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Common variants at 7p21 are associated with frontotemporal lobar degeneration with TDP-43 inclusions

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Competing Financial Interest

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Author Contributions

VVD, PS, MML, and ACP contributed equally to this manuscript. The overall study was designed and implemented by VMYL, JQT, VVD, HH and MML and discussed with GDS, LSW, and ACP. Coordination, sample handling, DNA extraction, genetic analysis and data management were done primarily by VVD and MML. Genotyping and additional testing, including immunohistochemical analysis, was done by PS, HH, VVD, or MML. Data analysis and quality control were performed by PS, LSW, and ACP and discussed and reviewed by VVD, HH, GS, VMYL, JQT, and MML. Expression analysis was designed and implemented by ACP. The manuscript was prepared by ACP, VVD, and PS and reviewed by MML, GDS, HH, JQT, VMYL, and LSW. The members of the International FTLD Collaboration, which includes all other authors, contributed cases, evaluated pathology, performed genetic studies and reviewed the manuscript.

A patent application on TMEM106B has been submitted. Dr. James Lah is currently involved in clinical trials involving: Elan, Janssen, Medivation, and Ceregene. Dr. Howard Feldman has been a full time employee in Neuroscience Global Clinical Research and Development at Briston-Myers Squibb.

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Abstract

Frontotemporal lobar degeneration (FTLD) is the second most common cause of presenile dementia. The predominant neuropathology is FTLD with TAR DNA binding protein (TDP-43)

inclusions (FTLD-TDP)1. FTLD-TDP is frequently familial resulting from progranulin (*GRN*) mutations. We assembled an international collaboration to identify susceptibility loci for FTLD-TDP, using genome-wide association (GWA). We found that FTLD-TDP associates with multiple SNPs mapping to a single linkage disequilibrium (LD) block on 7p21 that contains *TMEM106B* in a GWA study (GWAS) on 515 FTLD-TDP cases. Three SNPs retained genome-wide significance following Bonferroni correction; top SNP rs1990622 (P=1.08×10⁻¹¹; odds ratio (OR) minor allele (C) 0.61, 95% CI 0.53-0.71). The association replicated in 89 FTLD-TDP cases (rs1990622; P=2×10⁻⁴). *TMEM106B* variants may confer risk by increasing *TMEM106B* expression. *TMEM106B* variants also contribute to genetic risk for FTLD-TDP in patients with *GRN* mutations. Our data implicate *TMEM106B* as a strong risk factor for FTLD-TDP suggesting an underlying pathogenic mechanism.

FTLD manifests clinically with progressive behavioral and/or language deficits with a prevalence of 3.5-15/100,000 in 45 to 64 year olds2⁻⁵. The clinical presentation of cases with FTLD pathology varies depending on the referral base6 and among these cases, ~50% are diagnosed as FTLD-TDP1. A family history of a similar neurodegenerative disease may be present in up to 50% of FTLD cases, supporting the existence of a genetic predisposition7. Autosomal dominant *GRN* mutations occur in ~20% of FTLD-TDP cases⁸-11. *GRN* mutations are loss-of-function mutations with most resulting in premature termination of the mutant transcript invoking nonsense-mediated RNA decay and with the ensuing haploinsufficiency causing disease11^{,12}. However, a substantial number of familial FTLD-TDP cases are not explained by *GRN* mutations. Further, patients with the same *GRN* mutation show variable clinical phenotypes or ages of disease onset which likely reflect additional genetic and environmental factors13.

