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A framework for the investigation of rare genetic disorders in neuropsychiatry

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Denovo and inherited rare genetic disorders (RGDs) are a major cause of human morbidity, frequently involving neuropsychiatric symptoms. Recent advances in genomic technologies and data sharing have revolutionized the identification and diagnosis of RGDs, presenting an opportunity to elucidate the mechanisms underlying neuropsychiatric disorders by investigating the pathophysiology of high-penetrance genetic risk factors. Here we seek out the best path forward for achieving these goals. We think future research will require consistent approaches across multiple RGDs and developmental stages, involving both the characterization of shared neuropsychiatric dimensions in humans and the identification of neurobiological commonalities in model systems. A coordinated and concerted effort across patients, families, researchers, clinicians and institutions, including rapid and broad sharing of data, is now needed to translate these discoveries into urgently needed therapies.

GDs are defined as genetic disorders affecting fewer than 200,000 individuals in the United States¹, or 1 in 2,000 individuals; however, a few highly penetrant genetic variants that occur more frequently (such as trisomy 21, affecting 1 in 600 individuals²) are often included. RGDs can be caused by the alteration of single nucleotides in one gene, entire chromosomes with hundreds of genes, or anything in between (Fig. 1a). Though they are individually rare, the contribution of RGDs to human morbidity en masse is substantial, with 18% of protein-coding genes currently implicated in such disorders (Supplementary Table 1). RGDs affect the central nervous system (CNS) disproportionately, with CNS symptoms documented in 74%³ (Fig. 1b); neuropsychiatric manifestations account for a substantial fraction of this CNS morbidity, with RGDs diagnosed in at least 40% of cases of developmental delay⁴ and up to 20% of cases of autism spectrum disorder (ASD)⁵.

Some RGDs have near-complete penetrance and distinctive features that enable syndromes to be described, affected and unaffected family members to be reliably distinguished, and the underlying genetic locus identified (as for Dravet syndrome, an RGD caused by SCN1A mutations resulting in a severe form of epilepsy^{6,7}). Newer genomic technologies, including chromosomal microarray, wholeexome sequencing (WES) and whole-genome sequencing (WGS) (see Box 1, Glossary), have identified many additional RGDs with less distinctive features, lower penetrance and/or more variable effects on phenotype. Many of these newly described RGDs have milder phenotypic impacts that may be largely brain specific (for example, 15q13.3 deletions increase risk for psychosis 18-fold8; Supplementary Table 2), in contrast to those with large effect sizes and multisystem involvement (for example, in cases of trisomy 21, intellectual disability (ID), global developmental delay and short stature are nearly universal⁹). Genomic technologies have also expanded the known manifestations of previously recognized RGDs (for example, many individuals with SCN2A mutations present with ASD rather than seizures10,11).

The full extent to which RGDs contribute to the manifestations of behavioral aberration, evident in several neuropsychiatric disorders,

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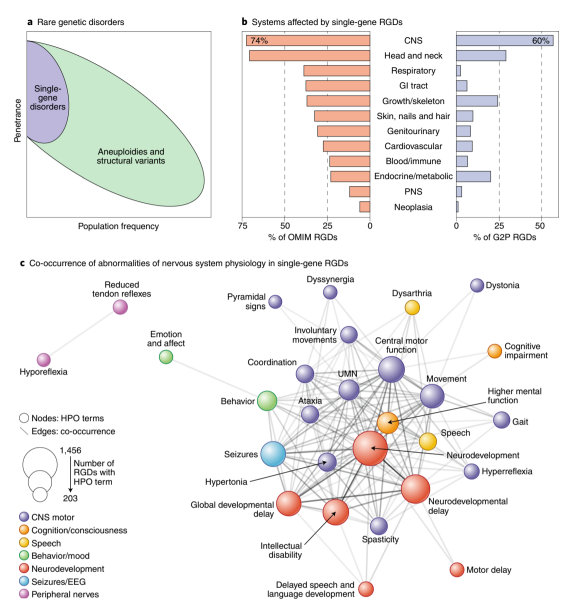


Fig. 1 Overview of rare genetic disorders (RGDs). **a**, RGDs may be caused by variants that affect one gene (purple) or many genes (green). Many aneuploidies and structural variants arise spontaneously at higher rates than single-gene disorders, leading to comparatively high population frequencies for a given penetrance^{33,124}. **b**, The CNS is involved in the majority of single-gene RGDs. **c**, Single-gene RGDs frequently affect multiple neuropsychiatric domains, as shown by extensive co-occurrence of Human Phenotype Ontology terms (Supplementary Table 1)¹²⁵. Terms that co-occur in at least 200 RGDs are shown as nodes (colored circles, size determined by the number of RGDs), with edge weight (gray lines) determined by the degree of co-occurrence of a term between RGDs (203–1,114). Network layout is based on the Compound Spring Embedder algorithm¹²⁶. OMIM, Online Mendelian Inheritance in Man; G2P, Gene2Phenotype; HPO, Human Phenotype Ontology; CNS, central nervous system; PNS, peripheral nervous system; UMN, upper motor neuron. Credit: Debbie Maizels/Springer Nature.

is still emerging. RGDs appear to contribute significantly to the genetic architecture of neuropsychiatric disorders characterized by high heritability, early age of onset, reduced fecundity, impaired cognition and behavioral deficits^{12–14}. In pediatric patients referred for developmental delay, ID and/or ASD, the diagnostic yield of clinical genetic testing (microarray and/or whole-exome sequencing) is more than 30%^{15–19}. It is important to note, however, that individuals with ASD with comorbid ID show a significantly higher rate of rare, de novo damaging variants than those with normal IQ, whereas there appears to be a greater contribution from common variants in those with higher IQ^{5,20,21}. Therefore, RGDs contribute significantly to neurodevelopmental disorders, such as ID, ASD and epilepsy^{2,2,23}; contribute modestly to child- or adolescent-onset neuropsychiatric disorders (attention deficit hyperactivity disorder

(ADHD)²⁴, schizophrenia^{8,25} and Tourette's syndrome²⁶); and contribute less to later-onset neuropsychiatric disorders (bipolar disorder and major depressive disorder^{8,14,27,28}).

