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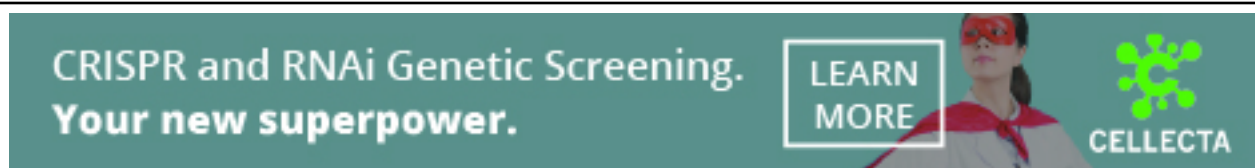
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Method

Whole-genome analysis of noncoding genetic variations identifies multiscale regulatory element perturbations associated with Hirschsprung disease

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It is widely recognized that noncoding genetic variants play important roles in many human diseases, but there are multiple challenges that hinder the identification of functional disease-associated noncoding variants. The number of noncoding variants can be many times that of coding variants; many of them are not functional but in linkage disequilibrium with the functional ones; different variants can have epistatic effects; different variants can affect the same genes or pathways in different individuals; and some variants are related to each other not by affecting the same gene but by affecting the binding of the same upstream regulator. To overcome these difficulties, we propose a novel analysis framework that considers convergent impacts of different genetic variants on protein binding, which provides multiscale information about disease-associated perturbations of regulatory elements, genes, and pathways. Applying it to our whole-genome sequencing data of 918 short-segment Hirschsprung disease patients and matched controls, we identify various novel genes not detected by standard single-variant and region-based tests, functionally centering on neural crest migration and development. Our framework also identifies upstream regulators whose binding is influenced by the noncoding variants. Using human neural crest cells, we confirm cell stage-specific regulatory roles of three top novel regulatory elements on our list, respectively in the *RET*, *RASGEF1A*, and *PIK3C2B* loci. In the *PIK3C2B* regulatory element, we further show that a noncoding variant found only in the patients affects the binding of the gliogenesis regulator NFIA, with a corresponding up-regulation of multiple genes in the same topologically associating domain.

[Supplemental material is available for this article.]

Hirschsprung (HSCR) disease is a rare, complex genetic disease characterized by missing enteric ganglia in various portions of the hindgut (Amiel et al. 2008). It is caused by failed migration, proliferation, differentiation, or colonization of enteric neural crest (NC) cells, which disrupts enteric nervous system (ENS) development (Badner et al. 1990; Chakravarti et al. 2006). Phenotypic severity of the disease is determined by the length of colonic aganglionosis and can be classified into short-segment HSCR (S-HSCR; 80% of cases), long-segment HSCR (L-HSCR; 15%–20% of cases), and total colonic aganglionosis (TCA; up to 5% of cases) (Udassin et al. 1981; Chakravarti et al. 2006).

HSCR is long recognized to be highly heritable (80%–97% for S-HSCR and ~100% for L-HSCR), while around 80% of HSCR cases are sporadic (>95% for S-HSCR) (Badner et al. 1990; Tam et al. 2019). The incidence of HSCR varies across ethnic groups, from 1.4 to 2.8 in every 10,000 new births, with the highest incidence

rate in Asia (Chakravarti et al. 2006; Amiel et al. 2008; Tam and Garcia-Barceló 2009). The genetic etiology of HSCR is multifactorial, involving rare and common, coding and noncoding variants in genes playing different roles in ENS development. Among the different subtypes of HSCR, S-HSCR is genetically most complex. Whereas L-HSCR and TCA cases are mostly autosomal dominant, the less severe S-HSCR subtype follows a complex, non-Mendelian inheritance pattern and has a male-to-female ratio of 4:1 (Badner et al. 1990; Emison et al. 2005), with the reason behind it not fully known.

