Genome-Wide TDT Analysis in French-Canadian Families with Tourette Syndrome

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BRIEF COMMUNICATIONS

Tourette syndrome (TS) is a complex neurodevelopmental disorder with an estimated prevalence of 1% that is characterized by motor and vocal tics as well as psychiatric comorbidities, such as obsessive-compulsive disorder (OCD) and attention deficit hyperactivity disorder (ADHD).10-12 Despite a strong genetic contribution, no common variants have been clearly associated with the disorder, possibly because of allelic and non-allelic genetic heterogeneity. Several candidate genes involved in dopaminergic neurotransmission have been analyzed based on the observation that neuroleptics are used to treat TS patients. Positive association results between TS and some of these genes have been reported. However, because other studies failed to replicate these results, the role played by dopaminergic in TS remains unclear.10-12 Founder populations from families with a high risk of breast and ovarian cancer.16 For within a heterogeneous population.13 For instance, the G2019S substitution in the LRRK2 gene accounts for 20–40% of North-African Arab and Ashkenazi patients with Parkinson disease.14,15 In the French-Canadian (FC) population, founder mutations in the BRCA1 and BRCA2 genes were identified in 40% of patients from families with a high risk of breast and ovarian cancer. For several reasons, the FC population of Quebec displays all the characteristics of a population isolate. An estimated 2,600 pioneers who settled in Quebec before 1680 account for two thirds of the modern FC gene pool and a vast majority of FC pioneers who settled in Quebec before 1760. Given that these founders rarely mixed with other immigrants over three centuries and that there was a sustained demographic growth in the FC population, the ~6 million FC individuals currently living in Quebec inherited most of their genes from a relatively small pool of founders.17

Based on a cohort of 217 FC trios (father, mother and proband) presenting with TS, the goal of this study was to assess the presence of frequent and highly penetrant alleles predisposing to TS in the FC population. Affected individuals and their relatives were recruited through the Montreal General Hospital and the Sainte-Justine Hospital (Montreal). Experienced clinicians performed diagnostic evaluations of TS, chronic tics and ADHD using the Diagnostic and Statistical Manual of Mental Disorders-IV criteria. Obsessive-compulsive disorder was evaluated using the Yale–Brown Obsessive-Compulsive Scale. We obtained approval from the ethics committee of our institution, as well as informed consent from all participants. The DNA was extracted from whole blood using standard procedures. For the purpose of the family-based genome-wide association analysis, we selected 95 trios among the 217 nuclear families recruited. To avoid the inclusion of sporadic cases, we selected only patients with a positive familial history of TS, tics or comorbid psychiatric disorders. Among the 95 selected TS cases, 73 were male, 31 presented with OCD and 53 received a diagnosis of ADHD. The DNA samples from the probands and their parents were sent to the deCODE genotyping service (http://www.decode.com/genotyping). Five hundred fifty-one highly polymorphic microsatellites covering the 22 autosomes and the X chromosome with an average marker density of 8 centimorgans were genotyped using standard methods. The successful genotyping rate was 95.8% and Mendelian inconsistencies were systematically removed. We performed a family-based transmission disequilibrium test (TDT) using the Family-Based Association Test (FBAT) program, version 2.0.2C. The TDT determines whether an excess or a lack of transmission of alleles to the affected offspring occur. Multiallelic tests of association were performed for each marker. The additive genetic model was applied, as it often performs best, even when the genetic model is not additive. The significance level was set at 0.00009 after Bonferroni correction for multiple testing. Employing the TDT Power Calculator software, we assessed the statistical power of the TDT from 95 trios22. Table 1 displays the number of trios...
estimated to be necessary to achieve 90% statistical power with a significance level of 0.00009, according to variable disease susceptibility allele frequencies and degrees of linkage disequilibrium between a marker and a disease susceptibility locus. Although none of the markers reached the significance level for association, we obtained $P$ values < 0.005 for four markers (D7S2485, D13S271, D15S1016 and D19S605; Figure and Table 2). None of the markers are located in any of the chromosomal loci proposed to be linked to TS by the Tourette Syndrome Association International Consortium for Genetics.23,24 To attempt to replicate our findings, the four markers were genotyped in the remaining 122 FC trios from our initial cohort of 217 FC trios with TS. Unfortunately, the analysis of the genotype data from the 217 FC trios revealed that none of the markers remained associated with TS (Table 2).

Our data suggest that common and highly penetrant founder mutations are unlikely to predispose to TS in the FC population. Our inability to replicate the association when the whole TS cohort was included in the analysis suggests that the peaks observed in the initial analysis were likely false-positive signals, which is strengthened by the fact that no single marker reached the genome-wide significance threshold. One may argue that the lack of replication of the association in the whole cohort may stem from the fact that we selected the 95 trios based on positive familial history of TS, and that the 122 remaining trios may comprise a significant proportion of non-genetic cases of TS. This is unlikely, as the vast majority of the cases in our cohort have first-degree relatives with TS, tics, ADHD or OCD, which indicates that sporadic cases with TS are the exception, rather than the rule. Our results suggest that, even in the case of a founder population (such as the FC), the genetically complex nature of TS renders this type of analysis powerless; the lack of significant association results in the analysis presented here may be explained by the complex inheritance pattern of TS and its comorbidities25, which have a great negative impact on family-based studies. We are aware that our study presents several limitations, which include the relatively small number of families and the low marker density of the genome-wide scan. As presented in Table 1, statistical power computation indicates that our sample size was probably large enough to detect an allele segregating with a founder mutation carried by 20 to 40% of patients, but the study was most likely underpowered for founder mutation frequencies of 10% or less, unless the marker allele was in perfect linkage disequilibrium with a founder mutation. A high-density single nucleotide polymorphism genome-wide association study using a large cohort of patients may be necessary to verify whether TS originates from the combined effect of several common and low penetrant alleles. It would also

<table>
<thead>
<tr>
<th>Marker</th>
<th>Chr</th>
<th>Position (bp)</th>
<th>95 trios $P$ value</th>
<th>217 trios $P$ value</th>
</tr>
</thead>
<tbody>
<tr>
<td>D7S2485</td>
<td>7q21.11</td>
<td>83,997,436</td>
<td>0.000595</td>
<td>0.451994</td>
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<tr>
<td>D13S271</td>
<td>13q31.1</td>
<td>84,278,764</td>
<td>0.003924</td>
<td>0.325128</td>
</tr>
<tr>
<td>D15S1016</td>
<td>15q21.3</td>
<td>51,320,121</td>
<td>0.000341</td>
<td>0.234887</td>
</tr>
<tr>
<td>D19S605</td>
<td>19q13.42</td>
<td>60,443,625</td>
<td>0.003598</td>
<td>0.163923</td>
</tr>
</tbody>
</table>

“Chr” refers to the chromosome band. The position of markers is in base pairs (bp) and was derived from the publicly available human genome assembly (UCSC Genome Browser).
be feasible to use ADHD and OCD diagnoses as covariates, or to analyze subgroups of TS patients separately, based on their comorbidities. However, this approach would necessitate large cohorts of patients and would reduce the statistical power because of multi-test adjustments. If the genetic model for TS is that many cases are caused by individually rare, but highly penetrant mutations, this approach will not be successful, and a whole-genome resequencing analysis or large-scale screening of candidate genes will be warranted. Finally, this study illustrates the difficulty in identifying susceptibility genes for neuropsychiatric disorders and emphasizes the necessity to develop new gene-mapping approaches for these complex disorders.

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REFERENCES