The GWA phase of the study included 515 cases of FTLD-TDP and 2509 disease-free population controls genotyped on the Illumina HH550 or 610-Quad BeadChips as described14 (Table 1). A large population control cohort was acceptable since the general population incidence of FTLD is low4^{,5},15. Cases were obtained under institutional review board approval by members of the International FTLD Collaboration consisting of investigators from 45 clinical centers and brain banks representing 11 countries (United States, Canada, United Kingdom, The Netherlands, Belgium, Spain, Germany, Australia, Finland, France, and Sweden). All cases met either pathological (n=499) or genetic (n=16) criteria for FTLD-TDP which was confirmed by detecting TDP-43 inclusions using immunohistochemistry (IHC)1^{,16}. A genetic criterion for inclusion (*i.e.* presence of a known pathogenic GRN mutation) was used since GRN mutation cases are always diagnosed as FTLD-TDP 8'9'17'18. All cases were checked for relatedness using identity by state (IBS). The results confirmed that although some GRN-associated FTLD-TDP cases share the same mutation on chromosome 17 with similarity in the immediate vicinity of GRN, they are no more related in the remainder of the genome than individuals without GRN mutations. Detailed inclusion criteria are provided in Methods; cohort features are in Supplementary Table 1.

Cochran-Armitage trend test statistics were calculated at all markers following quality control filtering. In addition to self-reported ancestry, all cases and controls were initially screened at ancestry informative markers (AIM) using the STRUCTURE software package19 to reduce the risk of population stratification from self-reported ancestry alone. Each case was subsequently matched to four controls by 'genetic matching' by smartPCA20 as previously described21. The genomic inflation factor (λ) for this study was 1.05 indicating that background stratification was minimal as demonstrated in the quantile-quantile (Q-Q) plots (Supplementary Fig. 1).

Three SNPs reached genome-wide significance following Bonferroni correction (Figures 1a and b). All three SNPs (rs6966915, rs1020004, and rs1990622) mapped to a 68 kb interval (Supplementary Fig. 2) on 7p21.3 (top marker, rs1990622, minor allele frequency (MAF) 32.1% in cases and 43.6% in controls, OR = 0.61, [95% CI 0.53 - 0.71], *P*= 1.08×10^{-11}). For rs1990622, the more common (T) allele confers risk with an OR of 1.64 [95% CI 1.34-2.00]. The interval contained nine additional markers in strong LD ($r^2 > 0.45$) that were also associated with FTLD-TDP (*P*-value range = $8.9 \times 10^{-3} - 7.5 \times 10^{-7}$; OR range 0.63-0.77) (Table 2). All 12 associated SNPs map to a single LD block spanning *TMEM106B*, which encodes an uncharacterized transmembrane protein of 274 amino acids (Figures 1b and c). SNPs rs1020004 and rs6966915 lie within introns 3 and 5, respectively, of *TMEM106B*, while rs1990622 is 6.9 kb downstream of the gene. These findings argue strongly for the association of the 7p21 locus, and the gene *TMEM106B*, with FTLD-TDP.

The association with FTLD-TDP in the GWA was replicated by TaqMan SNP genotyping in 89 independent FTLD-TDP cases and 553 Caucasian control samples at two of the genomewide significant SNPs (rs1020004 and rs1990622) (Table 1 and Supplementary Table 1). A polymorphic variation adjacent to rs6966915 interfered with interpretation of TaqMan genotyping therefore precluding its use in the replication. The replication set was selected based on the same pathological criteria and had similar characteristics as the GWA phase cohort (Supplementary Table 1). In this replication cohort, the top SNPs again showed significant association (P=0.004 for rs1020004 and P=0.0002 for rs1990622) with the same directions of association as those found in the GWA phase (Supplementary Table 1). These results suggest that in the 7p21 locus, encompassing the gene TMEM106B, we have identified a common genetic susceptibility factor for FTLD-TDP. Of interest, this association was not confirmed in a cohort of 192 living patients with unselected FTLD (Supplementary Table 2). This likely reflects heterogeneity in neuropathological substrates underlying FTLD, with only ~50% of unselected clinical FTLD cases expected to have FTLD-TDP. Assuming that TMEM106B genetic variants confer risk of FTLD-TDP specifically, the power to detect this association in 192 clinical FTLD cases and 553 controls is ~30% for an alpha-value of 0.05. To have >90% power to detect this association, a clinical FTLD cohort would require more than 1400 clinical FTLD cases and an equal number of controls.