Understanding the contribution of RGDs to neuropsychiatric disorders is critical for patient care and for developing effective therapeutics. Diagnosing and understanding RGDs enables screening for specific risks, early detection, informed family planning, initiation of services and therapies, detailed prognosis and support. Notably, recent advances in genetically targeted therapies^{29,30} raise the possibility of treating the underlying pathology. At the same time, the importance of this research is not limited to those diagnosed with RGDs. These high-penetrance disorders also have the potential to provide key insights into the underlying biology of idiopathic neuropsychiatric disorders.

Box 1 | Glossary

Single-nucleotide variant (SNV) A genetic variant in which one nucleotide (for example, C) is changed to another (for example, T). Single-nucleotide polymorphism (SNP) A common SNV (for example, found in $\geq 1\%$ of a population).

Insertion or deletion (indel). The gain or loss of 1–50 nucleotides, usually detected by sequencing.

Missense variant. An SNV in a gene that changes one amino acid of the resulting protein.

Protein-truncating variant/premature termination variant (**PTV**). An SNV or indel that disrupts one copy of a gene, resulting in a premature stop codon that is expected to elicit nonsensemediated decay so that no protein is formed.

Loss-of-function (**LoF**) **variant**. Genetic variants predicted to seriously disrupt the function of protein-coding genes, for example, a missense variant at an amino acid critical to protein function or a PTV.

pLI (probability loss-of-function intolerant) score. Probability that a given gene is intolerant of PTVs based on lower-than-expected rates of PTVs in the general population, suggestive of selective pressure. Genes with high pLI scores (\geq 0.9) are extremely LoF intolerant, whereas genes with low pLI scores (\leq 0.1) may be LoF tolerant or there may be insufficient data to assess their tolerance.

Copy-number variant (CNV). The gain (duplication) or loss (deletion) of \geq 50 nucleotides (previously \geq 1,000 bp), but often thousands to millions of nucleotides. Can be detected by microarray (large CNVs only) or WGS.

Structural variant (SV). A large-scale rearrangement of DNA. Can include CNVs, but also inversions, translocations, or more complex rearrangements.

Germline de novo variant. A new genetic variant observed in a child but not in either parent.

Somatic de novo variant. A new genetic variant observed in some cells of an individual but not others.

Chromosomal microarray (CMA). Technology that enables the number of copies of DNA to be assessed at thousands of locations across the genome, enabling the detection of CNVs. Many SNP genotyping microarrays detect SNPs at these locations too.

Whole-exome sequencing (WES). Technology that assesses ~45 million individual nucleotides in regions of DNA that encode proteins (exons), enabling the detection of SNVs and indels.

Whole-genome sequencing (WGS). Technology that assesses ∼3.2 billion individual nucleotides across the genome, enabling the detection of SNVs, indels, CNVs and other SVs.

Locus (plural: loci). A region of DNA ranging from a single nucleotide to an entire chromosome. For RGDs, it often describes a single gene or structural variant.

Mendelian disorder. A disorder that follows a Mendelian pattern of inheritance. Changes at several loci may elicit the same disorder. Complex disorder. A disorder caused by multiple variants with a range of effect sizes at multiple loci, often in combination with environmental factors.

Neuropsychiatric. Describes a behavioral or emotional disturbance due to an abnormality in the structure or function of the central nervous system.

Neuropsychiatric domain. A psychological construct relevant to human behavior and mental disorders that can be measured along a continuum in health and disease (e.g., anxiety).

Autosomes. Chromosomes 1–22 (in humans), in contrast to the sex chromosomes (X, Y) or mitochondria (M).

Autosomal dominant (AD). Describes a pattern of Mendelian inheritance in which a single variant on one copy of an autosomal chromosome causes the disorder or trait.

Autosomal recessive (AR). Describes a pattern of Mendelian inheritance in which two copies of a variant, one on each copy of an autosomal chromosome, must be present in order to cause the disorder or trait.

Compound heterozygous. A pattern of Mendelian inheritance in which two different variants at the same locus, one on each copy of a chromosome, cause the disorder or trait.

X-linked recessive (XLR). A pattern of Mendelian inheritance in which one variant on chromosome X causes the disorder in males only. It is not uncommon for carrier females to manifest mild or tissue-localized symptoms of disease due to normal or skewed X-inactivation of the functional copy of the gene in individual cells.

A US National Institute of Mental Health (NIMH) workshop was held in September 2017 with the goal of identifying strategies for advancing the understanding and management of neuropsychiatric disorders through a focus on RGDs (see Supplementary Table 3 for list of attendees). This Perspective outlines guidelines and recommendations that emerged from this workshop. In particular, three priorities were identified (see Box 2).

