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Genome-Wide Association Study to Find Modifiers for Tetralogy of Fallot in the 22q11.2 Deletion Syndrome Identifies Variants in the GPR98 Locus on 5q14.3

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Background—The 22q11.2 deletion syndrome (22q11.2DS; DiGeorge syndrome/velocardiofacial syndrome) occurs in 1 of 4000 live births, and 60% to 70% of affected individuals have congenital heart disease, ranging from mild to severe. In our cohort of 1472 subjects with 22q11.2DS, a total of 62% (n=906) have congenital heart disease and 36% (n=326) of these have tetralogy of Fallot (TOF), comprising the largest subset of severe congenital heart disease in the cohort.

Methods and Results—To identify common genetic variants associated with TOF in individuals with 22q11.2DS, we performed a genome-wide association study using Affymetrix 6.0 array and imputed genotype data. In our cohort, TOF was significantly associated with a genotyped single-nucleotide polymorphism (rs12519770, P=2.98×10−8) in an intron of the adhesion GPR98 (G-protein–coupled receptor V1) gene on chromosome 5q14.3. There was also suggestive evidence of association between TOF and several additional single-nucleotide polymorphisms in this region. Some genome-wide significant loci in introns or noncoding regions could affect regulation of genes nearby or at a distance. On the basis of this possibility, we examined existing Hi-C chromatin conformation data to identify genes that might be under shared transcriptional regulation within the region on 5q14.3. There are 6 genes in a topologically associated domain of chromatin with GPR98, including MEF2C (Myocyte-specific enhancer factor 2C). MEF2C is the only gene that is known to affect heart development in mammals and might be of interest with respect to 22q11.2DS.

Conclusions—In conclusion, common variants may contribute to TOF in 22q11.2DS and may function in cardiac outflow tract development. (Circ Cardiovasc Genet. 2017;10:e. DOI: 10.1161/CIRCGENETICS.116.001690.)

Key Words: chromosomes ■ DiGeorge syndrome ■ genotype ■ ivelo-cardio-facial syndrome ■ tetralogy of Fallot

One of the greatest challenges in the area of human genetics is to understand the basis of phenotypic heterogeneity in known diseases. The 22q11.2 deletion syndrome (22q11.2DS; velocardiofacial syndrome/DiGeorge syndrome; Mendelian Inheritance in Man No. 192430, 188400) is one of the most common genomic disorders, occurring in 1 of 4000 live births.1 Over 90% of affected individuals have a de novo, hemizygous 3 million base pair (Mb) deletion on chromosome 22q11.2.2-4 All subjects with the deletion have features of the syndrome,
but the clinical presentation is quite variable. For example, 60% to 70% of patients have congenital heart disease (CHD) involving the cardiac outflow tract (OFT) and aortic arch, whereas the rest have apparently normal cardiac structures. Among the most serious defect observed in individuals with the 22q11.2DS is tetralogy of Fallot (TOF), which is defined by the presence of a ventricular septal defect, pulmonary stenosis, overriding aorta, and right ventricular hypertrophy. TOF is caused in part by failed migration or differentiation of second heart field mesodermal cells from the pharyngeal apparatus in embryos, needed to form or remodel the cardiac OFT. One hypothesis to explain variable phenotypic expression results in neonatal lethality with severe cardiac OFT defects. 9–11 Global inactivation or second heart field–specific inactivation of Tbx1 results in neonatal lethality with severe cardiac OFT defects. 

Among the genes in the deleted region on 22q11.2, TBX1, which encodes a T-box transcription factor, is the major candidate for CHD. Tbx1 is expressed in the second heart field mesoderm, which is disrupted in 22q11.2DS.9–11 Global inactivation of Tbx1 or second heart field–specific inactivation of Tbx1 results in neonatal lethality with severe cardiac OFT defects. 