Damaging rare variants at protein-coding sequences associated with HSCR have been found in many genes involved in ENS development. Among them, *RET*, encoding a receptor tyrosine kinase, is responsible for ~50% of familial cases and 10%–20% of sporadic cases (Chakravarti et al. 2006). Other genes previously reported include *BACE2*, *DNMT3B*, *ECE1*, *EDN3*, *EDNRB*, *FAT3*, *GDNF*, *GFRA1*, *KIFBP*, *L1CAM*, *NRG1*, *NRG3*, *NRTN*, *NTF3*,

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NTRK3, *PHOX2B*, *PROK1*, *PROKR1*, *PROKR2*, *PSPN*, *SEMA3A*, *SEMA3C*, *SEMA3D*, *SOX10*, *TCF4*, and *ZEB2* (Amiel et al. 1996; Garcia-Barcelo et al. 2009; Alves et al. 2013; Luzón-Toro et al. 2015; Tang et al. 2018; Tilghman et al. 2019), which together account for around 5% of all cases. These rare damaging variants often have incomplete penetrance and explain only a small portion of the phenotypic variance.

In contrast, the most well-known common variant, rs2435357, located in intron 1 of *RET* and affecting *SOX10* binding to a *RET* enhancer, explains 10–20 times of the phenotypic variance as compared to the rare damaging *RET* variants (Emison et al. 2005, 2010; Chatterjee et al. 2016). Recently, some additional noncoding variants have also been found associated with HSCR. For example, two other variants in *RET* enhancers act synergistically with rs2435357 on HSCR risk (Chatterjee et al. 2016), while variants in four noncoding elements around the *RET* and *SEMA3* loci were found in 48.4% of cases and 17.1% of controls, with the presence of five or more variants in these regions corresponding to a high disease risk (Tilghman et al. 2019). However, despite the ever-growing knowledge about HSCR, there is still a considerable portion of cases that cannot be explained by the cataloged coding or noncoding variants.

The emerging studies of HSCR-associated common variants in *RET* enhancers and other functional noncoding regions (Emison et al. 2010; Chatterjee et al. 2016; Tilghman et al. 2019) and the abundance of noncoding signals from genome-wide association studies of many human traits in general (Zhang and Lupski 2015) both suggest that noncoding regions would be a valuable territory to explore in the quest to explain the missing heritability in HSCR.

Existing methods for studying noncoding genetic variants can be broadly classified into site-based and region-based methods (Supplemental Materials). In general, both types of methods face several major challenges, namely (1) there is a large number of sites/regions to consider genome-wide, leading to low statistical power due to multiple hypothesis testing correction, (2) many genetic variants at noncoding regions do not have direct functional effects, such as those in linkage disequilibrium with the actual functional variants, (3) epistatic interactions can exist among genetic variants, making it unsuitable to consider each variant separately (Cheng et al. 2017), (4) genetic variants at different regulatory elements of the same gene/pathway may result in the same convergent effect, and therefore each of them may not have high recurrence in the study subjects, and (5) the loci of different genetic variants could be related to each other not by their genomic locations but by the upstream regulators that commonly bind them, which are more nontrivial to determine.

In this study, we propose a new noncoding variant prioritization framework, Multiscale Analysis of Regulatory Variants on the Epigenomic Landscape (MARVEL), that addresses these five challenges. Applying our framework to the whole-genome sequencing (WGS) data generated from an S-HSCR cohort (431 cases and 487 ethnically matched controls), we identify disease-associated enhancers, promoters, and genes based on epigenomic data generated from the human pluripotent stem cell (hPSC)-derived enteric NC-like cells (hNC). This is by far the largest WGS data set of S-HSCR with both cases and controls coming from a single population. By aggregating lower-level (genomic variants and TF binding motifs) information, marginally significant signals emerge as more significant associations at a higher level (enhancers, promoters, and genes) that converge to specific functional pathways and upstream regulators.