The overlap between rare genetic disorders and psychiatric disorders

Individuals affected by RGDs frequently have multiple neuropsychiatric symptoms, and specific RGDs frequently lead to multiple neuropsychiatric symptoms (grouped into neuropsychiatric domains) in different individuals (Fig. 1c). The co-occurrence of neuropsychiatric phenotypes within single-gene RGDs (Supplementary Table 1) reveals a highly connected network between developmental delay, behavioral abnormalities, seizures and abnormalities of the motor system (Fig. 1c).

Specific RGDs occupy different regions of this co-occurrence network, leading to widespread variation between RGDs in the domains affected (Fig. 2a,b). Notably, some RGDs appear to have strong effects on specific neuropsychiatric domains, as exemplified by the link between the RGD 22q11.2 deletion syndrome and

psychosis^{8,31} and the putative protective effect of the reciprocal 22q11.2 duplication against psychosis³² (Fig. 2b, Extended Data Fig. 1, Supplementary Tables 2 and 4). In contrast, the association between the frequency of RGD diagnosis and IQ in ASD suggests that there is a global effect on cognition across many RGDs (Fig. 2a), and this notion is supported by large-scale screens in the general population^{5,13,33,34}. In addition, recent evidence suggests that common genetic variation may influence the phenotypic expression of RGDs, and thus may contribute to variation in clinical presentation (Fig. 2a)³⁵. More data are required to more precisely address the contribution of these RGD-causing mutations to neuropsychiatric traits, ideally from cohorts with consistent genotype and phenotype data ascertained without bias toward selection of individuals with particular characteristics, such as population-based birth cohorts, or with a proper strategy to control for ascertainment bias, such as assessing the same clinically presenting disorder across multiple RGDs and/or using first-degree relatives as controls34,36 to account for additional genetic and environmental factors (see Fig. 2c,d).

Further confounding our understanding is that the phenotypic severity within each RGD varies widely (Fig. 2a). Presently, our knowledge of individuals with RGDs is biased toward those who are clinically identified and thus likely more severely affected, with milder cases (Fig. 2d) and atypical presentations under-represented ^{12,13}.

At the same time, studies of brain, cognition and behavior in RGDs (i.e., phenotyping studies) often have the opposite bias, assessing only high-functioning individuals. This ascertainment bias complicates genetic counseling, as it is not yet possible to provide an accurate representation of prognosis.

To address Priority 1 (Box 2), we think that proper characterization of RGD effects on brain-related phenotypes requires moving away from categorical diagnoses (such as ASD) to neuropsychiatric domains assessed as quantitative traits (such as social impairment)37, as advocated for by the NIMH Research Domain Criteria (RDoC) initiative^{38,39}, and assessing traits consistently across carriers who were referred to the clinic (probands) and carriers who were not (family members, individuals from unselected populations) regardless of presenting diagnosis (Fig. 2d)36,40,41. Capturing the full spectrum of phenotypic severity and controlling for ascertainment bias that often leads to inflated estimates of severity (Fig. 2d)⁴² may require conducting population-level analyses of hundreds of thousands of individuals, for example, using birth cohorts or health system registries 13,33,34,43, and applying this experimental design across generations to capture early- and lateonset manifestations^{34,36,44,45}

Prioritizing RGDs for research

We think that the allocation of resources for research into an RGD should be commensurate with the level of evidence for genetic association to a specific neuropsychiatric domain. In order for resources to be provided for research, genome-wide statistical association should generally be demonstrated for a neuropsychiatric domain^{4,46,47} in a defined cohort^{4,5,8,23,48–50}; however, for very rare neuropsychiatric disorders (such as childhood-onset schizophrenia), a lower level of evidence may be appropriate⁵¹. Further, in order to identify supporting mechanistic insights, we assert that it is important that functional data are used to enhance these statistical models (for example, protein-truncating variant (PTV) as opposed to missense mutation data, pLI score; see Box 1, Glossary)⁵². For genes for which clear functional assays are available, systematic evaluation of observed variants in both cases and controls11,53 may also contribute to statistical association. In the absence of clear biomarkers (see below) indicating a role for a genetic variant in the etiology of disease, observing downstream, nonspecific effects for individual variants, such as altered synaptic function or behavioral assays, is not a substitute for identifying a statistical association between the genetic variant and the human condition. For genetic loci containing multiple genes (such as CNVs), association evidence for a locus does not automatically imply that a single gene is driving the effect. For many recurrent CNVs, multiple genes with smaller individual effects appear to contribute to the overall risk (Supplementary Fig. 1)5,54. The additive contribution of polygenic, common risk should also be considered^{5,21,54,55}.

For the majority of RGDs associated with neuropsychiatric domains, the evidence for statistical association comes from the observation of multiple de novo mutations at the same gene or locus in independent cases^{4,5,8,23,48-50}. Statistical association is calculated from three metrics: (1) the number of independent cases (affected individuals) with de novo mutations at the gene or locus; (2) the denominator, i.e., the total number of cases assessed to find the mutations; and (3) the mutation rate, i.e. the chance of observing a similar de novo mutation in an unaffected individual. Mutation rate is highly dependent on gene size, with larger genes having higher mutation rates (Fig. 3a) and therefore requiring more de novo mutations in cases to demonstrate association (Fig. 3b)10,46,47. Defining these three metrics is straightforward in research cohorts^{4,52}; however, when considering multiple clinical reports⁵⁶, the denominator becomes all neuropsychiatric cases that have undergone genetic testing worldwide. Estimating this worldwide denominator as 100,000 at present, at least 5-10 confirmed and independent

Box 2 | Identified priorities for advancing neuropsychiatric research through a focus on RGDs

- Priority 1: Develop new strategies to consistently characterize the effects of RGDs on cognitive and behavioral traits that may result in disabilities and clinical referral, across the rapidly expanding spectrum of neuropsychiatric-associated RGDs routinely identified in the clinic.
- Priority 2: Prioritize RGDs for exploration of pathophysiology through bottom-up approaches: building experimental models starting from the lower levels of the mechanistic hierarchy, such as gene expression and synaptic function, will help guide investigation and explain phenomena higher up in the hierarchy, such as macroscopic brain and behavioral alterations leading to psychiatric diagnoses.
- Priority 3: Integrate the knowledge resulting from Priorities 1 and 2, above, to accelerate the development of new drugs and other therapeutic interventions in those with and without RGDs.

de novo PTV mutations are required to achieve genome-wide association, depending on gene size (Fig. 3b, Extended Data Fig. 2).