We next evaluated TMEM106B gene expression in different human tissues to identify phenotype-associated differential expression and also any potential genetic regulators of expression. We queried the mRNA-by-SNP browser (http://www.sph.umich.edu/csg/liang/ asthma/, last accessed June 6, 2009), for genetic regulators of TMEM106B expression (eSNPs) in lymphoblastoid cell lines22. The top SNP, rs1990622, was significantly correlated with *TMEM106B* average expression levels (LOD 6.32; $P=6.9\times10^{-8}$), as was SNP rs1020004 (LOD 5.16, P=1.10×10⁻⁶). The risk allele (T) of rs1990622 was associated with a higher level of mRNA expression, indicating that TMEM106B may be under cisacting regulation by either the FTLD-TDP associated SNPs or another SNP(s) in LD with the associated variants. As the expression data in the publicly available database is derived from lymphoblastoid cell lines from normal individuals22, and the diseased organ in FTLD-TDP is brain, we asked if a similar correlation between genotype and expression phenotype for *TMEM106B* is also present in tissue types affected by disease, and in diseased individuals themselves. Accordingly, we used total RNA isolated from FTLD-TDP postmortem brains (n=18) and neurologically normal control brains (n=7) to evaluate TMEM106B expression in frontal cortex, which is severely affected in FTLD-TDP, by quantitative reverse-transcription PCR (QRT-PCR). All RNA samples used were confirmed to be of equivalent high quality as described23 (Supplementary Table 3). For the same individuals for which we obtained expression data, we genotyped SNPs rs1020004 and rs1990622 using allelic discrimination assays.

Corroborating results from the cell lines, expression of *TMEM106B* was significantly correlated with *TMEM106B* genotype, with risk allele carriers showing higher expression (overall P=0.027, TT vs. TC P=0.017, TT vs. CC P=0.03, for rs1990622, Fig. 2a and Supplementary Fig. 3a). Strikingly, however, expression of *TMEM106B* was >2.5 times higher in FTLD-TDP cases compared to normal controls (P=0.045, Fig. 2b). In addition, the effects of genotype and *TMEM106B* expression on risk of developing disease are at least partly independent, as are the effects of genotype and disease status on *TMEM106B* expression (Supplementary Table 4a and b). Thus, these data suggest that increased *TMEM106B* brain expression might be linked to mechanisms of disease in FTLD-TDP, and that risk alleles at *TMEM106B* confer genetic susceptibility by increasing gene expression.

The primary criterion for inclusion in the GWAS was a neuropathological diagnosis of FTLD-TDP; therefore we studied all cases together regardless of *GRN* mutation status. Nevertheless, *a priori* it was difficult to predict whether additional genetic susceptibility loci would be identified in a group with Mendelian inheritance of highly penetrant mutations. We therefore separately evaluated FTLD-TDP cases with (n=89) and without (n=426) *GRN* mutations. Association to the 7p21 locus persisted in both the *GRN* negative and positive clusters and there was no significant heterogeneity in the ORs for the disease/SNP association between the clusters (Fig. 3 and Supplementary Tables 2 and 5). Using family history status as a covariate in a logistic regression showed that the 7p21 association is independent of family history (Supplementary Table 6). Thus, *TMEM106B* variants may act as a modifier locus in the presence of *GRN* mutations, as the *APOE* locus has been shown to modify age of onset in patients with *PSEN12*4 or *PSEN225* mutations.

Additionally, in the whole GWA cohort, we observed a correlation between rs1020004 genotype and disease duration (P=0.03) with homozygotes for the risk allele (AA, wild-type) having shorter duration of disease (i.e. more severe disease) than individuals homozygous for the minor allele (GG, Supplementary Figure 4). These results provide strong confirmatory evidence for association of the 7p21 locus with increased risk for FTLD-TDP in both *GRN* positive and negative cases.