One hypothesis to explain variable phenotypic expression in the 22q11.2DS is the presence of pathogenic variants in Tbx1 on the haploid allele of 22q11.2. Previously, we tested whether common or rare single-nucleotide variants (SNVs) in the coding region of Tbx1 in the remaining allele of 22q11.2 were associated with CHD in 22q11.2DS subjects, but we did not find an association. 13 Another hypothesis is that there are copy number variations elsewhere in the genome that could explain differences in phenotypes. We previously found that a commonly occurring genomic duplication encompassing the glucose transporter gene, SLC2A3, was associated with CHD (P = 2.68×10−6).14 This copy number variation occurred in 5.8% of individuals with 22q11.2DS and CHD and 1.1% of those with 22q11.2DS and no CHD. Recently, a partial duplication of a chromatin modifier, KANSL1, was associated with CHD in a Chilean 22q11.2DS cohort.15 However, these copy number variations occurred in only some deleted subjects with CHD and thus do not explain the basis of phenotypic variability in the majority of patients.

Our goal was to identify common single-nucleotide polymorphisms (SNPs) that are associated with TOF in individuals with 22q11.2DS. We restricted our analyses to TOF because it is the largest single phenotypic category of severe CHD in our cohort. Restricting our analyses in this way may reduce heterogeneity in the genes that contribute to CHD in individuals with 22q11.2DS and thus may increase the power of a GWAS.

Methods

Human Subjects and Phenotype Data

We assembled a cohort of subjects with 22q11.2DS (Tables I and II in the Data Supplement). Subjects were previously recruited by the International Chromosome 22q11.2DS Consortium, the International 22q11.2 Brain Behavior Consortium (http://22q11-ibbc.org), and clinical groups that specialize in the treatment of individuals with 22q11.2DS. All subjects within the cohort had a clinical diagnosis of 22q11.2DS that was confirmed by the presence of a 22q11.2 deletion using fluorescence in situ hybridization or multiplex ligation-dependent probe amplification (SALSA MLPA kit P250 DGe George; MRC Holland, The Netherlands). Informed consent was obtained for all participants, and this study was conducted under an Internal Review Board-approved protocol at the Albert Einstein College of Medicine (CCI 1999-201). For this study, we used previously collected genomic DNA and phenotypic and demographic information. We obtained echocardiogram and cardiology reports to confirm the specific CHD diagnosis (eg, TOF).

SNP Array Genotype and Data Quality Control

Genomic DNA from 1244 study subjects was array genotyped using Affymetrix GeneChip Genome-Wide SNP 6.0 array. The majority of samples were genotyped at the Genomics Facility core laboratory of Albert Einstein College of Medicine. However, 37 samples were genotyped at the Advanced Genomics laboratory core at the Children’s Research Institute (Milwaukee, WI) for clinical purposes, and 191 Chilean samples were genotyped in the Center for Human Genetics, Clínica Alemana Universidad del Desarrollo, Santiago, Chile.

The raw data from all arrays were processed through the same pipeline using the same criteria. Genotype data from arrays with contrast quality control scores ≤0.4 per sample, contrast quality control <1.7 per batch, and Median Absolute Pairwise Difference metric >0.35, were excluded. Genotypes were called using the Birdseed V2 Genotyping Algorithm (call rate: 99.02±0.02%). To account for batch effects, BEAGLECALL Version 1.0.1 software was used to rescore genotypes.17 SNPs with call rates <95%, minor allele frequency <1%, or Hardy–Weinberg equilibrium P value <10−6 were excluded. In addition, samples that showed second-degree relatedness or closer, based on identity by state, were removed. For each subject, deletion size was determined using the log, intensity ratio as estimated by the Copy Number Analysis Module of Golden Helix Powerseat Package. The CEL files and genotype data are being deposited to National Center for Biotechnology Information database of Genotypes and Phenotypes phs001339.v1.p1.

We performed imputation to increase the number of SNPs available for analysis. Only genotyped SNPs with minor allele frequency >1% were used for imputation. Haplotypes were prephased using SHAPEIT software;18,19 and imputation was performed using IMPUTE2 with the 1000 Genomes Phase I data set as the reference panel.20 Imputed SNPs with minor allele frequency ≤1% or imputation quality (INFO) scores ≤0.8 were excluded from the GWAS.

Statistical Methods

We used a case–control approach, in which individuals with 22q11.2DS and TOF were considered cases and individuals with 22q11.2DS without CHD were considered controls. We conducted principal component analyses to identify the PCs of race/ethnicity. Potential associations between TOF, sex, and deletion size were assessed using logistic regression adjusted for the first 4 PCs. A P value <0.05 was considered significant.