Once a genetic locus meets the threshold for genome-wide association, other factors should be considered in selecting which RGDs to study, including the effect size on behavioral domains, level of risk for a categorical disease manifestation, population frequency, tractability for experimentation and therapeutic potential. Given the complexity of neurobiology, there are important roles for both detailed assessment of individual genes and consistent assessments across multiple genes.

Progressing from genotype to mechanisms

The second priority we identified is the design of bottom-up approaches in order to progress from genetic etiology to mechanism. Because RGDs are genetically defined, they provide an unparalleled opportunity to understand the pathophysiology of psychiatric domains and conditions. However, making the link between a genetic variant and the underlying mechanism (Fig. 4a) for a psychiatric domain or disorder presents a significant challenge^{57,58}. Conceptually, the problem can be divided into two parts: first, to link a genetic variant to a biochemical and/or biological consequence in a specific type of neuron or glial cell, and second, to connect a functional change in a set of cells to a defect in a circuit in a specific anatomical region of the brain at a particular time in development. Although linking these diverse levels seems daunting, the prospects for success are greatly improved by the ongoing development of experimental and conceptual tools, from wet-bench to bioinformatic approaches^{58,59}.

To pinpoint the subcellular location of a specific gene implicated in a disorder, it is possible to use genetic tagging with fluorescent proteins and imaging using ultra-resolution microscopy in human neurons created from stem cells in vitro. The function of the gene can be determined by using CRISPR-Cas9 genome engineering tools to knock out or overexpress the gene and observing the cellular consequences using a combination of high-content microscopy, calcium and voltage imaging, high-throughput electrophysiology and RNA sequencing. Although these tools have traditionally not been amenable to the study of multiple variants, the development of automated microscopes and patch clamps that are used for high-throughput drug screening is rapidly changing this. To establish the time in development and the anatomical location that is affected by a particular genetic variant, a set of gene atlases of the developing human brain have been generated by the Allen Institute and

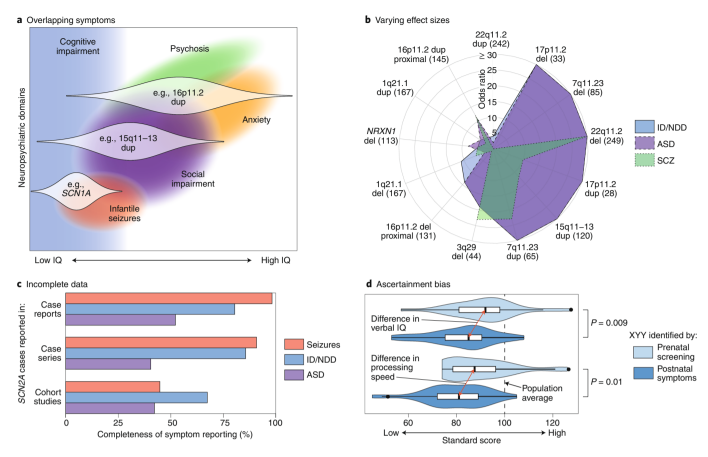


Fig. 2 | Cross-domain impact of RGDs and limitations of current evidence. a, A theoretical model of how three RGDs (16p11.2 dup, 15q11-13 dup and *SCN1A*) impact multiple neuropsychiatric domains across different individuals (distribution across affected individuals shown as white violin plots)^{34,127,128}. **b**, A polar plot showing the varying effect sizes (odds ratios) of different CNVs on the diagnosis of ID/NDD, ASD and SCZ (Supplementary Table 2, Extended Data Fig. 1). The number of CNV cases are shown in parentheses^{5,8,32,129-131}; the UK Biobank was used for controls³³. **c**, The completeness of symptom reporting for *SCN2A* mutations varies widely between publications¹³². Case reports describe a single *SCN2A* mutation in one case or family; case series describe multiple *SCN2A* cases; cohort studies describe hundreds of cases with the same disorder (for example, ID/NDD), some of which are found to have *SCN2A* mutations. This reporting bias, which is likely to be present for most RGDs, complicates comparisons across neuropsychiatric domains and between RGDs. **d**, The severity of symptoms in XYY aneuploidy varies between cases ascertained by prenatal screening (light blue) and those ascertained on the basis of clinical symptoms (dark blue)¹³³. This ascertainment bias, which is also likely to be present for most RGDs, also complicates cross-disorder comparisons and potentially inflates estimates of effect size and penetrance. ID, intellectual disability; NDD, neurodevelopmental delay; ASD, autism spectrum disorder; SCZ, schizophrenia; IE: infantile epilepsy; CNV, copy-number variant. Credit: Debbie Maizels/Springer Nature.

other groups that allow gene expression to be ascribed to specific types of cells in specific regions of the brain at particular times in development (http://portal.brain-map.org). Moreover, recent progress in recapitulating neural development in vitro using pluripotent stem cells offers a powerful platform to manipulate and investigate the roles of specific disease genes in human neurons and glial cells⁶⁰. This recent work demonstrates the high reproducibility and validity of these systems for studying many human brain developmental processes.

