Open questions: What has genetics told us about autism spectrum disorders?

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Abstract

Some of the most interesting questions in biology today, in my view, derive from the real advances in neuropsychiatry that have come largely from human genetics. Research in autism spectrum disorders (ASDs) has been leading the way, mainly because it has become especially well funded and has recently attracted many outstanding scientists. (I must make it clear that I am an outsider in this field, as I have never worked on any neuropsychiatric disorder).

Autism spectrum disorder genetics

More than 25 mutations that have a large effect on risk have now been strongly implicated in ASDs. Some are inherited, whereas others are de novo [1-5]. All of them are rare: no one of the affected genes appears to be involved in more than 1% of ASDs. Interestingly, so far, none of them has been found to be specific for ASDs, in that most have also been implicated in other neuropsychiatric disorders, including, for example, intellectual disability, schizophrenia and epilepsy. Even in the same family, the same large-effect mutations can contribute to different neuropsychiatric disorders. There thus seems to be a deep genetic connection between these disorders that is not understood. One possibility is that they all reflect synaptic dysfunction: there is increasing evidence, for example, that mutations that affect genes that encode proteins that work only or primarily at synapses can greatly increase the risk of ASDs and of other neuropsychiatric disorders [5].

From analyses of copy number variants (CNVs) and exome sequencing, it is now estimated that mutations in hundreds of genes can substantially increase the risk of developing ASD, the predisposing mutant genes varying from one individual to another [1-5]. Thus, ASDs are genetically heterogeneous, and their well known clinical heterogeneity is presumably at least in part a reflection of this genetic heterogeneity. There is also evidence that more than one rare, large-effect mutation can contribute to ASD in a single individual. So although ASDs are common (approximately 1% prevalence), the genetic underpinning in each individual is rare, and may even be unique, much as an individual cancer is genetically unique. All of this is remarkable progress, given that, until recently, very little was known about ASD genetics.

Reversibility in monogenetic animal models

Once a large-effect mutation is identified as a risk factor for a neuropsychiatric disorder, one can make a genetic animal model of it, in which the underlying neurobiological basis of the disorder can be studied. Such approaches have already yielded important insights into the nature of ASDs. The studies on mouse models of the monogenetic, or syndromic, forms of these disorders - such as Rett and fragile X syndromes - have been particularly informative. These single-gene conditions are neurologically complex, and the core features of autism occur in about half of the affected individuals. The most dramatic and surprising finding has been that both conditions can be largely reversed in adult mice carrying the mutations: if the brain problem is corrected either genetically (in Rett, where the mutation is in MeCP2) [6] or with drugs that correct an abnormal downstream signaling pathway (in fragile X, where the mutant gene is FMR1) [7], most of the behavioral and physiological abnormalities go away. This was totally unexpected, as these are developmental brain disorders: but these remarkable results demonstrate that the serious neurological defects seen in these mutant mice reflect reversible functional defects in the adult brain. This is potentially very good news for the future of drug therapies for humans with ASDs (and probably for some other neuropsychiatric disorders as well), although it is still uncertain whether ASDs in humans are similarly reversible in adults.

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If neuropsychiatric disorders are primarily the result of synaptic dysfunction, however, reversibility should not be so surprising: synaptic plasticity is, after all, presumably the basis for all of normal brain function that depends on memory, and it persists well into adulthood, which is why even as we age we can still learn and remember - at least some things. Moreover, some human neuropsychiatric disorders spontaneously show dramatic, if temporary, reversibility. Bipolar disorder is the clearest example: in this condition, individuals can switch from extreme mania to deep depression and then back to apparent normality, and can do this repeatedly. It is clear that the problem in bipolar disorder is an abnormality in mood control, but we don’t know much about the normal mood control system, let alone how it goes wrong in bipolar disorder. The identification of large-effect mutations that greatly increase the risk of this disorder, however, could lead to useful animal models, which might then shed light on both the underlying brain abnormality in bipolar individuals and the normal mood control system.

If there is a synaptic problem in ASDs, what is it?
If synaptic dysfunction is at the heart of ASDs, what exactly is the synaptic problem? Given the clinical and genetic heterogeneity of ASDs, there is unlikely to be one answer. There is evidence that there are defects in the ratio of excitatory to inhibitory synaptic transmission in some mouse ASD models, but it seems that the ratio can be abnormally increased or decreased, and the abnormality can vary from one brain region to another, even in the same mouse model [8].

Even if a consistent synaptic defect were found, it would be a major challenge to relate it to a particular behavioral phenotype. And how would you know whether the defect is a primary result of the genetic defect or a secondary effect of the brain’s attempt to compensate? After all, the nervous system is a specialist in compensatory adjustments.

Are abnormal genes the whole story?
Although genetic abnormalities are likely to play an important part in most ASDs, they are unlikely to be the only contributor. If one identical twin develops an ASD, there is a 70 to 90% chance that the other will too - but it’s not 100%. Why not? The answer isn’t known, but the acquisition of somatic mutations in the developing brain could, in principle, be part of the answer.

Could environmental influences play a part? Although there is no compelling evidence that adverse postnatal environmental factors contribute to the risk of ASDs, there is strong evidence that stressful conditions in utero can do so: maternal infection early in pregnancy, for example, can greatly increase the risk of ASDs (and schizophrenia) [9]. It remains a mystery how such stresses in utero interact with predisposition genes in the fetus.

Why the ASD ‘core’?
ASD individuals have two defining characteristics: impaired social communication and interactions; and restricted and obsessive interests and behaviors. Together, these constitute the autistic ‘core’. In addition, most individuals on the autistic spectrum have a number of other neurological and psychiatric problems, which can include intellectual disability, language impairment, sensory and motor abnormalities, and so on.

A central mystery in ASDs is why mutations in so many genes result in the core features of ASD. It is not obvious why these features should occur together. What connects them? There are various possible explanations. One possibility is that the primary problem is the strong drive for sameness and predictability (which is an important part of the restricted interests and obsessive behaviors phenotype), and this leads to a secondary social withdrawal; after all, people are the least predictable objects in a developing child’s environment. Another possibility is that the primary problem is an electrically ‘noisy’ brain, which leads to a secondary social withdrawal and restricted interests - to keep the noise level to a minimum.

Although many mysteries remain, considering the little that was known up to a decade or so ago, there has been staggering recent progress, which is bringing the neurobiology of ASDs and neuropsychiatric disorders out of the darkness into the light. Much of what has been learned so far is encouraging, and, as more high-risk genes are discovered, progress can only accelerate.

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