A common form of the hyperkinetic movement disorder dystonia is caused by mutations in the gene \textit{TOR1A} (located within the \textit{DYT1} locus), which encodes the ATPase torsinA. The underlying neurobiological mechanisms that result in dystonia are poorly understood, and progress in the field has been hampered by the absence of a dystonia-like phenotype in animal models with genetic modification of \textit{Tor1a}. In this issue of the \textit{JCI}, Liang et al. establish the first animal model with a dystonic motor phenotype and link torsinA hypofunction to the development of early neuropathological changes in distinct sensorimotor regions. The findings of this study will likely play an important role in elucidating the neural substrate for dystonia and should stimulate systematic neuropathological and imaging studies in carriers of \textit{TOR1A} mutations.
will likely be required to definitively elucidate how P2Y14 mediates the HSPC stress response (10, 22). If P2Y14-deficient HSCs within the animal model system developed by Cho, Yusuf, and colleagues do indeed prove to be functionally senescent, then this could be an exciting model system for studying the onset of stress-induced senescence within the HSC compartment. A better understanding of the regulators of HSC stress response has important implications for regenerative medicine.

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D.J. Rossi is a New York Stem Cell Foundation Robertson Investigator.

Address correspondence to: Derrick J. Rossi, 200 Longwood Ave., Warren Alpert Building, Room #149e, Boston, Massachusetts 02115, USA. Phone: 617.713.8900; Fax: 617.713.8910; E-mail: derrick.rossi@childrens.harvard.edu.


Twisting mice move the dystonia field forward

Ása Petersén1 and Deniz Kirik2

1Translational Neuroendocrine Research Unit and 2Brain Repair and Imaging in Neural Systems (B.R.A.I.N.S.) Unit, Department of Experimental Medical Sciences, Lund University, Lund, Sweden.

A common form of the hyperkinetic movement disorder dystonia is caused by mutations in the gene TOR1A (located within the DYT1 locus), which encodes the ATPase torsinA. The underlying neurobiological mechanisms that result in dystonia are poorly understood, and progress in the field has been hampered by the absence of a dystonia-like phenotype in animal models with genetic modification of Tor1a. In this issue of the JCI, Liang et al. establish the first animal model with a dystonic motor phenotype and link torsinA hypofunction to the development of early neurodevelopmental changes in distinct sensorimotor regions. The findings of this study will likely play an important role in elucidating the neural substrate for dystonia and should stimulate systematic neurodevelopmental and imaging studies in carriers of TORA mutations.

Neurological disorders and the need for animal models

For many brain disorders, identification and characterization of the underlying neurobiological mechanisms remains a challenge for clinicians and scientists. Lack of defined neural substrates and an understanding of the pathways responsible for neurological and psychiatric symptoms has limited the development of novel therapies, which are urgently needed to improve the care and quality of life of affected individuals. In this issue, Liang and collaborators present animal models that recapitulate the major clinical symptomatology of dystonia (1). The study by Liang and colleagues represents an important leap forward for the dystonia research field.

Dystonia: the twists and turns

Dystonia is characterized by sustained or intermittent muscle contractions that cause abnormal, often repetitive, twisting movements and postures and is now recognized as a heterogeneous group of hyperkinetic movement disorders. The term dystonia was coined in 1911 by Herman Oppenheim, who used “dystonia musculorum deformans” to describe a childhood-onset form of generalized dystonia (2). These disorders have traditionally been classified as either primary or secondary dystonias. Primary dystonia is considered to only present with tremor or myoclonus as an additional neurological symptom.
in the absence of any neuropathological changes, whereas secondary dystonia is considered the consequence of a hereditary neurodegenerative disorder or an insult to the brain, and other neurological symptoms can be manifest. More recently, the dystonia classification system has been called into question. In particular, the division into primary and secondary dystonia has been criticized as neuroimaging and neuropathological studies have begun to indicate the presence of structural changes in primary dystonia (3). The recent international consensus classification has instead proposed to use a two-axes classification based on clinical characteristics and etiology (4).