Results

A novel association framework for the noncoding genome

MARVEL is designed to address the five challenges listed above (Fig. 1; Methods; Supplemental Materials). First, we use cell type-specific epigenomic data to define a target set of active regulatory elements in the relevant cell types, in order to limit the number of statistical tests to be performed (Fig. 1A). Second, in these target elements, we use sequence motifs as a proxy to evaluate the functional potential of the genetic variants (Korhonen et al. 2017), which helps prioritize the genetic variants and their corresponding regulatory elements among those in linkage disequilibrium (Fig. 1B). Third, for each target element, we consider all genetic variants simultaneously to reconstruct the sample-specific sequence for evaluating their joint functional effects (Fig. 1C). For instance, if two genetic variants have opposite effects on the binding strength of a TF according to its sequence motif, the net expected effect would be smaller than the sum of their individual absolute effects. Fourth, our framework considers multiple occurrences of a motif in the same regulatory element (Fig. 1D) or different regulatory elements of the same gene (Fig. 1E) in a joint manner. As a result, a gene would be considered significantly perturbed when its regulatory elements have frequent motif-disrupting genetic variants even if in different subjects these regulatory elements contain different sets of genetic variants (Corradin et al. 2014; Chatterjee et al. 2016). The joint effect of multiple motifs is further captured by first selecting an important combination of the motifs using a feature selection procedure (Fig. 1F) followed by an association test (Fig. 1G), which together identify motifs whose match scores in these regulatory elements are significantly associated with the target phenotype, adjusted for covariates such as age and sex of the subjects. The identified phenotype-associated regulatory elements and genes are further investigated by additional analyses such as gene set enrichment and single-cell expression analyses (Fig. 1H). Fifth, by using sequence motifs as functional proxy, we also identify the potential upstream regulators whose binding to different target elements is frequently perturbed, which can be either loss or gain of binding. By identifying those whose binding motifs are commonly perturbed in the regulatory elements of different genes (Fig. 1I), we identify both the core regulators involved in the phenotype (Boyle et al. 2017) and genes potentially involved in the same pathways.

Using simulated data (Supplemental Materials), we verified that MARVEL is able to select the simulated informative motifs associated with the phenotype, including motifs whose match scores are affected by genetic variants at different allele frequencies, effect sizes, and correlations among each other (Supplemental Fig. S1). We also verified that, in the *P*-value and effect size distributions produced by MARVEL, the simulated disease-associated regions were clearly separated from the background (Supplemental Fig. S2).

Novel noncoding regulatory elements and gene loci associated with S-HSCR

We applied MARVEL to analyze our S-HSCR WGS data (Supplemental Materials). Briefly, target enhancers and promoters were defined using chromatin accessibility and histone modification data from hNC, which were, respectively, used in an enhancer-based and a promoter-based analysis. A gene-based analysis was also performed by considering at the same time all the promoters and enhancers that potentially regulate each gene, with the motif scores of the different regulatory elements weighted according to their