The association between TOF and each SNP was assessed by logistic regression analysis under an additive genetic model using data from all study subjects. These analyses were performed using SNPTTEST v2.5.2 https://mathgen.stats.ox.ac.uk/genetics_software/snptest/snpstest.html and accounted for the genotyping accuracy and first 4 PCs of race/ethnicity.21 A P value of 5×10−8 was used as the genome-wide significance cutoff for single association tests. For a meta-analysis, the cohort was split into groups determined by principal component analyses. Each group was analyzed separately using logistic regression, and the results were meta-analyzed using the inverse-variance method. Power for these analyses was assessed using QUANTO (http://biostats.usc.edu/Quanto.html). Manhattan plots and quantile–quantile (Q–Q) plots were generated using Golden Helix Powerseat. For regions of interest identified in the GWAS, regional association plots were generated using LocusZoom software (http://locuszoom.sph.umich.edu/locuszoom/).23

Conditional logistic regression analyses were performed to determine whether multiple variants within a region are independently associated with TOF in individuals with 22q11.2DS. Specifically, within a region, we conditioned on the genotyped SNP with the
lowest $P$ value and the first 4 principal components of race/ethnicity and individually evaluated the association of TOF with each additional SNP in the region. Conditional analyses were conducted using SNPTEST v2.5.2.

**Linkage Disequilibrium Analysis of Whole-Genome Sequence to Identify Variants in Linkage Disequilibrium With GWAS Findings**

Whole-genome sequencing of a subset (n=397) of our 22q11.2DS samples was performed using the Illumina HiSeq2000 and HiSeq X Ten platform at Hudson Alpha Institute for Biotechnology (Huntsville, Alabama) as part of the International 22q11.2 Brain and Behavior Consortium to find genes for schizophrenia. Variant calling was performed using PEMapper software for read mapping to the hg38 (GRCh38) reference genome and PECaller software for variant calling.24 CrossMap (http://crossmap.sourceforge.net/) was used to convert genome coordinates between hg38(GRCh38) and hg19 (GRCh37).25 To follow-up on the top result from the GWAS (Results), the genomic region (chromosome 5 [chr5]:88,703,723–91,409,593) from MEF2C through ARRD3C was extracted from the whole-genome sequencing data. Functional annotation of SNVs was performed using the Variant Classification and the Annotate and Filter tools in the Golden Helix software. Nonexonic SNVs (intronic, intergenic) were removed, and predicted functional SNVs were used to generate an linkage disequilibrium (LD) matrix using Haploview. LD measurements of $r^2 > 0.8$ were used to define LD haplotype blocks.

**Mouse Embryo Analysis and Whole-Mount RNA In Situ Hybridization**

Gene expression profiling was previously performed to identify differentially expressed genes in the second heart field mesoderm of wild-type mouse embryos.12 Data from wild-type embryos were extracted for evaluation of specific expression levels in this tissue. For in situ hybridization, RNA probes were generated from mouse embryo cDNA using digoxigenin-uridine triphosphate (Roche Diagnostic Corp, Indianapolis, IN; Table III in the Data Supplement). In this cohort, neither sex nor race was significantly associated with TOF in this cohort, these variables were not included in the logistic models. The genomic inflation factor ($\lambda=1.02$) and the Q–Q plot (Figure II in the Data Supplement) provided little evidence of a systematic deviation from the expected distribution of the test statistic.

Three SNPs mapping to intron 61 of GPR98 (G-protein–coupled receptor 98) were significantly associated with TOF. The genotyped SNP, rs12519770 ($P=2.98 \times 10^{-4}$),

**GWAS to Identify Genetic Loci for TOF**

The TOF phenotype comprised the largest individual group of subjects with severe intracardiac anomalies in our 22q11.2DS population (36%; Table 1; Figure 1). We conducted a GWAS to identify genetic variants associated with TOF. This analysis was based on data from 326 subjects with 22q11.2DS and TOF and 566 subjects with 22q11.2DS and normal cardiac anatomy. This study had a power of 80% to detect an odds ratio of >1.9 for a common SNP with an allele frequency >0.3 under a log-additive model at $P < 5 \times 10^{-8}$.