In addition to the 7p21 locus, analysis of the *GRN* cases alone showed highly significant association with SNPs near the GRN locus on 17q21 (Fig. 3 and Supplementary Table 7). Not unexpectedly, haplotype analysis of the cases indicated that the chromosome 17 association was driven by a shared haplotype among the p.R493X (NM_002087.2:c. 1477C>T) mutation carriers which represented 20.2% (18/89) of the GRN mutation cases. To determine if the observed association at the GRN locus was dependent on the association at the TMEM106B locus we carried out a logistic regression analysis conditioning on the most significantly associated SNP at the 7p21 locus, rs1990622, in the patients with GRN mutations. The conditional analysis had no effect on the association at the GRN locus suggesting that the associations with 17q21 and 7p21 are independent. IBS analysis confirms that these individuals are unrelated and therefore the identified association on chromosome 7 cannot be discounted in GRN mutation carriers. Indeed, conditioning on the top SNP at the GRN locus, rs8079488, also had no effect on the TMEM106B association (results not shown). In addition to the 7p21 locus, the GWAS of GRN negative cases showed a trend for association at five other loci (Supplementary Table 8) including a locus on chromosome 9p21.2 that falls within a 7.7 Mb critical interval defined from five previous linkage studies, representing a potential refinement of that region26. We observed no association at the GRN locus in the cases without GRN mutations.

We then evaluated mRNA expression of *TMEM106B* in FTLD-TDP with and without *GRN* mutations separately, and the *GRN* mutants showed increased expression (overall P=0.0009), compared to controls (P=0.0005) and FTLD-TDP without *GRN* mutations

(P=0.002) (Figure 2c). Furthermore, controlling for rs1990622 genotype and focusing on heterozygotes (n=14), the presence of a *GRN* mutation remained significantly associated with increased *TMEM106B* expression (P=0.039, Fig. 2d) compared to normal controls. These results are compatible with a model in which mutations in *GRN* are upstream of increased *TMEM106B* expression in increasing risk for FTLD-TDP.

A mechanistic understanding of the pathogenesis of FTLD has been hampered the heterogeneity in clinical and pathological features. With the discovery of TDP-43 as a major FTLD disease protein, the pathologically-defined entity of FTLD-TDP emerged16. Identification of *GRN* mutations as a major genetic cause of FTLD-TDP, led to definition of a genetic subgroup of FTLD-TDP. This study identifies *TMEM106B* as a genetic risk factor for FTLD-TDP. We speculate that the homogeneous pathologically-defined study population used here enabled us to detect a robust signal with relatively small case numbers.

Our data suggest a potential disease mechanism in which risk-associated polymorphisms at 7p21 increase *TMEM106B* expression, and elevated *TMEM106B* expression increases risk for FTLD-TDP. Additionally, we show that *TMEM106B* genotypes are a significant risk factor for FTLD-TDP even in *GRN* mutation carriers implying that *GRN* mutations may act upstream of *TMEM106B* in a pathogenic cascade. Future directions of research on this novel genetic risk factor will include a detailed evaluation of the *TMEM106B* locus by sequencing, collection of more pathologically-defined FTLD-TDP cases for a genome-wide replication, and studies of expression profiles in additional tissues and brain regions. A better understanding of this gene may in turn provide an opportunity to intervene in an otherwise fatal and devastating neurodegenerative disease.

METHODS

Inclusion criteria

Individuals of European descent with dementia clinically +/- motor neuron disease (MND) and an autopsy diagnosis of FTLD-TDP confirmed by TDP-43 IHC were included. Mixed pathologies were not excluded. Living individuals with a pathogenic *GRN* mutation were also included18. Only a single proband per family was permitted. Appropriate informed consent was obtained. 598 unique FTLD-TDP cases met inclusion criteria; 515 were used for the GWAS after PCA matching to controls. Characteristics described in Supplementary Table 1. Whole genome amplification (WGA), performed in duplicate and pooled, was used for 15 (Repli-g Mini, Qiagen), but only 6 cases ultimately passed quality control parameters for the GWAS. The replication, using SNP genotyping, included cases of insufficient quality or quantity for the GWA phase (n=27), cases available only as formalin-fixed paraffinembedded tissue (n=6), and cases randomly not used for GWA phase (n=56). Three FTLD-TDP cases with mutations in valosin-containing protein (VCP) gene were included (two in GWA and one in replication)18.