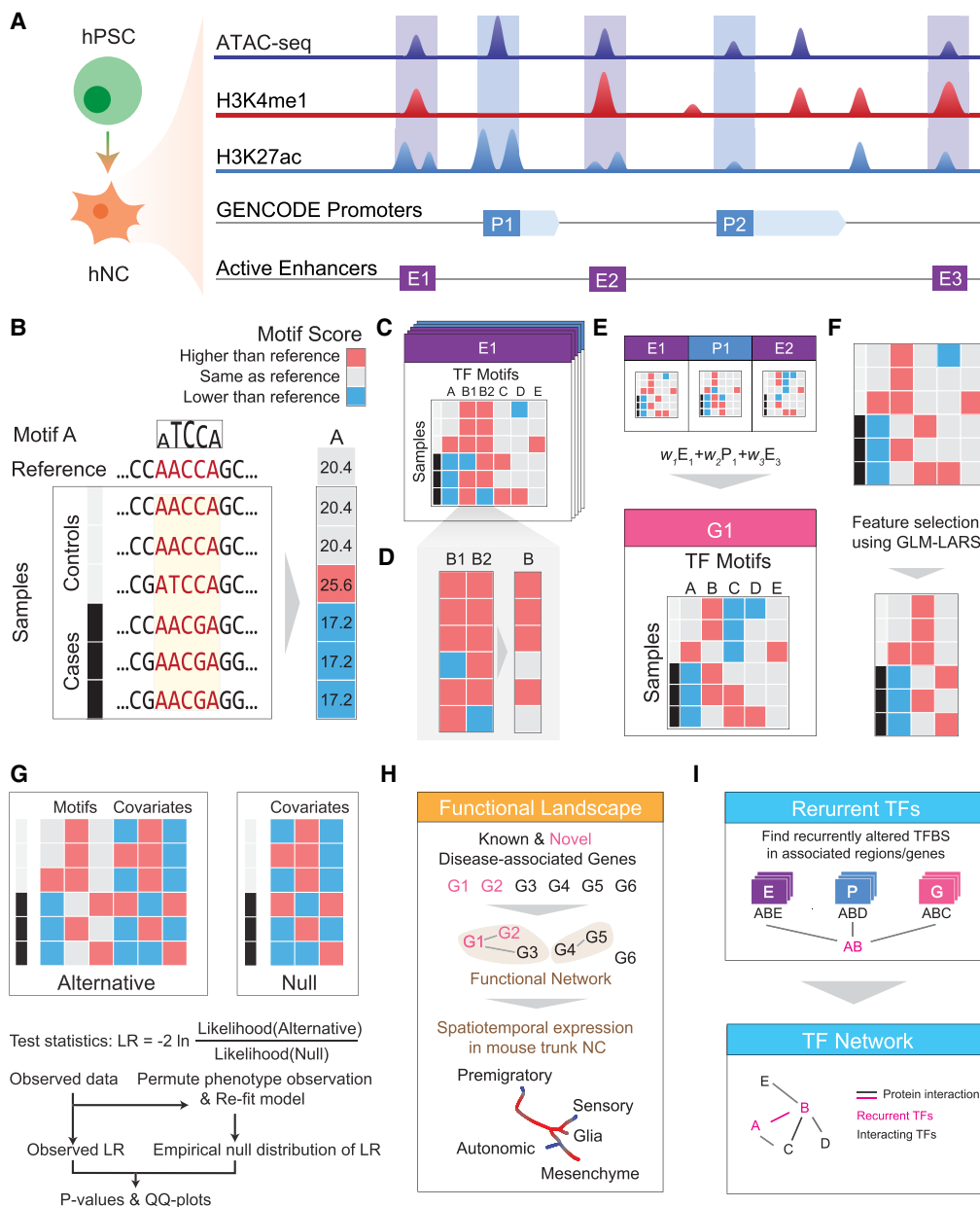


Figure 1. Schematic overview of MARVEL. (A) Epigenomic data of relevant cell types (hNC in the case of HSCR) are integrated with a gene annotation set to identify the active regulatory elements relevant to the phenotype of interest. (B) In each regulatory element, the functional significance of genetic variants is evaluated by their perturbation to TF sequence motifs. (C) Since the perturbation effects of multiple genetic variants may not add up linearly, they are considered together to reconstruct the sample-specific sequences, based on which the overall change of TF motif match scores is determined. (D) For motifs with multiple appearances within the same regulatory element, their match scores are aggregated to give a single score. (E) At a higher level, if a gene involves multiple regulatory elements, the aggregated match scores of a motif in the different elements can be further aggregated into a single score. This is done in the gene-based analysis. (F,G) The aggregated match score matrix of all the motifs for a regulatory element/gene is used as the input of an association test, which selects a subset of the most informative motif features (F) and compares a model involving both these selected features and the covariates with a null model that involves only the covariates using likelihood ratio (LR) test (G). (H) The regulatory elements and genes identified to be significantly associated with the phenotype can be further studied by other downstream analyses, such as gene set enrichment and single-cell expression analyses. (I) TFs with recurrently perturbed match scores in different regulatory elements are collected to infer a network that highlights the phenotype-associated perturbations.

expected chromatin contact frequencies with the transcription start site (TSS) in hPSCs (Supplemental Materials; Supplemental Table S4). This gene-based analysis allowed the detection of genes with frequent perturbations by multiple low-recurrence genetic variants at different regulatory elements. Association tests were

then performed based on the motif scores in each of these three sets of target regions in turn, which contained 150,828 enhancers, 87,461 promoters, and 24,557 genes, respectively.

We defined all regions above the 95% confidence interval of the null in the quantile-quantile plot as loosely associated

with S-HSCR, and all regions passing the FDR threshold of 0.1 (including a subset of the loosely associated regions) as significantly associated (Fig. 2A; Supplemental Table S1). Quantified by the area under the receiver operating characteristic (AUROC) (Methods), the loosely associated regions had generally larger effect sizes than background hNC enhancers, hNC promoters, and genes (Fig. 2B). Comparing the three sets of target regions, there were more enhancers and genes significantly associated with S-HSCR as compared to promoters. Noncoding variant association methods that focus on the promoter regions would have missed the many associated enhancers and aggregated regulatory regions of the genes. As a negative control, we repeated the procedures on a background set of enhancers that do not overlap the hNC enhancers (Methods) and found none of them either significantly or loosely associated with S-HSCR (Supplemental Fig. S3).