Subjects of all races and ethnicities were included, and associations were assessed using logistic regression adjusted with the first 4 PCs of race/ethnicity. As neither deletion size nor sex was significantly associated with TOF in this cohort, these variables were not included in the logistic models. The genomic inflation factor ($\lambda=1.02$) and the Q–Q plot (Figure II in the Data Supplement) provided little evidence of a systematic deviation from the expected distribution of the test statistic.

Three SNPs mapping to intron 61 of GPR98 (G-protein–coupled receptor 98) were significantly associated with TOF. The genotyped SNP, rs12519770 ($P=2.98 \times 10^{-4}$),

<table>
<thead>
<tr>
<th>Table 1. Characteristics of Study Subjects</th>
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<tbody>
<tr>
<td><strong>Subject Characteristics</strong></td>
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<tr>
<td><strong>Congenital heart defect</strong></td>
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<tr>
<td>TOF</td>
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<td>Other CHD</td>
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<tr>
<td>Normal (controls)</td>
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<td><strong>Deletion size</strong></td>
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<tr>
<td>Typical 3 Mb (LCR22A-D)</td>
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<tr>
<td>Nested LCR22A-B</td>
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<td>Nested LRC22A-C</td>
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<td>Other</td>
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The cohort of 1472 subjects is shown categorized based on self-reported race, ethnicity, and sex. The numbers and percentages in each category are indicated. A total of 906 subjects have congenital heart disease (CHD), whereas the rest have normal structures. The deletion sizes are indicated. LCR indicates low copy repeats; and TOF, tetralogy of Fallot.
and imputed SNPs, rs7720206 \((P=2.22\times10^{-8})\) and chr5: 90067043:D \((P=2.10\times10^{-8})\), showed the strongest association (Figure 2A, Table 2). These three SNPs seem to be in complete LD (Figure 2B and 2C). For rs12519770, the A allele was the risk allele with a frequency of 0.58 in TOF cases and 0.45 in controls, conferring an odds ratio of 1.69 \((P=3.2\times10^{-4})\) per copy of the A allele in the 22q11.2DS cohort (Table 2; Figure 2). There was also suggestive evidence of association between TOF and 2 additional groups of SNPs in GPR98. The top genotyped SNP in each cluster (rs6889138, rs6893710) is listed in Table 2 and illustrated in Figure 2B and 2C.

Although our initial GWAS adjusted for the first 4 PCs of race/ethnicity, the observed associations may still reflect bias because of uncontrolled confounding resulting from population stratification. Consequently, we repeated our analyses for the top SNP, rs12519770, after separating the cohort into 3 groups: white, Admixed, and African, as determined by principal component analyses (Figure I in the Data Supplement). The \(P\) value for this SNP was significant in the meta-analysis \((P=4.43\times10^{-8};\) Table V in the Data Supplement), suggesting that the observed association is unlikely to be the result of population stratification.

Within the 5q14.3 region, there seemed to be 3 clusters of SNPs that were associated with TOF. We refer to these as clusters 1, 2, and 3, and the clusters are ranked in ascending order based on the \(P\) value for the top SNP within the cluster. To determine whether >1 variant was independently associated with TOF, we performed conditional analyses in which we conditioned on the genotyped SNP in GPR98 with the smallest \(P\) value in cluster 1 (rs12519770; \(P=2.98\times10^{-8}\)) and individually evaluated the association of TOF with each of the additional SNPs in the 5q14.3 region \((n=1344\) SNPs, Table VI in the Data Supplement). In these conditional analyses, there was suggestive evidence for association with 1 SNP (rs6893710, \(P=3.92\times10^{-4}\)). This variant was the top SNP in the third cluster of associated genes (Table 2). The association of the top SNP in the second cluster (rs6889138) was attenuated in the conditional analysis \((P=0.002)\).