Controls

GWAS controls consisted of 2509 samples, including 1297 self-reported Caucasian children of European ancestry recruited from CHOP Health Care Network and 1212 samples from the 1958 birth cohort genotyped by the WTCCC27. Although the controls were not selected for absence of neurodegenerative disease, the large size of the cohort relative to the low population frequency of FTLD overrides this potential concern. Furthermore, the minor allele frequencies at the 7p21 loci are very similar (<1-2% variation) between CHOP and WTCCC cohorts suggesting they accurately reflect the control allelic frequencies in the general population (Supplementary Table 9). To reduce the risk of population stratification all internal controls were screened using the STRUCTURE package19 at 220 AIMs. To

improve clustering the samples were spiked with 90 CEPH, Yoruban and Chinese/Japanese individuals genotyped as part of the HapMap project. Cases were excluded if their inferred proportion of ancestry was less than 90% that of the CEU cluster.

For the replication 553 controls were as follows: 275 from Coriell Institute (Neurologically Normal Caucasian control panels, Camden, NJ), 155 clinical controls from neurology clinics at University of Pennsylvania (UPenn), 28 brain samples of neurologically normal individuals > 60 years from the UPenn Center for Neurodegenerative Disease Research (CNDR), and 95 population controls from CHOP.

DNA extraction and quality assessment

Samples sent as DNA from external sites were extracted using different methods. Remaining samples (376) were extracted at UPenn from frozen brain tissue or blood. Genomic DNA was extracted from frozen brain tissue (50 mg) by the Qiagen MagAttract DNA Mini M48 Kit on the M48 BioRobot. Genomic DNA was purified from whole blood using FlexiGene kit (Qiagen). High quality DNA was required for the Illumina genotyping. All DNA samples were evaluated for purity by spectrophotometric analysis (Nanodrop) and for degradation by 1% agarose gel electrophoresis (Invitrogen).

TDP-43 IHC

Autopsy cases were confirmed to have TDP-43 pathology by IHC performed by the sending institution or at UPenn CNDR as previously described16. TDP-43 negative cases were excluded.

GRN sequencing

To stratify the analysis according to *GRN* mutation status, exons 1-13 (with exon 1 representing exon 0 in Gass *et al.*10) and adjacent intronic regions were sequenced as described13 in cases not previously evaluated. *GRN* sequencing was not possible due to limited sample quantity in a few cases (n=13 in GWA, n=15 in replication). Novel variants identified in this study not predicted to cause a frameshift or premature termination and previously described variants of uncertain significance were grouped with *GRN* mutation negative cases. The most common mutations identified are given in Supplementary Table 1.

Illumina genotyping and quality control

The FTLD-TDP cases and CHOP control samples were genotyped on either the Illumina HH550 BeadChip or the Illumina human610-quad BeadChip at the Center for Applied Genomics at CHOP as previously described14. The 1958 birth cohort samples were genotyped on the HH550 BeadChip by the WTCCC27. Sixteen individuals, 13 cases and 3 controls, were excluded from GWA phase for low genotyping (<98% chip-wide genotyping success). We further rejected 13,316 SNPs with call rates <95%, 23,552 SNPs with MAF < 1% and 1,940 SNPs with Hardy Weinberg equilibrium *P*<10⁻⁵ in the controls samples; the λ was 1.05. Cases and controls were screened for relatedness using the IBS estimations in *plink* (http://pngu.mgh.harvard.edu/~purcell/plink/index.shtml) on 100,000 randomly distributed markers throughout the genome. Pairwise Pi-hat values in excess of 0.01 were indicative of relatedness.