By looking for cell types that are enriched in the expression of specific genes, it has been possible to determine that the phenotypes associated with particular RGDs are caused by dysfunction in a specific type of cell. For example, phenotypes associated with mutations in the gene *SCN1A* that cause Dravet syndrome are due to defects in fast-spiking interneurons⁶¹, and glutamatergic neurons in the cerebral cortex are implicated in ASD^{52,62-64}. A similar strategy can be used to identify the developmental window during which a particular genetic variant leads to a disorder by narrowing down the expression of the gene to particular periods during development, as is the case for mutations in *FOXG1*, which likely lead to neurodevelopmental disorders by affecting the differentiation of cortical glutamatergic

neurons during brain development⁶⁵. Similar approaches have been applied to structural and functional neuroimaging data, testing multiple brain regions and circuits to identify those that are distinct versus shared across disorders^{66–68}, as well as neuroanatomic patterns that predict subsequent disease^{69–71}. Advances in data sharing have enabled the analysis of large-scale, multisite cohorts^{72–74}. Large-scale harmonized phenotypic data across RGDs could be analyzed in a similar manner and integrated with imaging and genomic data.

Identifying the circuits

The development of new, effective treatments is hindered by our current very limited understanding of the circuit abnormalities that underlie different neuropsychiatric disorders. RGDs offer a unique roadmap to the identification of these circuits because distinct underlying biochemical defects have now been shown to manifest with common phenotypes. For example, genes with roles in glutamatergic receptors (such as *GRIN2B*), sodium channel activity (such as *SCN2A*) and chromatin remodeling (such as *CHD2*) can all manifest both in the behaviors of ASD and in seizures^{5,23,52}. In all likelihood, these mechanistically distinct genetic causes of

disorders have common phenotypic manifestations through convergent effects on key circuits. The convergent effects of the mutations responsible for these circuit abnormalities therefore offer a unique set of probes to identify those circuits.

Although in vitro tools are essential for identifying the biochemical and cell biological consequences of genetic variants, identifying the neuronal circuits that lead to changes in behavior requires a more intact system. Well-defined and replicable genetic loci associated with RGDs provide a unique and indispensable opportunity to generate experimental animal models. These experimental models do not replicate the complexity of human neuropsychiatric disorders, but are essential as scientific tools to interrogate the effects of disorder-associated genes and variants on circuit properties⁷⁵. In fact, it is at this juncture of genetics and neurobiology that RGDs present an unmatched advantage to investigate pathophysiological mechanisms across multiple levels (Fig. 4a). Using a combination of recently developed technologies such as optogenetics, chemogenetics, trans-synaptic tracers and single-cell sequencing in both animal models and human cells⁷⁶⁻⁷⁹, we now have the ability to determine the effect of genetic variants on specific cells and networks with unprecedented detail.

Genetically engineered mouse lines can offer important insights into gene function that provide a basis for probing pathological mechanisms in RGDs. Although behavioral phenotypes of mouse models may not resemble human disorders, the molecular, cellular and circuit phenotypes are more likely to be informative. Crossspecies validation of animal models provides evidence for evolutionarily conserved electrophysiological signatures and cellular phenotypes that may enhance confidence in the relevance of the genetic perturbation (Fig. 4b–d)^{80–82}. Regardless of the approach, replicable and rigorously collected data are beneficial, including replication of key findings in a second cohort⁸³.

Nonhuman primate (NHP) experimental systems, such as are used in studying depression⁸⁴ and schizophrenia^{85–88}, offer several advantages, including structural and functional similarity of the prefrontal cortex to humans, presence of similar cortical–subcortical circuits and complex behavior repertoires⁸⁹. These benefits come at the expense of lower throughput and substantially greater costs and ethical concerns. As with other models, validation of resulting findings in humans will be essential.

Alongside animal models, advances in cell reprogramming and differentiation, as well as CRISPR-Cas9 methods, enable the study of human neurons and glial cells in specific RGDs^{90,91}. In particular, three-dimensional (3D) cultures, also known as organoids, resembling specific brain regions and in-vitro-assembled 3D cultures to model inter-regional cell-cell interactions⁹² hold promise in capturing previously inaccessible aspects of human development. Transplantation of human-derived neural cells into animals could reveal defects associated with circuit-wide integration in RGDs. Moving forward, further improvements (for example, cellular maturation, spatiotemporal control, scalability) and validation of human cellular models will be needed⁹⁰.

The discovery of biomarkers for patient selection for clinical trials, confirmation of target engagement (i.e., determining that the intervention has the predicted effect on its hypothesized mechanism of action) or prediction of treatment response that translates between species, including humans, would transform the utility of such experimental systems 81,82,93,94. Without such biomarkers, our best hope is to look for convergence across multiple RGDs in multiple experimental systems. Rather than advocating for one animal or cellular experimental system, we should encourage the development of multiple avenues of investigation for the RGDs that rank highest in the prioritization criteria (above) and for assays that can be compared across multiple systems (Fig. 4b–d)⁹⁵. In all animal or cellular experimental systems, genetic background can affect the results profoundly⁹⁶, and