**Definition of the 5q14.3 Locus**

To identify nonsynonymous variants that may be in LD with the rs12519770 and to narrow the region containing the association signal on 5q14.3 based on LD, we performed an LD analysis using existing whole-genome sequence data from 397 individuals with 22q11.2DS (http://22q11-ibbc.org; unpublished data, International 22q11.2 Brain and Behavior Consortium authors in Supplementary Table 1, 2017). These individuals comprise a subset of the samples genotyped on Affymetrix 6.0 arrays and were selected based on psychiatric but not cardiovascular phenotype. There were 9680 SNVs identified in the 2.7 Mb region around GPR98 \((chr5: \text{8799640–90704983})\). There were 161 coding SNVs (102 nonsynonymous, 58 synonymous, and 1 splicing) and 115 SNVs in the 3’ or 5’-untranslated regions for a total of 276 SNVs (Table VII in the Data Supplement). None of these SNVs were in LD with rs12519770. The SNP, rs6893710, was in weak LD with the synonymous variant, rs41304884 \((GPR98, \text{NM_032119.3, c.16164G>A; D'=0.859, r²=0.389; Figure 2B})\). There were no nonsynonymous variants related to our association signal.

Most GWAS signals that have been previously discovered are in intergenic regions and may mark transcriptional regulatory regions in the genome rather than genes themselves. To test this for TOF in 22q11.2DS, we examined the LD pattern from available whole-genome sequencing data on the same 397 22q11.2DS subjects to narrow the interval with SNPs showing the strongest association. We narrowed down the association signal to a 104.7 kb region on chromosome 5 \((chr5: \text{90057563–90162285};\) Figure 2C and 2D, red block). Most of the common SNPs with \(P\) values \(<10^{-5}\) were located in this region (Figure 2C). A similar LD pattern has been observed in the white subset from the 1000 Genomes Project (Figure III in the Data Supplement). Thus, we were able to narrow the region of the association signal.

**Genes Mapping to 5q14.3**

Because the SNPs found within the intron of GPR98 might affect its regulation or, instead, the regulation of other genes in the region, we examined local chromosome conformation forming topologically associated domains (TADs).\(^{26–28}\) To identify higher-order chromatin-mediated looping, we extracted data for the 5q14.3 interval from the Hi-C browser.\(^{29}\) There was chromatin interaction data for 28 different cell lines ranging from H1 embryonic stem cells to cancer cell lines.\(^{29–31}\) We focused on TAD contact domains in a 3.3 Mb region including GPR98 (Figure 3; Figure IV in the Data Supplement). Because the 104.7 kb region with genetic association is within the GPR98 locus, it is possible that variants might affect its expression or that of nearby genes. In addition to GPR98, this region includes 6 additional genes (Figure 3; Figure IV in the Data Supplement), including 5 protein coding genes within the 2.3 Mb TAD: MEF2C (Myocyte enhancer factor 2C), CETN3 (Centrin 3), MBLAC2 (Metallo-β-lactamase domain containing 2), POLR3G (RNA polymerase III subunit G), and LYSMD3 (LysM, putative peptidoglycan-binding, domain containing 3). One gene, ARRD3 (Arrestin domain

Figure 1. Distribution of cardiovascular phenotypes in 1472, 22q11.2 deletion syndrome (22q11.2DS) subjects. The number of subjects (y axis) sorted into phenotypes (x axis) is shown in the bar graph. All individuals have a hemizygous 22q11.2 deletion. The most serious cardiovascular diagnoses with the largest number of subjects is tetralogy of Fallot (TOF; n=326; black bar) among a total with congenital heart disease (CHD; n=906; gray bar) when compared with those with no intracardiac or aortic arch anomalies as detected by echocardiogram summary and cardiology report (white bar).
Figure 2. Genome-wide association results for tetralogy of Fallot (TOF) in 22q11.2 deletion syndrome (22q11.2DS). A, Values in the Manhattan plot for TOF vs controls were plotted against their respective positions on the autosomal chromosomes. The red line represents the genome-wide significance threshold ($P=5\times10^{-8}$). The blue line represents the threshold for suggestive association ($P=1\times10^{-5}$). A single locus marked by the GPR98 (G-protein–coupled receptor V1) gene reached genome-wide significance. B, LD matrix of selected, predicted functional SNPs with top $P$ values on the 5q14.3 region from WGS (Methods). The LD with respect to associated single-nucleotide polymorphisms (SNPs) with highest $P$ values in the region is shown. The LD plot is based on $r^2$ values. Key: $r^2=0$ is given in white, $0<r^2<1$ is given in shades of grey and $r^2=1$ is given in black. The pairwise $D'$ values are provided in the boxes. Nine SNPs are in modest LD ($D'=0.80$, $r^2=0.23$) with rs12519770. The G allele of the top genotyped SNP, rs6889138, located in intron 74 of GPR98 and had a minor allele frequency (MAF) of 0.30 in TOF cases, and 0.21 in controls giving an odds ratio (OR) of 1.68 ($P=1.72\times10^{-7}$) per copy in the 22q11.2DS cohort. There were 14 SNPs in the second group that had suggestive association with TOF and were in modest LD with SNP, rs12519770 ($D'=0.84$, $r^2=0.02$). The top SNP, rs6893710, is located in intron 47 of GPR98 and had a MAF of 0.058 in TOF cases, but 0.015 in controls, giving an OR of 4.05 ($P=1.04\times10^{-8}$) per copy of the C allele in the 22q11.2DS cohort (Table 2). C, LocusZoom plot of region of association at rs12519770 on 5q14.3 indicating $-\log_{10} P$ values (y axis) against the chromosomal positions of SNPs (x axis). The genotyped (Continued)
containing 3), maps downstream of GPR98, but it is in a different TAD (Figure 3). The same domain structure marked by TAD triangles occurred in most of the cell lines that were examined (Figure IV in the Data Supplement).