As expected, all three analyses identified various regions near the *RET* locus as significantly associated with S-HSCR (Table 1; Supplemental Materials). Moreover, we also identified nine enhancers and four genes significantly associated with S-HSCR that are not close to *RET* (Table 1).

Among the nine enhancers, the one on Chromosome 1 (Chr 1: 204,456,576–204,457,577) overlaps intron 10 of *PIK3C2B*. *PIK3C2B* is a phosphoinositide 3-kinase (PI3K) family protein, which is believed to have important roles in the signal integration and transduction in hNC (Frank and Tsai 2009). Binding sites of *TWIST1* and *NFIA* are altered in this enhancer (Table 1), both with altered match scores in six cases but not in any controls. *TWIST* is an hNC specifier (Simoes-Costa and Bronner 2013) while *NFIA* has been shown to regulate gliogenesis, which is also crucial to a functional ENS (Kang et al. 2012; Glasgow et al. 2014; Tchieu et al. 2019). Another significantly associated enhancer, on

Chromosome 17 (Chr 17: 8,915,658–8,916,659), is also close to two other PI3K family genes, *PIK3R5* and *PIK3R6*. The two enhancers on Chromosomes 18 and 16 (Chr 18: 73,918,014–73,919,015 and Chr 16: 86,954,235–86,955,236) are within/close to F-box genes (*FBXO15* and *FBXO31*, respectively). F-box proteins bind to *CUL1* to form SCF (SKP-CUL1-F-box protein) E3 Ubiquitin Ligase complexes, which mediate ubiquitination of proteins that regulate the cell cycle and diverse neuronal activities (Liao et al. 2004; Frank and Tsai 2009). In particular, *FBXO31* has been shown to regulate neural morphogenesis and migration (Vadhvani et al. 2013). Another enhancer on Chromosome 17 (Chr 17: 958,411–959,412) is close to *ABR*, which encodes a protein that is able to promote *RHO* activity and reduce *CDC42* activity (Vaughan et al. 2011), both of which are important for actin regulation and cell polarization during migration (Vaughan et al. 2011; Cohen et al. 2018). The enhancer on Chromosome 9 (Chr 9: 100,937,755–100,938,756) is close to *PLPPR1*, which encodes a member of the plasticity-related gene (PRG) family. Members of the PRG family mediate lipid phosphate phosphatase activity in neurons and are known to be involved in neuronal plasticity (Savaskan et al. 2004; Trimbuch et al. 2009), potentially also involved in ENS development (Schäfer et al. 2009). The enhancer on Chromosome 11 (Chr 11: 120,020,704–120,021,705) is close to *TRIM29*, a gene that regulates epithelial-to-mesenchymal transition (EMT), which is an important process shared by both NC migration and cancer metastasis (Sun et al. 2019). Finally, the genes closest to the enhancer on Chromosome 22 (Chr 22: 37,128,477–37,129,478) are not clearly related to HSCR, which could represent new findings. In general, many of the top S-HSCR-associated enhancers identified by MARVEL cannot be identified by commonly used single-variant and region-based association (Supplemental Materials; Supplemental Fig. S4).