Following the quality control measures cases were matched to controls by 'genetic matching' as previously described21. We computed principal components for our dataset by running smartpca, a part of the EIGENSTRAT package, on 100,000 random autosomal SNPs and applied a matching algorithm implemented in MATLAB to the output. The matching algorithm assigns each sample a coordinate based on k eigenvalue-scaled principal components. It then matches each case to m unique controls within a distance d, keeping

only cases that match exactly m controls. The distance thresholds were manually optimized to minimize λ and maximize power (i.e. number of cases). We matched each case to four controls, using the first three principal components and a distance threshold of 0.025.

Statistical Analysis for Association

Statistical tests for association were performed using *plink*. Single marker analyses for the genome-wide data were done using the Cochran-Armitage trend test. The genomic inflation factors were 1.05 for the complete case set and 1.03 for the GRN mutation carriers, indicating only minor background stratification. The Breslow-Day test within *plink* was used to test for heterogeneity of odds ratio for the disease/SNP association between GRN mutation carriers and non-carriers. Conditional SNP regression analyses were completed in *plink*, the allele dosages of the conditioning SNP were included as covariates in the logistic regression models. To determine if the association at the TMEM106B locus was dependent on family history we included family history status as a covariate in a logistic regression model using *plink*. Haplotypes were reconstructed and population frequencies estimated using the EM algorithm implemented in the program fastPHASE28. For the age of onset and disease duration analyses we performed an analysis of variance (ANOVA) with the general linear models procedure in R (www.r-project.org). Independent variables for each ANOVA were the log transformed age of onset or disease duration in years and the individual SNP genotype with additive encoding (ie three categories where 0 is homozygous for the ancestral allele, 1 is heterozygous and 2 is homozygous for the minor allele). Power calculations were based on the rs1990622 allele frequencies observed for cases and controls in the GWAS, using a two-tailed test. We assumed that clinical FTLD cases without TDP-43 pathology as the neuropathological substrate would have allele frequencies similar to controls.

SNP Genotyping for Replication

For the replication, genotyping was performed using TaqMan chemistry-based allelic discrimination assays (Applied Biosystems (ABI), Foster City, CA) on the ABI 7500 Fast Real-Time System followed by analysis with SDS 7500 software v2.0.1. The ABI assays used were: rs1020004, C_7604953_10 and rs1990622, C_11171598_10. A nearby novel genetic variation (possible deletion) was found to interfere with correct genotyping of the T allele of SNP rs6966915 using ABI reagents C_31573289_10 (as well as by DNA sequencing), thus this SNP was not used further.

Human samples for expression analysis

Frontal cortex human brain samples from the CNDR Brain Bank characterized following consensus criteria1^{,3} were dissected as previously described23. Neurologically normal controls (n=7), FTLD-TDP cases with (n=8), and without (n=10) *GRN* mutations were sampled (Supplementary Table 10). *GRN* mutations were confirmed to be absent from control cases. RNA quality was verified using an Agilent 2100 Bioanalyzer (RIN>6 for inclusion) as previously described23. mRNA expression was quantified by QRT-PCR on the ABI7500 using the delta-delta C_T method, and the geometric mean of two housekeeping genes (β -actin and Cyclophilin A), shown to have stable expression in frontal cortex samples from FTLD-TDP and normal individuals23. Detailed information on primers is available on request.

