining a wider spectrum of disorders
- Using more model organisms
- Using neuroimaging biomarkers
- Tracing developmental trajectories

Drug 1
Drug 2
Drug 3
In-vitro
Animal models
Clinical NDDs
<table>
<thead>
<tr>
<th>Disorders</th>
<th>Major symptoms</th>
<th>Availability of animal models</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intellectual Disabilities (ID):</td>
<td>Impaired mental functions in conceptual (language, reading, writing, knowledge, interpretation), social (empathy, compassion, judgment, communication, harmony) and practical (personal care, financial management, hobby) aspects</td>
<td>+</td>
</tr>
<tr>
<td>Communication Disorders</td>
<td>Difficulties in language, speech, phonetic fluency or social communication</td>
<td>?*</td>
</tr>
<tr>
<td>Autism Spectrum Disorder (ASD):</td>
<td>Persistent deficits in reciprocal social communication and interaction, as well as restricted, repetitive patterns of behavior, interests or thoughts</td>
<td>+</td>
</tr>
<tr>
<td>Attention-Deficit/Hyperactivity Disorder (ADHD):</td>
<td>Impaired attention with bursts of hyperactivity/impulsivity</td>
<td>+</td>
</tr>
<tr>
<td>Specific Learning Disorder:</td>
<td>Difficulties with learning skills like reading, writing or spelling</td>
<td>Not possible</td>
</tr>
<tr>
<td>Motor Disorders</td>
<td>Impaired execution of coordinated motor skills, or repetitive motor behaviors</td>
<td>+</td>
</tr>
<tr>
<td>Tic Disorders</td>
<td>Habitual sudden, rapid, recurrent and non-rhythmic motor movements or vocalizations (including Tourette’s syndrome)</td>
<td>+</td>
</tr>
</tbody>
</table>

* While animals do not have language, their vocalizations (e.g., rodent USVs, bird songs or primate ‘calls’) may potentially be relevant to modeling neurobiological bases of communication
Table 2. Selected animal (rodent) models of neurodevelopmental disorders. ID – intellectual disabilities, ASD – Autism Spectrum Disorder, ADHD – Attention Deficit/Hyperactivity Disorder, SLD – Specific Learning Disorder, CD – communication disorders (see Table 1 for details).

<table>
<thead>
<tr>
<th>Rodent models</th>
<th>Availability in other model organisms</th>
<th>Clinical relevance</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Zebrafish</td>
<td>Chicks</td>
</tr>
<tr>
<td>Social/preference tests</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Social recognition</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Open field test</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Various memory tests</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Various attention tests</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Various impulsivity tests</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Self-grooming test</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aggression test</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Marble burying test</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ultrasonic vocalizations</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>
Table 3. Main drugs currently used to treat major neurodevelopmental disorders (NDDs). ASD – autism spectrum disorder, ADHD – attention deficit hyperactivity disorder

<table>
<thead>
<tr>
<th>Drugs</th>
<th>Main profile and mechanism of action</th>
<th>NDDs treated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Risperidone (Risperdal)</td>
<td>Atypical antipsychotic, blocks D2 and 5-HT2A receptors</td>
<td>Tics, ASD***</td>
</tr>
<tr>
<td>Haloperidol (Haldol)</td>
<td>Typical antipsychotic (neuroleptic), blocks D2 receptors</td>
<td>Tics*</td>
</tr>
<tr>
<td>Pimozide (Orap)</td>
<td>Typical antipsychotic (neuroleptic), blocks D2,D3 and D4 receptors</td>
<td>Tics*</td>
</tr>
<tr>
<td>Clonidine (Catapres)</td>
<td>Sympotolytic α2 adrenergic- and imidazole receptor agonist</td>
<td>Tics, ADHD*</td>
</tr>
<tr>
<td>Aripiprazole (Abilify)</td>
<td>Atypical antipsychotic, partial agonist at dopamine and 5-HT receptors</td>
<td>Tics, ASD***</td>
</tr>
<tr>
<td>Atomoxetine (Strattera)</td>
<td>Non-stimulant norepinephrine reuptake inhibitor</td>
<td>ADHD*</td>
</tr>
<tr>
<td>Methylphenidat (Ritalin)</td>
<td>Stimulant, dopamine-norepinephrine reuptake inhibitor</td>
<td>ADHD*</td>
</tr>
<tr>
<td>Lisdexamfetamine (Vyvanse)</td>
<td>Stimulant prodrug, blocks monoamine uptake</td>
<td>ADHD*</td>
</tr>
<tr>
<td>Olanzapine (Zyprexa)</td>
<td>Binds to α1, dopamine, histamine H-1, muscarinic, and 5-HT2 receptors</td>
<td>ASD</td>
</tr>
<tr>
<td>Ziprasidone (Geodon)</td>
<td>Blocks dopamine-serotonin receptors and monoamine reuptake</td>
<td>ASD</td>
</tr>
<tr>
<td>Adderall***</td>
<td>Stimulant amphetamines, block dopamine-norepinephrine reuptake</td>
<td>ADHD*</td>
</tr>
<tr>
<td>Buspirone (Buspar)</td>
<td>Anxiolytic, serotonin 5-HT1A receptor partial agonist</td>
<td>ADHD, ASD</td>
</tr>
<tr>
<td>Naltrexone</td>
<td>Antagonist of μ-opioid receptor</td>
<td>ASD</td>
</tr>
<tr>
<td>SSRIs****</td>
<td>Antidepressants, selective serotonin reuptake inhibitors</td>
<td>ASD, ADHD</td>
</tr>
</tbody>
</table>