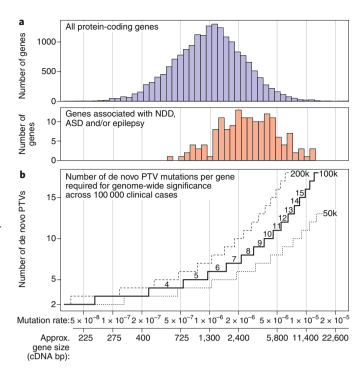


Fig. 3 | Thresholds for genome-wide significant association. a, Mutation rates vary across genes based primarily on gene size (cDNA) but also on sequence (for example, GC content)^{10,47}. Genes associated with neurodevelopmental delay, ASD and/or epilepsy to date tend to be large, with higher mutation rates^{4,23,52}. **b**, Tens of thousands of individuals with neuropsychiatric disorders have been sequenced to date. To estimate the number of independent de novo protein-truncating variants (PTVs) across multiple case reports that are required for reliable association of a gene with a neuropsychiatric disorder, we used a Poisson distribution. Expected mutation rates in 50,000, 100,000 and 200,000 individuals are estimated from controls across the range of gene sizes (see Supplementary Methods and Extended Data Fig. 2)^{5,47}. ASD, autism spectrum disorder; NDD, neurodevelopmental delay; GC, guanine-cytosine¹³⁴. Credit: Debbie Maizels/Springer Nature.

experiments that replicate results across multiple strains or clones should be encouraged 97.

Therapeutic development

Aside from some metabolic disorders 98, very few RGDs currently have treatments guided by genetic findings. Recent advances in genetic therapy, including the landmark improvement of infant mortality and neurological function in patients with spinal muscular atrophy (SMA) treated with the antisense oligonucleotide nusinersen²⁹ or Zolgensma gene replacement³⁰, offer grounds for cautious optimism. However, potential therapies have yet to modify cognitive or behavioral symptoms in humans in fragile X syndrome⁹⁹, tuberous sclerosis¹⁰⁰ or Rett syndrome¹⁰¹, despite promising results in mice¹⁰²⁻¹⁰⁴. Gene dosage is important in many such disorders, with too much or too little causing symptoms^{11,105}, presenting challenges in titrating therapy. Consequently, recessive loss-of-function disorders, such as SMA, may be easier to treat than dominant lossof-function disorders. Delivery of a therapeutic to the brain presents an additional challenge, further complicated if the therapy needs to be targeted to a specific cell type or brain region.

The extent to which neuropsychiatric symptoms are modifiable in humans varies by domain, age and RGD. Although some domains can be modified years after onset (for example, seizures, psychosis, attention), others may require early treatment, as suggested

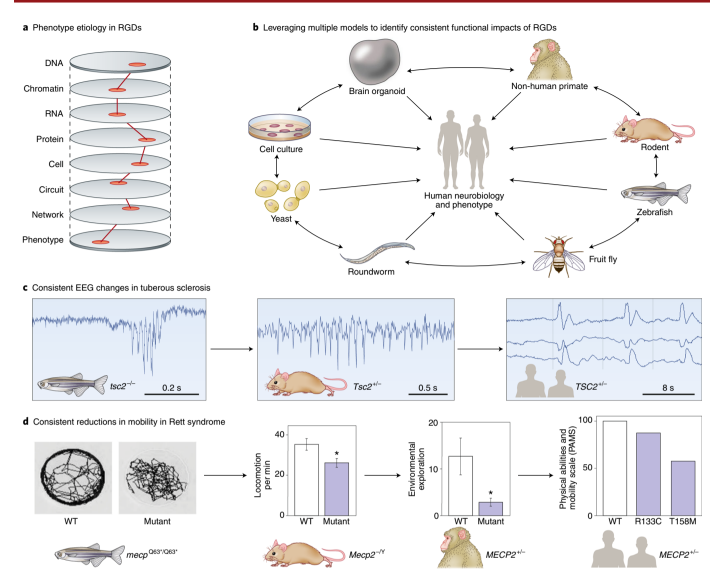


Fig. 4 | Functional assays across disorders and models. a, To understand the etiology of neuropsychiatric disorders, we need to identify the minimal 'causal path' by which the effects of the RGD lead to the phenotype, as shown by the hypothetical red line. Future therapeutics or biomarkers would be expected to interact with this causal path. **b**, No model experimental system perfectly recapitulates the human brain. By performing similar assays across multiple models, we can identify consistent consequences of RGDs, while leveraging the strengths of each model. These need to be related back to humans through similar assays or testing model predictions. **c**, Seizure activity is consistently observed in models of tuberous sclerosis (*TSC2*), though a homozygote model is used in zebrafish¹³⁵⁻¹³⁷. **d**, Mobility is consistently reduced in models of Rett syndrome (*MECP2*); as with *TSC2*, the specific genetic lesion assessed varies between models¹³⁸⁻¹⁴¹. Credit: Debbie Maizels/Springer Nature.

by the critical period of visual cortex development¹⁰⁶ or the relationship between early management of phenylalanine hydroxylase deficiency (PKU) and adult IQ⁹⁸. With this in mind, therapies that have failed in adults and adolescents are being tested in 2–6-year-olds with fragile X syndrome¹⁰⁷ and infants with tuberous sclerosis^{108,109}. Prenatal screening for many RGDs is feasible with current technology, and there is precedent for neonatal treatment for highly penetrant RGDs²⁹.

