2016 in Review and Message from the Editors to our Reviewers

For this Helix, the editors of Neurology® Genetics have chosen some of their favorite articles of the past year. It is an eclectic collection drawing on different journals including Neurology: Genetics and varied topics.

Two articles report major advances in the understanding of how the C9orf72 GGGGCC repeat expansion exerts its pathogenic effect.1,2 This mutation is a major cause of familial amyotrophic lateral sclerosis (ALS) and frontotemporal dementia (FTD) as well as the cause of 5%–10% of what appears to be sporadic ALS. Understanding the molecular and cellular mechanisms of its pathogenicity is an essential step toward finding therapies for these devastating diseases. The work reported in these articles is remarkable as it identifies effects of both toxic RNA and repeat associated non-AUG (RAN)-translation products generated by this repeat and links them to pathogenic mechanisms shared by other ALS/FTD-causing mutations.

A landmark publication this year describes the generation and characterization of a C9orf72 bacterial artificial chromosome transgenic mouse.3 This mouse model will be important in helping to clarify the pathogenesis of ALS and FTD and can be used to explore the efficacy of potential treatments.

Two articles analyzed white matter changes in FTD due to GRN and C9orf72 mutations and in primary familial brain calcifications due to a novel PDGFB mutation.4,5 The white matter changes preceded atrophy/calcifications and could be a pre-manifest marker in primary familial brain calcification. White matter changes could therefore exacerbate the pathogenic cascades in neurodegenerative diseases characterized by motor and cognitive signs. These changes may include oligodendroglial dysfunction, as suspected for the GRN haplinsufficiency. Extensive white matter changes are present also in mouse models of neurodegeneration, and effects of altered oligodendrocyte function on disease progression are suspected.

Adams et al.6 provide an example of how “crowdsourcing” data collection may provide enough power to detect genetic variants affecting normal variation as well as disease susceptibility. Combining data from many contributors allowed them to identify genetic variants affecting intracranial volume, strictly related to brain volume, which in turn affects the risk of cognitive impairment.

The use of gene therapy to treat peripheral neuropathies faces a number of challenges including ones related to delivery. A recent study describes rescue of an X-linked demyelinating form of Charcot-Marie-Tooth disease (CMT1X) in gap junction beta 1 gene (GJB1) null mice, which lack connexin32 (CX32) protein.7 One intrathecal injection of a lentiviral vector expressing GJB1 gene under the control of the myelin protein zero (MPZ) promoter led to stable and cell-specific expression of CX32 in up to 50% of Schwann cells in multiple lumbar spinal roots and peripheral nerves. The treated mice had improved motor function and electrophysiologic indices, as well as less pathology. The virus vector presumably diffused from the spinal fluid in the subarachnoid space into the epineurium of the peripheral nerve, demonstrating that this delivery route could be used in the treatment of other peripheral neuropathies.

Phuah et al.8 extended the clinical phenotype of APOE ε4 carriers, long known as the major risk factor for Alzheimer disease. They found that APOE ε4 carriers had a significantly greater decline in serum total cholesterol and low-density lipoprotein in the 6 months preceding an intracerebral hemorrhage, relative to APOE ε2 carriers. APOE remains one of the most interesting genes in the CNS, and one of the most important. Despite being identified as a risk factor for Alzheimer disease in 1989, its function in that regard is still not fully understood. Mutations in the gene in humans and mice support its role in

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lipoprotein metabolism. It is fascinating to note that almost 30 years after its rise to clinical importance, we continue to identify its association with major neurologic events.

An impressive multinational effort characterized the variability of dysferlin mutations, including symptoms and rate of progression of their associated disease. Missense mutations were a frequent cause of the disease. The characterization of the mutation, which may cause a spectrum of disease syndromes, is increasingly important in neuromuscular disease as novel therapeutics may be broadly applicable across phenotypes. Indeed, it is likely that the mutation (e.g., missense, nonsense, etc.) is more likely to select the therapeutic trial population rather than the phenotype. Additional, similar, studies in other limb-girdle muscular dystrophies are now possible as the Jain Foundation, Muscular Dystrophy Association, and other patient advocacy foundations have subsidized genetic testing for limb-girdle muscular dystrophies. These resources will extend the knowledge of the genotype-phenotype variability and offer a more accurate understanding of the prevalence of these conditions.

Genome-wide association studies have identified many risk variants, most of them not changing the coding sequence of a gene, but potentially regulating gene expression. Several DNA variants upstream of the SNCA gene encoding α-synuclein have been identified, but their direct relationship to regulating SNCA expression was not known. Soldner et al. made use of a novel method to manipulate the genome, called CRISPR/Cas9, to delete or change specific sequences encompassing previously identified variants. They accomplished these genetic engineering feats in embryonic stem cells that they differentiated into neurons so that SNCA expression could be measured in the correct cell type. Indeed, the variant with the greatest associated risk for Parkinson disease increased SNCA expression in neurons.

The final selection does not directly relate to the nervous system, but is relevant in the context of neurogenetics, as it deals with the evaluation of DNA variants as causative for disease. The Kohane group examined variants that had previously been considered causal in hypertrophic cardiomyopathy. Variants that were falsely classified as pathogenic occurred predominantly in African Americans. Simulations showed that the inclusion of even small numbers of African Americans in control cohorts probably would have prevented these misclassifications, highlighting the importance of diversity in variant databases. The problem of erroneously assigning mutation status to rare, sometimes population-dependent benign DNA variants is widespread. It is of most actionable importance in the clinical arena, but as editors of Neurology: Genetics, we face similar challenges when deciding on acceptance of manuscripts reporting novel mutations in established disease-causing genes.

We wish to acknowledge the individuals who have completed reviews for the journal since its launch in April 2015. Your thoughtful comments are tremendously helpful and highly appreciated. We are also grateful for your cooperation in returning reviews in a timely manner. Please follow the guidelines for reviewing articles accessed by selecting the Information for Reviewers (IFR) link on the Neurology.org/ng website. The IFR provides information on expectations of reviewers regarding confidentiality, timeliness, and reviewer conflicts of interest; it also provides instructions for formatting the comments to editors and authors to enable the most effective communication with authors. Please email ngjournal@neurology.org if you would like to do more reviews or if you have never reviewed for the journal but are interested in doing so. Include a description of your credentials and expertise in the areas in which you are qualified to review. We look forward to hearing from you!

Our 2016 reviewers are listed at the end of this article.

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