Statistical analyses of expression data and replication cohort

For all brain expression and replication cohort analyses, statistical tests were performed using open source R software packages. R-scripts are available upon request. For evaluations of the effect of disease status, SNP genotype, and gender on *TMEM106B*

expression, linear regressions were used to compute p-values in univariate models. We evaluated assumptions of linearity by checking QQ plots (observed vs. predicted under normal distribution). For pairwise comparisons within the linear models, risk allele homozygotes and *GRN* mutants, respectively, were designated the reference group for marginal t-tests evaluating genotype effects and the effects of *GRN* mutations on expression. Normalized gene expression sample genotype and gender data are provided in Supplementary Data 1 and 2. For evaluations of the independent contributory effects of SNP genotype and *TMEM106B* expression on disease state, logistic regressions were used to compute AIC values in multivariate vs. univariate models (Supplementary Table 4a). For evaluations of the independent contributory effects of SNP genotype and disease state on *TMEM106B* expression, linear regressions were used in multivariate vs. univariate models (Supplementary Table 4b). For analyses of association of SNP genotypes with disease in our TaqMan replication cohort, Cochran-Armitage trend tests were used to compute *P*-values under a codominant model.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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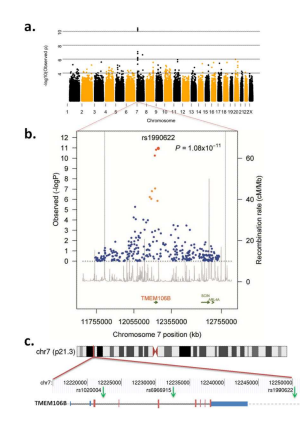


Figure 1. Region of genome-wide association at 7p21

a. Manhattan plot of $-\log 10$ (observed *P*-value) across genome demonstrating region of genome-wide significant association on chromosome 7; **b.** Regional plot of the *TMEM106B* associated interval. **Foreground plot:** Scatter plot of the $-\log_{10} P$ -values plotted against physical position (NCBI build 36). **Background Plot:** Estimated recombination rates (from phase 2 of the HapMap) plotted to reflect the local LD structure. The color of the dots represents the strength of LD between the top SNP rs1990622, and its proxies (red: r2 0.8; orange 0.8 < r2 0.4; blue < 0.4). Gene annotations were obtained from assembly 18 of the UCSC genome browser; **c.** Location of 3 highest associated SNPs (green arrows) relative to the gene structure of *TMEM106B* (blue bars, 3' and 5'-untranslated regions; larger red bars, coding exons; thick gray line, intronic regions; gray dashed line, downstream chromosome sequence) and chromosome 7 location.

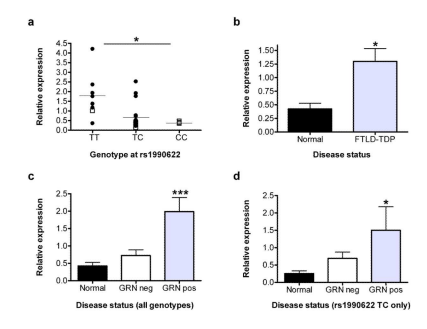


Figure 2. TMEM106B expression variation by genotype and disease state

a. TMEM106B mRNA expression by QRT-PCR in frontal cortex differed significantly by genotype at rs1990622 (overall P=0.027, genotype TT vs. TC P=0.017, TT vs. CC P=0.03). Black circles, FTLD-TDP (n=18); open squares, normal (n=7); horizontal lines, group mean. Significance of P-values are denoted by the numbers of asterisks. b. TMEM106B mRNA expression in frontal cortex was significantly higher in samples from FTLD-TDP patients compared to normal controls (P=0.045). c. TMEM106B expression in frontal cortex samples in FTLD-TDP with (GRN pos, n=8) or without (GRN neg, n=10) GRN mutations compared to normals (n=7). GRN mutation carriers had significantly higher levels of TMEM106B expression (overall P=0.0009, GRN pos vs. controls P=0.0005, GRN pos vs. GRN neg P =0.002). **d.** When only cases heterozygous at rs1990622 (n=14) were evaluated, GRNmutations remained significantly associated with a higher level of TMEM106B expression (P=0.039) in frontal cortex. QRT-PCR was performed in triplicate for all expression studies. Expression values were normalized to the geometric mean of two housekeeping genes and are shown relative to a single reference normal control sample23. Error bars represent the standard error of the mean. Normalized gene expression data and sample genotype and gender data used for these analyses are provided online in Supplementary Material.