There is a considerable genetic contribution for many forms of dystonia. The first dystonia locus (DYT1) was identified in 1990 and localized to chromosome 9 (5). The gene, which was identified seven years later and named TOR1A, encodes for the ubiquitously expressed torsinA protein (6). TorsinA is a member of the AAA family of ATPases present in the endoplasmic reticulum/nuclear envelope space and is thought to play a role in structural integrity and protein trafficking in the cell (reviewed in ref. 7). The DYT1/TOR1A mutation is responsible for around 50% of early onset primary dystonia cases (8). Most often, the genetic mutation in DYT1-related dystonia is an in-frame GAG deletion (termed ΔE mutant), which is thought to result in at least a partial loss of torsinA function. The penetrance of the mutation is around 30% to 40%, and in these patients the symptoms are variable, ranging from a mild focal presentation to disabling generalized dystonia (8). Although a few of the available studies suggest the presence of rather unspecific structural and morphological alterations in the substantia nigra and the cerebellum (9, 10), there is a general paucity of more comprehensive clinical studies examining structural changes in the brain of DYT1-dystonia patients.

A dystonia model with the right moves

Studies of animals with genetic modification of the Dyt1/Tor1a gene have suffered from the inability of these models to display overt dystonia; therefore, these preclinical models lack validity for studying overt clinical symptoms. Previous models have been engineered with a heterozygous knockin of the Tor1a ΔE mutation or with brain region-specific knockout of the Dyt1 locus; however, these mutations do not result in a clear dystonic motor phenotype (recently reviewed in ref. 11). On the other hand, homozygous knockout of Tor1a or homozygous knockin of the Tor1a ΔE mutation in mice results in neonatal lethality, and therefore, these models are not useful (12). Based on previous attempts to develop a murine dystonia model, Liang and collaborators hypothesized that a dystonia mouse would require torsinA deficiency in critical brain areas during postnatal CNS development in order to develop a clinically relevant phenotype. Toward this goal, Liang and colleagues generated mice that lacked the Tor1a gene in the entire CNS and found that these mice developed striking abnormal twisting movements that are indicative of dystonia. The severe early-onset phenotype observed in mice lacking torsinA in the CNS was accompanied by premature death at postnatal day 16. Liang et al. took their dystonia mouse model one step further and created mice that are heterozygous for the Tor1a ΔE mutation and a floxed WT Tor1a allele, which can be deleted in select tissues, allowing expression of torsinA ΔE in the absence of the WT protein in the CNS. Liang et al. elegantly demonstrate that these animals develop a dystonia-like phenotype as early as the second postnatal week, but survive to adulthood. Interestingly, the CNS-specific Tor1a ΔE mutant mice showed an age-related improvement in behavioral phenotypes, suggesting that the residual function of torsinA may partly compensate for the early deficits in the maturing CNS (1).

Importantly, the behavioral abnormalities in the mouse models developed by Liang and colleagues were accompanied by perinuclear accumulation of ubiquitin as well as neuropathological changes, including gliosis, caspase-3 activation, and ER stress in selective sensorimotor regions. Neuropathology was observed in deep layers of sensorimotor cortex, ventral posterior thalamus, globus pallidus, deep cerebellar nuclei, red nucleus, and the facial nerve nuclei. Thus, a second and equally important aspect of the study by Liang and colleagues is the linkage of neuropathological changes in the brain to a model of “primary” dystonia, which provides further support in the debate on the presence of structural brain changes in this disease. Although Liang and collaborators detected a selective vulnerability of certain sensorimotor cells to torsinA deficiency, further neuropathological analyses will be important to determine the full extent of structural and morphological changes in these models. Nevertheless, the findings by Liang et al. call for further systematic analyses of postmortem tissue from individuals with DYT1-related dystonia in order to elucidate the level of neuropathology in the clinical setting.