* US Federal Drug Administration (FDA)-approved drugs
** Mostly used to treat aggression associated with ASD
*** A combination of amphetamine and dextroamphetamine
**** Mostly used to treat some (e.g., aggression, affective deficits) aspects of NDDs, but can trigger neurodevelopmental deficits in off-spring
**Table 4. Examples of various tests to characterize neural development and behavior of young and adult rodents.**

<table>
<thead>
<tr>
<th>Category</th>
<th>Tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>Simple reflexes</td>
<td>Sensorimotor (pupillary, salivation, lacrimation) reflexes, acoustic startle*, limb grasping, somatomotor (loss of crossed-extensor reflex and rooting response, vibrissae response)</td>
</tr>
<tr>
<td>Sensory and motor</td>
<td>Cliff avoidance*, olfactory discrimination*, righting, gait analysis, rotarod, grip strength test, nest building, rope climbing test, vertical screen, self-grooming analyses and the open field test*</td>
</tr>
<tr>
<td>Attention</td>
<td>Various attention tests*</td>
</tr>
<tr>
<td>Cognitive</td>
<td>Olfactory conditioning*, T-maze*, Morris water maze, passive and active avoidance tests*, operant conditioning schedules*, homing and object recognition tests*</td>
</tr>
<tr>
<td>Social</td>
<td>Ultrasonic vocalization, social interaction* and social preference tests*</td>
</tr>
<tr>
<td>Emotionality</td>
<td>Elevated plus maze, light-dark box*, social interaction test*, self-grooming analysis</td>
</tr>
</tbody>
</table>

* Conceptually similar tests are also available for aquatic (zebrafish) models, illustrating evolutionarily conserved trains across species
Table 5. Summary of selected additional strategies to improve drug development to treat neurodevelopmental pathogenesis

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Innovate drug selection and screening</td>
<td>Use a wider spectrum of compounds, especially those beyond traditional mechanisms of anti-NDD therapy</td>
</tr>
<tr>
<td>Develop disorder-specific drugs</td>
<td>Drugs targeting ASD vs ADHD or depression</td>
</tr>
<tr>
<td>Develop sex- and age-specific drugs</td>
<td>For different disorder subtypes and cohorts</td>
</tr>
<tr>
<td>Parallel behavioral changes with electrophysiological biomarkers</td>
<td>Use neuroimaging approaches (e.g., functional magnetic resonance imaging or mismatch negativity analyses) sensitive to NDDs*</td>
</tr>
<tr>
<td>Focus on additional disorders and their comorbidity</td>
<td>E.g., with PTSD, aggression, anxiety and psychoses</td>
</tr>
<tr>
<td>Focus on ‘adult’ developmental disorders beyond DSM-5</td>
<td>E.g., child and adult PTSD and depression</td>
</tr>
<tr>
<td>Uncover what NDD phenotypes are shaped by neurodevelopmental vs. neurochemical changes</td>
<td>Use of conditional transgenic animals and/or prenatal environmental factors</td>
</tr>
<tr>
<td>Establish developmental timelines of phenotypes preceding the full-blown onset of NDDs</td>
<td>Apply longitudinal studies</td>
</tr>
<tr>
<td>Examine gene-environment correlations</td>
<td>Refine analysis of mother- and father-offspring, as well as early ‘peer-peer’ social interactions</td>
</tr>
<tr>
<td>Assess developmental genomic responses in CNS in a region-specific manner</td>
<td>Use anatomically comprehensive databases of the developing human brain, including in situ hybridization, and microarray analyses**</td>
</tr>
<tr>
<td>Include more research and evidence generated by complementary alternative medicine (CAM)</td>
<td>Examine anti-NDD potential of nutrients, natural products and plant extracts, including those used in traditional Chinese medicine, Indian Ayurveda and Native/Latin-American or African folk medicine.</td>
</tr>
</tbody>
</table>

*See, for example, 80, 81, 101-102 for details.

** See 102-105 for details of NDD-associated susceptibility genes’ upregulation in the prefrontal cortex, and prenatal abundance of NDD-related (e.g., ASD-related) gene expression.
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