Old meets new: identifying founder mutations in genetic disease

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From the first arrivals over 10 000 years ago to the most recent immigrants, every person coming to Canada has carried with them a unique genetic contribution. Once here, forces such as the size of founder populations and isolation by geography or cultural practices have led to a high prevalence of specific genetic conditions in many diverse ethnic groups. Although an appreciation of such variation in disease frequency and population genetics principles has been around for some time, the application of modern methods for gene identification now has the potential to improve outcomes for patients, their families and communities.

Examples of this approach are well illustrated by two linked CMAJ articles. In the first, Macardier and colleagues document the finding of a common mutation in the SI gene responsible for congenital sucrase–isomaltase deficiency in Inuit, a condition highly prevalent in this population. In the second, Rousseau-Nepton and colleagues describe the use of whole-exome sequencing in children with glycogen storage disease type IIIa to identify a founder mutation in the AGL gene in Inuit from Nunavik.

Both of these studies deal with autosomal recessive disorders that are common in Inuit populations. Both conditions are caused by homozygous mutations in their respective genes, and yet, despite identical genotypes, affected patients can present with variable manifestations. In both diseases, early diagnosis and dietary interventions may help ameliorate the clinical consequences of the deleterious genotype.

However, although it is apparent that the SI mutation is widespread in Inuit populations, suggesting an ancient origin, the AGL mutation in the Inuit population has so far only been identified in those from Nunavik. Inuit from other regions may also be gene carriers, but the lack of reports of glycogen storage disease type IIIa in other areas suggests a more recent mutational event. Haplotype analysis can provide information on the relative age of founder mutations, because regions of homozygosity around a gene of interest become shorter with time as a result of meiotic recombination. Furthermore, the SI mutation appears to be unique and restricted to the Inuit, whereas the AGL mutation was found to be the cause of glycogen storage disease type IIIa in a North African Jewish population, and its heterozygous state has been found in genome databases. Rousseau-Nepton and colleagues suggest that this coincidence is likely due to mutations at a genetic “hot spot” rather than to any shared ancestry. However, introduction of alleles from other groups into indigenous populations is always possible. For example, evidence exists from both molecular data and oral history that the founder mutation for autosomal recessive infantile leukoencephalopathy, seen in Cree and Chippewayan populations in Manitoba and Cree in Northern Quebec, may have come via British fur traders for the Hudson Bay Company in the 1770s.

Globally, founder effects have lead to similar patterns of high prevalence of rare diseases in specific populations. Other Canadian examples include infantile hypophosphatasia among Mennonites and Bowen–Conradi syndrome in Hutterites. Founder alleles can also contribute to risk for more common adult-onset disorders. Among the Oji-Cree in northwestern Ontario, a population with high rates of type 2 diabetes mellitus, each copy of a unique mutation in HNF1A (hepatocyte nuclear factor 1 homeobox A) lowers the median age of onset by about...
seven years. Similarly, specific alleles in cancer predisposition genes may be more common in certain ethnic groups, such as 999del5 in BRCA2 (breast cancer 2, early onset) among Icelandic families with breast cancer or founder MSH2 (MutS protein homolog 2) mutations in Lynch syndrome in Newfoundland and Labrador.

Awareness of such variation in distribution of disease has allowed an important preventative health strategy: using ethnic background to identify people who should be offered screening for conditions such as sickle cell anemia, thalassemias and Tay–Sachs disease. Taking a detailed three-generation family history may provide useful clues as to whether a condition is familial and what the reasons for that might be — genes, shared environment or interactions between them. However, assumptions cannot be made that certain genetic disorders only occur in particular ethnic groups. Cystic fibrosis, although more common in white people, occurs in other populations, such Canadian and Australian indigenous groups owing to either admixture or spontaneous mutation. Qureshi and Kai tell a cautionary tale of how ignorance that people with white skin can carry sickle-cell mutations led to delayed diagnosis in an affected child and the erroneous assertion of nonpaternity.

Attention to ancestral background will help assess the probability of genetic disease. In addition, the possibility of parental consanguinity should be explored because it increases risks for both recessive and multifactorial disorders. However, as the linked articles document, in many genetic isolates, close consanguinity is not common, and such populations may in fact be “outbred” (i.e., marriage partners are often less closely related than might happen by chance). Consanguinity increases the likelihood of homozygous genotypes, both wild type and deleterious ones, but it has no impact on allele frequency. Usually, the high prevalence of recessive disease is a consequence of a high carrier frequency due to founder effect and, potentially, heterozygote advantage.

Consanguinity has, however, been very beneficial to the discovery of new recessive mutations using homozygosity mapping. When the culprit gene is already suspected, sequencing may identify the mutation, though this can be challenging, as was initially seen with AGL in the Nunavik population. Next-generation molecular tools have shown great potential to accelerate discovery in this area, and it should be a source of pride that Canadian initiatives to find mutations for rare genetic disorders (such as the Finding of Rare Disease Genes in Canada [FORGE Canada] Consortium) have been so productive.

As suggested in both articles, once a causative mutation is discovered, opportunities exist to use molecular rather than more invasive diagnostic testing, to start treatment early and to offer carrier testing to family members at risk. In collaboration with the community, targeted population screening can be considered, as has already been done for carnitine palmitoyltransferase I A (CPT1 A) deficiency in Hutterite populations, the ultimate goal being to reduce the burden of disease and provide patients and their families with more personalized genetic health care.

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