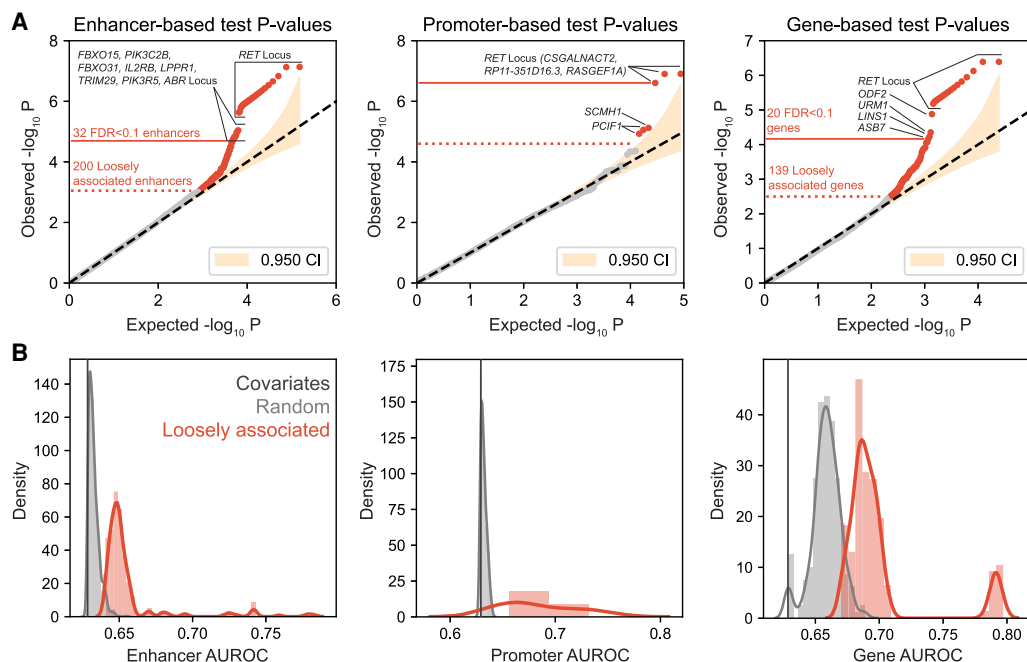


Figure 2. Association test results. (A) Q-Q plots of association P -values in the enhancer-based, promoter-based, and gene-based tests. In each plot, the yellow shaded area shows the 95% confidence interval. The dotted red line marks the demarcation point of the loosely associated regions, above which all the regions are outside the 95% confidence interval. The solid red line marks the threshold for significantly associated regions, above which all regions have an FDR Q -value < 0.1 . The significantly associated regions are a subset of the loosely associated regions. (B) Comparison of the AUROC value distributions of the models of the loosely associated regions with those of the models of the background regions and the covariate-only model.

Table 1. Significant S-HSCR-associated enhancers, promoters, and genes

Region	<i>P</i> -value(s)	Altered motifs of TFs expressed in hNC	
		Gain	Loss
Direction of binding change associated with increased disease risk:			
Significantly associated enhancers (and three corresponding nearest genes of each)			
23 enhancers near <i>RET</i>	7.42×10^{-8a} – 2.35×10^{-6}		
Chr 18: 73,918,014–73,919,015 (<i>FBXO15</i> , <i>TIMM21</i> , <i>CYB5A</i>)	9.10×10^{-6}	CPEB1, NFATC1, NFIC, ZNF322	LEF1, SP3, TBX1, WT1
Chr 1: 204,456,576–204,457,577 (<i>PIK3C2B</i> , <i>PPP1R15B</i> , <i>MDM4</i>)	9.77×10^{-6}	EGR2, HOXC6, TWIST1, ZFP82, ZNF528	EGR1, HAND1, NFIA, ZNF816
Chr 16: 86,954,235–86,955,236 (<i>C16orf95</i> , <i>FOXL1</i> , <i>FBXO31</i>)	1.06×10^{-5}	ETS2, NFAT5, PITX2	FLI1, SP2, TAF1, ZEB1, ZNF317, ZNF418
Chr 22: 37,128,477–37,129,478 (<i>IL2RB</i> , <i>TMPRSS6</i> , <i>C1QTNF6</i>)	1.33×10^{-5}	EGR2, HAND1, KLF16, ZNF263	NHLH1, ZNF322
Chr 9: 100,937,755–100,938,756 (<i>PLPPR1</i> , <i>MURC</i> , <i>BAAT</i>)	1.51×10^{-5}	FOXH1, TBX15, ZNF816	AR, ETV7, TGIF2LX, ZNF335
Chr 9: 100,937,771–100,938,772 (<i>PLPPR1</i> , <i>MURC</i> , <i>BAAT</i>)	1.51×10^{-5}	FOXH1, TBX15, ZNF816	AR, ETV7, TGIF2LX, ZNF335
Chr 11: 120,020,704–120,021,705 (<i>TRIM29</i> , <i>OAF</i> , <i>POU2F3</i>)	1.59×10^{-5}	ARID5B, SOX9, TGIF1, ZNF816	IRF2, NKX6-1, TEAD2, ZNF394
Chr 17: 8,915,658–8,916,659 (<i>PIK3R5</i> , <i>PIK3R6</i> , <i>NTN1</i>)	1.79×10^{-5}	NR2F1, POU2F2, SP3, UBP1, ZNF148	ETV3, NR2F6, ZNF317
Chr 17: 958,411–959,412 (<i>NXN</i> , <i>TIMM22</i> , <i>ABR</i>)	1.9×10^{-5}	EVX1, FOXG1, PITX2, ZNF770	FOXD2, PAX5, ZNF667
Significantly associated promoters			
Three promoters near <i>RET</i>	1.24×10^{-7a} – 2.48×10^{-7}		
Significantly associated genes			
16 genes near <i>RET</i>	4.07×10^{-7a} – 6.05×10^{-6}		
<i>ODF2</i> (Chr 9: 128,455,155–128,501,292)	9.27×10^{-6}	E2F4, MYOD1, ZBTB4, ZNF274	NR113, SMAD3, TGIF2, ZFP64
<i>URM1</i> (Chr 9: 128,371,319–128,392,016)	4.35×10^{-5}	E2F4, ELK3, MYOD1, ZBTB4, ZNF274	NR113, SMAD3, ZFP64
<i>LINS1</i> (Chr 15: 100,559,369–100,603,230)	5.89×10^{-5}	FOXC1, FOXF1, ZNF394	HAND1, MECP2, MTF1, ZSCAN31
<i>ASB7</i> (Chr 15: 100,602,534–100,651,705)	6.81×10^{-5}	FOXC1, FOXF1, ZNF394	HAND1, MECP2, MTF1, ZSCAN31