We next determined whether any of the genes are expressed in the pharyngeal apparatus or heart in embryos. Probes were generated, and in situ hybridization was successfully performed for Gpr98, Cetn3, Lysmd3, and Mef2c in mouse embryos at E9.5 and E10.5, when the cardiac OFT is expanding (Figure 3). Although Gpr98 is weakly expressed at E9.5, it is strongly expressed in the neural tube region, particularly the hindbrain\(^2\) (Figure 3). Cetn3, important in the cilia\(^3\) for centrosome reproduction,\(^4\) and Lysmd3, of unknown function, are ubiquitously expressed, although Cetn3 has lower expression levels in the heart itself (Figure 3). Mef2c encodes a MADS box transcription factor, and it is expressed in the pharyngeal apparatus including the second heart field mesoderm,\(^3\) as indicated in Figure 3. Among the genes in the TAD, MEF2C is the only one specifically expressed in cardiac progenitor cells known to be required for heart development.\(^6\)

The second heart field mesodermal progenitor cell populations forming the cardiac OFT lie within the distal pharyngeal apparatus. We then examined expression levels of genes in the 5q14.3 region in existing Affymetrix microarray data from the microdissected distal pharyngeal apparatus.\(^12\) The purpose was to determine whether any of the genes are expressed in this critical tissue. We compared expression levels of genes on 5q14.3 to the highest expressed genes (Actb, Gapdh, Tbx1 expression, and the lowest expressed genes (Il6, Olfr299) in this tissue as shown in Figure 3C. All of the genes are expressed in the pharyngeal apparatus, albeit Gpr98 is expressed at the lowest level, as can be also seen in Figure 3B.

The Encyclopedia of DNA Elements functional genomics data were examined in the 104.7 kb region of LD with SNP, rs12519770, to identify possible regulatory regions. We found 3 possible regulatory regions defined by binding of multiple transcription factors, with one close to rs12519770 (Figure V in the Data Supplement). None of the SNPs that were genotyped in our study lie within these putative regulatory regions; however, critical embryonic regulatory regions could be different, and they have not yet been defined.

### Discussion

We identified genome-wide significant associations between TOF and several SNPs in an intron of GPR98 in our 22q11.2DS cohort. We narrowed the associated region to a 104.7 kb interval. This interval may harbor functional variants that are in LD with the associated SNPs or noncoding variants that regulate the expression of genes within a broad TAD on 5q14.3.

GPR98 contains 90 exons, spans over 610 kb, and encodes a member of the adhesion-G protein-coupled receptor family of receptors. The GPR98 protein binds calcium and is weakly expressed early in mouse embryonic development at E9.5, but it becomes more strongly expressed in the future brain and neural tube by E10.5. It is the largest of the 7-transmembrane receptors and has important functions in hearing and vision.\(^37,38\) Recessive mutations cause Usher syndrome type 2C (Mendelian Inheritance in Man No. 605472), which is characterized by congenital hearing loss and progressive retinitis pigmentosa.\(^39\) There are multiple splice variants present in GPR98, and it is not known if all isoforms have similar functions. Thus, it is possible that ≥1 splice variants could have a function in neural crest cells deriving from the neural tube. Neural crest cells are a migratory population of progenitor cells in the pharyngeal apparatus, which contribute to cardiac OFT septation. There are no reports of a possible function of GPR98 in the cardiovascular system and no known connections to human TOF.