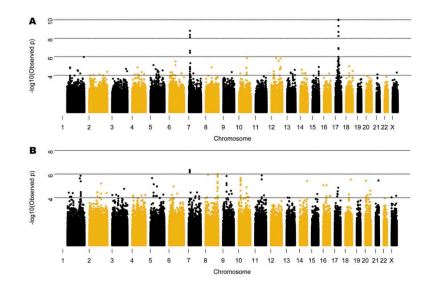


Figure 3. Manhattan plot in cases with and without *GRN* mutations

Manhattan plot of $-\log 10$ (observed *P*-value) across genome in cases with (a) and without (b) GRN mutations. The subset of cases with *GRN* mutations demonstrates regions of genomewide significant association on chromosomes 7 and 17. The chr 17 association is confirmed to be driven by a shared haplotype in c.1477C>T (p.R493X) *GRN* mutation carriers representing ~20% of mutation positive cases, however the chromosome 7 association is not related to any single *GRN* mutation and remains when the cases with c.1477C>T are removed (*P*=1.446×10⁻¹⁰). The same locus on chr 7 identified in the *GRN* mutation cases is also the strongest signal in the *GRN* negative cases, although it does not reach genome-wide significance. A list of the SNPs with the highest signals in b is given in Supplementary Table 8.

Table 1

Summary of samples and controls used for GWA and replication phases

Phase	Case Numbers	Case Study Source	Control Numbers	Control Study Source	Method of Testing	λ
GWA	515	International FTLD Consortium	2509	1297 CHOP European- Caucasian, 1212 WTCCC	Illumina HH550 or 610-Quad BeadChips	1.05
Replication	89	International FTLD Consortium	553	3 Penn Autopsy, Penn ADC, Coriell Neurologically Normal panel, CHOP European-Caucasian 2 SNPs		

CHOP, Children's Hospital of Philadelphia; Penn ADC, University of Pennsylvania Alzheimer's Disease Center; WTCCC, Wellcome Trust Case Control Consortium; λ , genomic control inflation factor.

Table 2

SNPs on chromosome 7 in region with highest association in the GWAS

SNP rs ID	BP	Minor Allele	MAF case	MAF cont	CA P-val	OR	Lower 95% CI	Upper 95% CI
rs1006869	12071795	G	0.148	0.184	5.82×10^{-3}	0.77	0.64	0.92
rs1990602	12088321	G	0.133	0.166	8.97×10 ⁻³	0.77	0.64	0.94
rs10226395	12101859	С	0.154	0.191	4.90×10 ⁻³	0.77	0.64	0.93
rs1003433	12130625	G	0.287	0.351	9.45×10 ⁻⁵	0.75	0.64	0.86
rs6952272	12166585	Т	0.154	0.207	9.88×10 ⁻⁵	0.70	0.58	0.84
rs12671332	12182087	С	0.173	0.245	7.50×10 ⁻⁷	0.64	0.54	0.77
rs1468915	12194417	С	0.171	0.242	9.49×10 ⁻⁷	0.65	0.54	0.77
rs1020004	12222303	G	0.233	0.338	5.00×10 ⁻¹¹	0.60	0.51	0.70
rs6966915	12232513	Т	0.321	0.435	1.63×10 ⁻¹¹	0.61	0.53	0.71
rs10488192	12243606	Т	0.139	0.204	1.46×10 ⁻⁶	0.63	0.52	0.76
rs1990622	12250312	С	0.321	0.436	1.08×10^{-11}	0.61	0.53	0.71
rs6945902	12252934	А	0.183	0.230	7.17×10 ⁻⁴	0.75	0.63	0.89

SNPs are listed in genomic order based on location on chromosome 7. SNPs in bold text have the lowest P-values.

Chr, chromosome; BP, base pairs (NCBI build 36); MAF, minor allele frequency; cont, controls; CA P-val, Cochrane-Armitage P-value; OR, odds ratio; CI, confidence interval.