Conclusions and future directions

The development of animal models for the study of brain disorders in general has been important for the progress of neuroscience research. Initially, nonspecific lesions in different brain areas of mice helped lead to knowledge about their function. More specific neurotoxic lesions mimicked neuropathological changes observed in clinical conditions and provided a basis both for testing hypotheses related to mechanisms of disease and evaluating restorative therapies. With the identification of the genetic causes of neurodegenerative disorders, the generation of animal models with better construct validity has been possible; however, studies in mice that replicate the genetic mutations identified in patients have seldom resulted in an exact phenocopy of clinical symptoms. In many instances, it appears that the genetic burden in an animal model needs to be exaggerated in order to produce a robust and clinically relevant phenotype. In the case of dystonia, Liang and collaborators engineered mice with alterations of the Dyt1/Tor1a gene that are more severe than what is present in patients and therefore compromised construct validity in favor of the face validity (1). A similar approach to murine model development and analysis has also been beneficial for Huntington disease (HD), a neurodegenerative movement disorder caused by an expanded CAG repeat in the gene coding for the huntingtin protein (13). In the case of HD, the animal models with severe phenotypes were obtained only when mice were engineered to express either a mutant HD protein with a much longer polyglutamine stretch than seen in any patients or a truncated and more toxic form of the protein (14). Nevertheless, the similarities found between clinical manifestations in murine models and patients, such as in the HD field, support the value of these models for examining...
Cardiac fibroblasts have been long recognized as active participants in heart disease; however, their exact physiological and pathological roles remain elusive, mainly due to the lack of specific markers. In this issue of the JCI, Moore-Morris and colleagues used a fibroblast-specific collagen1a1-GFP reporter to demonstrate that fibroblast accumulation after aortic banding in murine hearts arises almost exclusively from proliferation of resident fibroblasts originating from both the epicardium and a previously unrecognized source, the endocardium. Further characterization of fibroblast origin and function in different types and stages of heart disease could lead to development of improved fibroblast-targeted cardiac therapies.

**Lack of specific cardiac fibroblast markers**

Cardiac fibroblasts comprise 30% to 70% of all the cells in the healthy adult heart (1). The number of fibroblasts in the heart is not constant and changes dynamically with development, disease, and aging (2, 3). Traditionally, cardiac fibroblasts have been thought to play passive roles in the heart and to be solely responsible for maintaining homeostasis of extracellular matrix proteins, including type I and III collagens and fibronectin. Due to their ubiquitous presence in the heart, fibroblasts are well poised to actively regulate and modify cardiac function through their direct contacts with other cardiac cells and matrix as well as through secretion of different cytokines, matrix proteins, and proteases. Over the last decade, the pleiotropic roles of fibroblasts in cardiac biology and disease have been studied extensively (reviewed in refs. 4, 5); however, the lack of specific and comprehensive markers of fibroblast phenotypes has hampered the progress in this important research field. In particular, vimentin, the most inclusive marker of cardiac fibroblasts, also labels all other mesoderm-derived cells in the heart. Thy-1 cell antigen-1 (Thy-1, also known as CD90), discoidin domain receptor-2 (DDR2), prolyl-4-hydroxylase (P4H), transcription factor 21 (TCF21, also known as epicardin, Pod1, and capsulin), peristin, cadherin-11, and fibroblast-specific protein-1 (FSP1, also known as S100A4) have all been used to study cardiac fibroblasts, but all of these markers label only a subset of fibroblasts, have poor expression in the healthy adult heart, or nonselectively label endothelial, smooth muscle, or immune/inflammatory cells (6). Because the pathological tissue remodeling that is secondary to cardiac injury and inflammation involves contributions from both resident and extracellular cells, the lack of adequate markers for cardiac fibroblasts may lead to erroneous conclusions about their origin, roles, and potential to be therapeutically targeted in fibrotic heart disease.

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**Address correspondence to:** Deniz Kirik, B.R.A.I.N.S. Unit, BMC D11, 221 84 Lund, Sweden. Phone: 46.46.2220564; Fax: 46.46.2223436; E-mail: deniz.kirik@med.lu.se.