CETN3 is another gene of note in the 5q14.3 region because it encodes a centrin protein that functions in the cytoskeleton of centrosomes and cilia.\(^40\) Cilia are critically important for conferring left right asymmetry during embryonic development and when disrupted is associated with human cardiac anomalies.\(^25\) Because laterality defects are not commonly found in association with 22q11.2DS, more work would need

### Table 2. Top SNPs in GPR98 Associated With TOF

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<thead>
<tr>
<th>SNP</th>
<th>Chr:Position (bp)</th>
<th>Cluster</th>
<th>Risk Allele</th>
<th>Other Allele</th>
<th>RAF</th>
<th>OR (95% CI)</th>
<th>P Value</th>
<th>Conditional P Value*</th>
<th>Conditional P Value†</th>
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<tbody>
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<td>5:90073277</td>
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<td>A</td>
<td>G</td>
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<td>1.69 (1.39–2.06)</td>
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<td>A</td>
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<td>1.68 (1.35–2.10)</td>
<td>1.72E-07</td>
<td>0.00199523</td>
<td>NA</td>
</tr>
<tr>
<td>rs6893710</td>
<td>5:90086041</td>
<td>3</td>
<td>C</td>
<td>T</td>
<td>0.03</td>
<td>4.05 (2.27–7.24)</td>
<td>1.04E-06</td>
<td>3.92E-05</td>
<td>6.11E-05</td>
</tr>
</tbody>
</table>

The top genotyped single-nucleotide polymorphisms (SNPs) in each cluster of SNPs identified for tetralogy of Fallot (TOF) are shown. Chr:Position (bp), chromosome and positions are indicated according to NCBI36/hg18 (March 2006), and allele coding was based on the positive strand. The cluster numbers that the SNPs belong to are indicated. GPR98 indicates G-protein–coupled receptor V1; NA, not associated; OR, odds ratio; and RAF, risk allele frequency in the whole cohort.

*Conditional P value: analysis was conditioned on SNP rs12519770.
†Conditional P value: individually evaluated the association of TOF with 1344 SNPs in the 5q14.3 region (Table VI in the Data Supplement).
to be done to provide additional support of the role of CETN3 as a modifier of TOF in these individuals.

Among the remaining 5 genes (MEF2C, POLR3G, MBLAC2, LYSMD3, and ARRD3C), MEF2C is of particular interest because it encodes a transcription factor required in the second heart field mesoderm of the pharyngeal apparatus during embryogenesis for cardiac OFT development.41,42 Because haploinsufficiency of TBX1 is important for 22q11.2DS, and both MEF2C and TBX1 are expressed in the second heart field progenitor cells, it is possible that TBX1 might act in the same genetic pathway as MEF2C. Further, studies in mouse models indicate that Tbx1 may be a negative regulator of Mef2c.43 This suggests that genetic variants in the 5q14.3 locus that may be associated with MEF2C expression levels could act as genetic modifiers of 22q11.2DS, with the
caveat that causation will require direct experimental support. Further, ISL1 transcription factor (ISL LIM homeobox 1 transcription factor) and GATA (transcription factor) proteins bind and regulate both Mef2c and Nkx2-5 (NK2 homeobox 5 transcription factor) cardiac development genes.44 One hypothesis to test in the future by direct experimentation would be that rare DNA variants in MEF2C, ISL1, or NKK2-5 affect cardiac OFT formation in individuals with 22q11.2DS.

