Taming Rare Variation With Known Biology in Long QT Syndrome

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Different techniques have been developed to narrow down or filter a large number of rare variants by excluding those that are unlikely to cause disease. Excluding noncoding variants by focusing on exonic sequences, discounting synonymous variants, and eliminating variants cataloged in publicly available databases are standard filters that allowed Boczek et al. to narrow down the list of possible disease-causing variants to 263. The next filtering step was to require that the causative variant be present in affected family members and absent in unaffected members. Although family trios typically consist of mother/father/child, the decision to sequence an affected aunt instead of the mother allowed for a more powerful filter because only one fourth of the genome will be shared between the index and the aunt. In autosomal dominant Mendelian inheritance patterns, inclusion of the father for purposes of segregation does not add significantly to the filtering power; however, this does help eliminate variants that might have been erroneously called because of systematic sequencing errors. Although the expected number of variants left at this point was 65 (approximately one fourth of 263), there were actually 110 variants observed.

Additional exome sequencing of all phenotyped family members would have added power to this filtering step. However, accurate filtering assumes perfect phenotyping, which can be challenging in LQTS because the prolonged QT phenotype is variably expressed. Besides adding to the cost of WES of additional family members, inaccurate phenotyping can result in erroneously filtering out causative variants.

Sanger sequencing and functional validation is burdensome for 100 variants, so investigators have looked to predefined lists of candidate genes. Similar approaches, combining WES with functional genomic filters, have been used to prioritize variants and identify novel genes linked to endometrial cancer. This method has the advantage of narrowing a large list of candidate variants substantially. The disadvantage of this approach is that if the causative variant resides in a gene from a completely novel pathway with no previous suspicion of involvement, then it would likely be filtered out. CACNA1C was fortunately included in the list of 1629 genes from the LQTS interactome, which was used in the next filtering step. It is notable, however, that this list was generated from a protein–protein interaction network based on the set of genes that cause LQT1-12, which includes CACNA1C. Therefore, in this study, CACNA1C could be considered as a candidate gene.

The use of the LQTS interactome list allowed for a helpful filtering step, which reduced the number of candidate variants from 110 to 8. In fact, the only variant that cosegregated completely in all 12 family members, possibly the most convincing evidence of causality, was the Pro857Arg CACNA1C variant. Complete cosegregation was achieved by classifying...
borderline-affected family members phenotype-positive, a reasonable strategy in the investigation of causality in a condition with variable expressivity of the phenotype.

Of the 8 variants identified after the final filters were applied, all 3 bioinformatics/systems biology algorithms (Endeavour, SUSPECTS, and ToppGene) ranked CACNA1C highest. In their ranking algorithms, 2 of these programs use resources such as semantic analysis of existing literature, in which CACNA1C is frequently included among the list of genes that are already linked to LQTS. The 3 LQTS-disease—causing genes (KCNQ1, KCNH2, and SCN5A) that were used as the algorithms’ training genes are well characterized with extensive ontologic labels. These factors likely helped CACNA1C rise to the top of the ranked lists. Creating systems biology—based filters with sets of genes that are not as well characterized in Mendelian disorders for which several causative genes do not already exist may not result in such accurate predictions.

Impressively, these investigators went beyond human genetic evidence to explore the functional effect of this variant on the expressed channel. Functional analysis of the Pro857Arg CACNA1C variant in HEK293 cells demonstrated gain-of-function of the CACNA1C channel, likely mediated through impaired degradation of Ca2+.12 These findings in addition to the cosegregation observation support the hypothesis that this variant in CACNA1C is LQTS-causing and suggest a novel mechanism by which CACNA1C trafficking disorders can lead to LQTS.

To further demonstrate that CACNA1C causes nonsyndromic LQTS and may play a more prominent role in LQT, Boczek et al6 performed mutational analysis in 102 previously undiagnosed families with LQT syndrome and found an additional family with rare variation in CACNA1C. Limited cosegregation was evident in 1 of the families whose CACNA1C variant affected the same amino acid as the index family. Presence of rare variation alone is not sufficient to demonstrate pathogenicity.16 However, the fact that 2 of these variants were located in the same domain (proline—glutamic acid—serine—threonine—rich domain [PEST]) responsible for degradation signaling is suggestive of common pathogenicity. Population variation reveals rare missense variation in CACNA1C coding regions in 38 out of 4300 (0.9%) unrelated Caucasian individuals presumably free of Mendelian disease. This rate is lower than the rate of 3 out of 102 (2.9%) patients with nonsyndromic LQTS reported in the mutational analysis by Boczek et al9 (χ2 = 0.031). This further supports the hypothesis that CACNA1C is linked to nonsyndromic LQTS.

Although CACNA1C is known to cause prolongation of the QT interval as 1 of several clinical manifestations of Timothy Syndrome,18 this is the first nonsyndromic LQTS family whose disease has been linked to a CACNA1C mutation. Although variants in genes linked to LQT4—15 are responsible for <5% of LQTS,19 this and other recent studies suggest that CACNA1C may play a more prominent role in nonsyndromic LQTS than was originally believed. Lieve et al20 reported that, in patients with suspected LQTS who underwent sequencing of the genes responsible for LQTS1—10, rates of rare variation of unclear significance were highest in CACNA1C. This could be either because rates of benign rare variation are higher in this gene, or these variants were determined of unclear significance because they have not been as frequently documented in the literature.

We expect that as the cost of sequencing continues to plummet, WES in many more families with uncharacterized Mendelian diseases will ultimately be performed. However, finding large families with clear positive and negative phenotypes for conclusive cosegregation analysis will remain challenging. Because of this, filters that use systems biology algorithms, such as those presented by Boczek et al,9 will become an important step in helping narrow down large lists of plausible variants. There will need to be a balance between using unbiased whole-genome approaches that are more likely to uncover genes in novel disease pathways and a candidate gene—based approach where effective filters link genes in well—established disease pathways.

Despite technical challenges in clinical grade coverage from current capture technologies and algorithmic challenges in calling small insertion/deletions and structural variation, exome sequencing continues to offer an exciting alternative for families where traditional panel—based sequencing has failed to identify a convincingly causal variant of disease. Boczek et al6 have demonstrated with cosegregation, functional, and mutational analyses that CACNA1C is linked to nonsyndromic LQTS. Because commercially available LQT panels now routinely include sequencing of most exons in CACNA1C, not just exon 8 that is responsible for Timothy Syndrome, we anticipate a growing number of reports of CACNA1C variants linked to nonsyndromic LQTS.

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