Clinical trials require predefined outcome measures. In neuropsychiatric disorders, this usually entails caregiver questionnaires regarding symptom severity and functioning. There is considerable need for more reliable, sensitive and objective measures. Measures of brain function that can be compared to model systems, such as EEG or functional MRI, should be included in trial protocols to help demonstrate target engagement by the therapy and refine dosing for future trials⁹⁴. Establishing multisite databases of indi-

viduals with RGDs, alongside genotype and longitudinal phenotype data, for example, by extracting knowledge from patient's clinical notes^{110,111}, would reduce the costs of trials and help identify subgroups with the greatest potential to respond. Finally, clinical trials need to be of sufficient size to yield clear positive or null results^{99,112,113}.

Phenotyping and harmonization

To date, most phenotypic descriptions have focused on individual RGDs, ranging from case series to quantitative assessments of large patient registries. Such studies provide vital, clinically relevant insights that can have important therapeutic implications^{114,115}. At the same time, the overlap among neuropsychiatric domains, within and between RGDs, provides the opportunity for cross-domain and cross-disorder analyses^{44,116}, which may identify biologically defined subcategories of neuropsychiatric domains¹¹⁷.

Achieving this vision will require a community-wide initiative to collate consistent, harmonized genetic data with quantitative measures of cognition and behavior across multiple RGDs and multiple neuropsychiatric domains. No single set of clinical data elements adequately describes all RGDs. The variables to collect in a natural history study should be broad and based on features of the disorder, including morbidities that are most important to patients, those that are most likely to be life limiting, potential prognostic characteristics and those that may help formulate sensitive clinical endpoints¹¹⁸. The resulting data repository should reflect existing National Institutes of Health (NIH) data-sharing policies and standards, and become part of the NIH data sharing ecosystem to secure access for researchers, clinicians and institutions, enabling a wide range of hypothesis-testing with no restrictions on data use and appropriate human subjects and privacy safeguards^{119,120}. This effort will also require standardized phenotypic ontologies, which can be achieved by adapting existing ontologies to ensure that adequate and comprehensive terminology exists for neuropsychiatric symptom manifestations across the lifespan.

Other stakeholder actions

The challenges and opportunities presented above require coordinated efforts among the community of stakeholders to fund, execute, coordinate and communicate research (see Supplementary Table 5). Patient- and family-based organizations fulfill an essential role of providing support and decreasing isolation experienced by patients and caregivers; these organizations also have a key role in forming interactive networks that connect them with researchers and health care systems (for example, to keep families informed about the latest clinical trials and research findings). Although some examples of interactive research and family-based networks for RGDs already exist, they are not all focused on or inclusive of neuropsychiatric disorders, and efforts need to be made to use these networks for such purposes. Such networks will contribute not only to larger sample sizes, which are critical for research progress, but also to organizing healthcare systems to collect consistent phenotypes that include neuropsychiatric measures to enhance data harmonization, as well as implementing standards for genotyping and phenotyping. Academic health systems, in partnership with other health care systems, can play a pivotal role in these networks by expanding the reach of their initiatives and imparting knowledge gained from the research. Some registries and databases will use a federated approach, and some will utilize a centralized repository approach. Both will be useful for specific purposes, with federated approaches allowing for more flexibility in individualization and 'real-time' accrual of data, and centralized repositories providing more digested, pre-harmonized data that require less work by the user before analysis, but also restricts usage. Both types of networks will clearly be useful, and will continue to be expanded upon with respect to data types included, with electronic health records (EHRs) quickly becoming an additional data source.

Recently, clinical notes from patient EHRs have been identified as important sources of clinical and demographic information. This has led to the development of a variety of knowledge-extraction tools that transform the raw text content of clinical notes into structured associations between patients and phenotypes defined in terminologies such as the Systematized Nomenclature of Medicine–Clinical Terms and the Human Phenotype Ontology. Knowledge extracted from clinical notes is of increased value to research into RGDs given the scarcity of patient data, necessitating the use of natural language processing engines to extract structured knowledge from clinical notes¹¹⁰. In addition, biobanks are increasingly being developed to support organized collections of biological specimens, genomic data and associated clinical information—including EHR data—from broadly consented, diverse patient populations contributing to RGD research^{121–123}.

Partnerships among patient and family organizations, academic sites and other health systems will provide an important platform for working with funding agencies to emphasize the need for the implementation of core, cross-diagnostic phenotypic measures and the use of broad neuropsychiatric measures to facilitate data harmonization. This approach will culminate in early collaboration and coordination across groups working in this space, resulting in the design of maximally effective initiatives, data sharing and dissemination of information and knowledge gained. Furthermore, partnerships will create a platform for collaboration with pharmaceutical companies, encouraging alliances in preclinical and translational research toward drug development. However, data harmonization for collaboration necessarily requires data sharing. The community should follow NIH and established standards in the field for phenomic, genomic and biosample sharing. Although this can be done retrospectively, optimal sharing will begin before data collection, and include sharing of variables and even study methodology, to ensure replication and maximal generalizability of findings.

Drug development programs require a strong scientific foundation, including detailed understanding of the natural history of a disorder. Because of the small numbers of patients affected, and because clinical experience is dispersed among clinical referral centers, the natural history of RGDs is rarely well characterized. The US Food and Drug Administration advises sponsors to evaluate the depth and quality of existing natural history knowledge early in drug development¹¹⁸. Close collaboration with patient and family groups and all stakeholders is essential to design and conduct impactful natural history studies for future potential drug design and approval.

Several examples (Supplementary Table 5) illustrate the potential contributions of such convergent efforts. The benefits of coordination across stakeholders are immense, and could create resources enabling investigators to merge and integrate available databases, examine existing data across disorders, and define the best data formats for prospective studies that use consistent broad and deep phenotyping to facilitate harmonization efforts. Furthermore, it fosters complementary approaches for building resources and establishing cohorts (for example, mining of electronic medical records, clinical recruitment, patient advocacy) with synergism that propels the field.