In recent years, higher-order chromatin structure technologies have demonstrated that chromatin interactions occur in a nonrandom manner along the chromosome arms, which are separated into regions of highly interacting chromatin.30,45-47 Relevant to this, the 104.7 kb region found with top associated SNPs to TOF shows a possible regulatory connection with MEF2C located 2 Mb upstream of GPR98 using available Hi-C, chromatin conformation data.29 One limitation of using Hi-C data to draw conclusions about gene regulation is that these data indicate that the 2 genes may reside within the same topological region but do not prove that there is a definitive regulatory connection. Further chromatin conformation data in cell progenitors relevant to cardiac OFT development, followed by direct experimental approaches, will be required to define this interaction further. In addition, a better understanding of the regulatory landscape in this region will be needed to identify the mechanism(s) responsible for the association to TOF we have observed in the 5q14.3 region.

Further support for MEF2C, as a possible modifier gene, comes from a recent GWAS of circulating VEGF (vascular endothelial growth factor) levels in blood in adults.46 Although this is a study of adults, factors that regulate VEGF levels may be similar throughout life and could affect fetal development. Previously, it was found that absence of one of the VEGF isoforms causes a phenocopy of 22q11.2DS in mouse models.49 In this recent GWAS of VEGF levels in adults, MEF2C and JMJD1C (Jumonji domain containing 1C) were found among the 6 loci with significant association to VEGF levels. JMJD1C is relevant because we previously found significant enrichment of rare predicted exonic variants in JMJD1C in whole-exome sequence from 184 22q11.2DS subjects in which the cases were enriched for TOF.50 This provides a potential biological connection between common and previous rare variant analyses for 22q11.2DS. However, in regards to the general population, neither haploinsufficiency nor mutation of MEF2C in humans has been associated with any type of CHD thus far.51-55 Further proof that MEF2C is a cardiac disease gene in the general population will require future genetic studies.

The number of samples we obtained, even after 25 years of collection, is quite small for a GWAS of a complex trait. Thus, one of the limitations of the study was the lack of a true replication cohort. Another limitation of our study is possible population stratification. The majority of our cohort was self-reported as white. Nevertheless, a subset had different ethnicities requiring statistical correction in the analysis. Further, we examined each ethnicity separately and combined each group by performing a meta-analysis. We found that the top SNP in GPR98 was still statistically significant. In addition, the odds ratio was in the same direction in all the subcohorts, supporting our findings despite the limitations.

In this report, we used available bioinformatic data to interpret our data. However, this study did not provide proof of causation. This will need to be done by performing functional studies including development of animal models. Despite the limitations of the study, this is the first GWAS to identify common variants that may modify the cardiac phenotype in a large cohort of individuals with 22q11.2DS.

Conclusions

A GWAS of TOF in 22q11.2DS has identified a significant locus on 5q14.3 harboring potential genetic risk factors. Several genes reside in this locus including MEF2C that is a known gene for cardiac OFT development in animal models. Further work needs to be done to ascertain whether MEF2C or other genes in this region act as modifiers of TOF in 22q11.2DS.

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Appendix

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Disclosures
None.

References
The 22q11.2 deletion syndrome (DiGeorge syndrome/velocardiofacial syndrome) occurs in 1 of 4000 live births, and 60% to 70% of affected individuals have congenital heart disease, ranging from mild to severe. In our cohort of 1472 subjects with 22q11.2 deletion syndrome, a total of 62% (n=906) have congenital heart disease and 36% (n=326) of these have tetralogy of Fallot, comprising the largest subset of severe congenital heart disease in the cohort. One of the main questions clinicians are interested in understanding is the basis for variable phenotypic expression. To address this question, using 22q11.2 deletion syndrome as a model, we performed a genome-wide association study and found tetralogy of Fallot was significantly associated with a genetic interval on chromosome 5q14.3. The associated region is within an intron of GPR98. This intron may contain regulatory sequences that could effect expression of GPR98 or other genes in the region. Among them, MEF2C (Myocyte-specific enhancer factor 2C) is the only gene that is known to affect heart development. On the basis of this work, common DNA variants on 5q14.3 may explain, in part, why phenotypic variation occurs despite the fact that they have the same deletion. Further work will need to be done to determine whether the same genetic variants can alter MEF2C expression and risk for congenital heart disease in the general population or alter clinical outcomes. In the future, understanding how genetic variations influence phenotype in both genetic syndromes and in more common disease will make it possible to improve genetic counseling, rehabilitation, clinical outcomes, and in the creation of novel therapeutics.