Conclusions

We think that RGDs hold important biological clues into the mechanisms underlying complex neuropsychiatric disorders, such as ASD and schizophrenia, that can be leveraged in model experimental systems, with the view to understanding the etiology underlying the full spectrum of these neuropsychiatric conditions. Although the individual features of each RGD are important, it is the commonalities across RGDs that hold the greatest promise to transform our understanding and management of neuropsychiatric disorders. Identifying these commonalities in phenotype and neurobiology will require coordination and sharing of methods, data and resources across individuals and institutions, following the lead of the NIH and genomics community. Although RGDs and the neuropsychiatric domains they impact pose substantial challenges, the opportunity these rare disorders offer for illuminating mechanisms is likely to be transformative for scientists, clinicians and, ultimately, patients.

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Competing interests

The authors declare no competing interests.

Additional information

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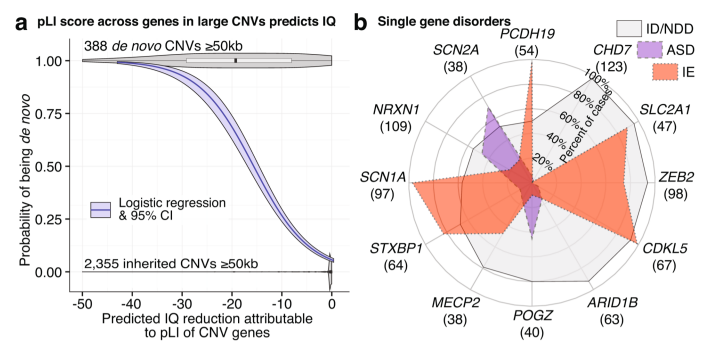
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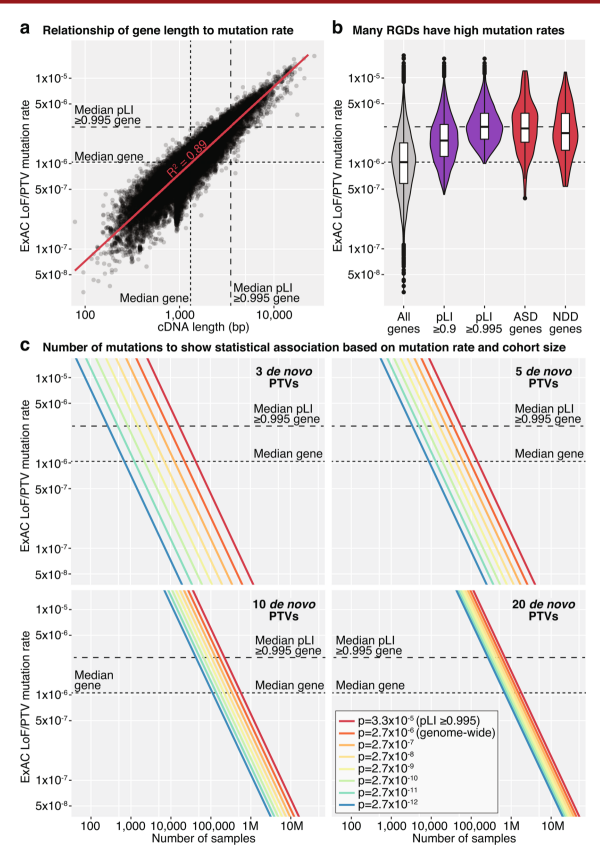
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Extended Data Fig. 1 | Impact of RGDs on neuropsychiatric domains. a, Many RGDs impact cognition, measured by IQ. For CNVs, the decrease in IQ (x axis) can be predicted by considering the pLI score of the genes within the CNV. CNVs that are predicted to markedly reduce IQ are more likely to be de novo (y axis), based on logistic regression (blue line) of 2,743 CNVs detected in patients with neurodevelopmental disorders and the general population (gray distributions at top and bottom). Updated analysis from ref. ⁵⁴. b, In Fig. 2, we show the odds ratio for ID/NDD, ASD and SCZ across different CNV loci. Here, we show an equivalent plot for single-gene RGDs. Insufficient control data exist to estimate odds ratio, and therefore we show the percentage of cases with ID/NDD, ASD, and IE based on curated publication review applied equally across genes (https://dbd.geisingeradmi.org) with the number of cases are shown in parentheses (see Supplementary Table 2 for numbers). Abbreviations: ID, intellectual disability; NDD, neurodevelopmental delay; ASD, autism spectrum disorder; SCZ, schizophrenia; IE, infantile epilepsy; pLI, probability loss-of-function intolerant.



Extended Data Fig. 2 | Thresholds for genome-wide significant association with de novo PTVs. a, Gene mutability is a function of gene length (cDNA) and sequence context (particularly GC content). **b**, RGD gene discovery from exome sequencing has been driven by de novo mutations, leading to a bias towards larger genes with higher mutability. **c**, Thresholds of statistical association (colored lines) are estimated for a given number of de novo PTV mutations (3, 5, 10, and 20) as cohort size (*x* axis) and gene mutability/size (*y* axis) varies. *P* values are estimated based on the rate of de novo PTV mutations in controls⁴ and a Poisson distribution (see Methods for details). Abbreviations: pLI, probability of loss-of-function intolerance; ASD, autism spectrum disorder; DDD: Deciphering Developmental Disorders; GC content, guanine-cytosine content.