For significant S-HSCR-associated enhancers, promoters, and genes, regions near *RET* are collapsed into one entry. For the enhancer list, the nearest genes of each enhancer were identified based on the distance between the center position of the enhancer and the TSSs of genes. Expressed TFs are defined based on our hNC RNA-seq data, with a cutoff threshold of FPKM (fragments per kilobase of exons per million reads) = 1.

^aThe smallest *P*-values possible from the permutation tests.

The four significantly associated genes not close to the *RET* locus come from two loci, respectively, on Chromosomes 9 and 15, with no loosely associated enhancers or promoters near them. Each of these genes was thus found to be significantly associated with S-HSCR due to weak association signals that distribute across multiple regulatory elements of it. Indeed, a significantly larger fraction of enhancers within 1 Mbp from the TSSs of these genes have *P*-values smaller than 0.1 as compared to the set of all enhancers ($P=0.025$, Fisher's exact test). One of the genes in these loci is *ODF2*, which encodes the cenexin protein. Previous studies have demonstrated that, in various mammalian cell lines, cenexin controls centrosome positioning during directional cell migration (Hung et al. 2016). Since the centrosome acts as a steering machine for cell movement and stabilizes the migration direction (Ueda et al. 1997; Hung et al. 2016), *ODF2* might help stabilize the direction of NC migration.

Loosely associated signals aggregate into functional pathways

Having found that genetic variants weakly associated with S-HSCR at different regulatory elements aggregated to give stronger signals at the gene level, we next investigated whether association signals could be further aggregated into functional pathways. We performed this investigation by taking the 417 genes loosely associat-

ed with S-HSCR from our enhancer-based, promoter-based, and gene-based analyses and connected those with functional interactions in Reactome (Supplemental Materials; Wu and Haw 2017). A total of 69 genes were connected, by 94 functional interactions. This number of functional interactions was significantly larger than those of random gene sets of the same size ($P=0.032$), suggesting that these S-HSCR genes were indeed functionally related.

To annotate the functions of these S-HSCR genes, we expanded this gene set by adding known HSCR genes and genes important in ENS development or NC migration (Methods) and found that the resulting genes formed five main clusters according to their functional interactions (Fig. 3A). These five clusters are, respectively, related to (1) chemotaxis and cell-cell signaling, (2) cell adhesion, migration, and interaction with extracellular matrix, (3) PI3K-PKC-MAPK signaling, (4) E3 Ubiquitin Ligase complexes, and (5) transcriptional regulatory factors. They are highly related to NC migration and its development (Fig. 3B), and many genes loosely associated with S-HSCR were involved in these functional clusters (Supplemental Table S2).

Chemotaxis signaling is used to direct NC to target tissue (positive signaling) and works against the negative signals from the neural tube or somite to promote NC delamination and migration (Szabó and Mayor 2018). Additionally, cell polarization is critical during the directional migration of NC. Under the regulation

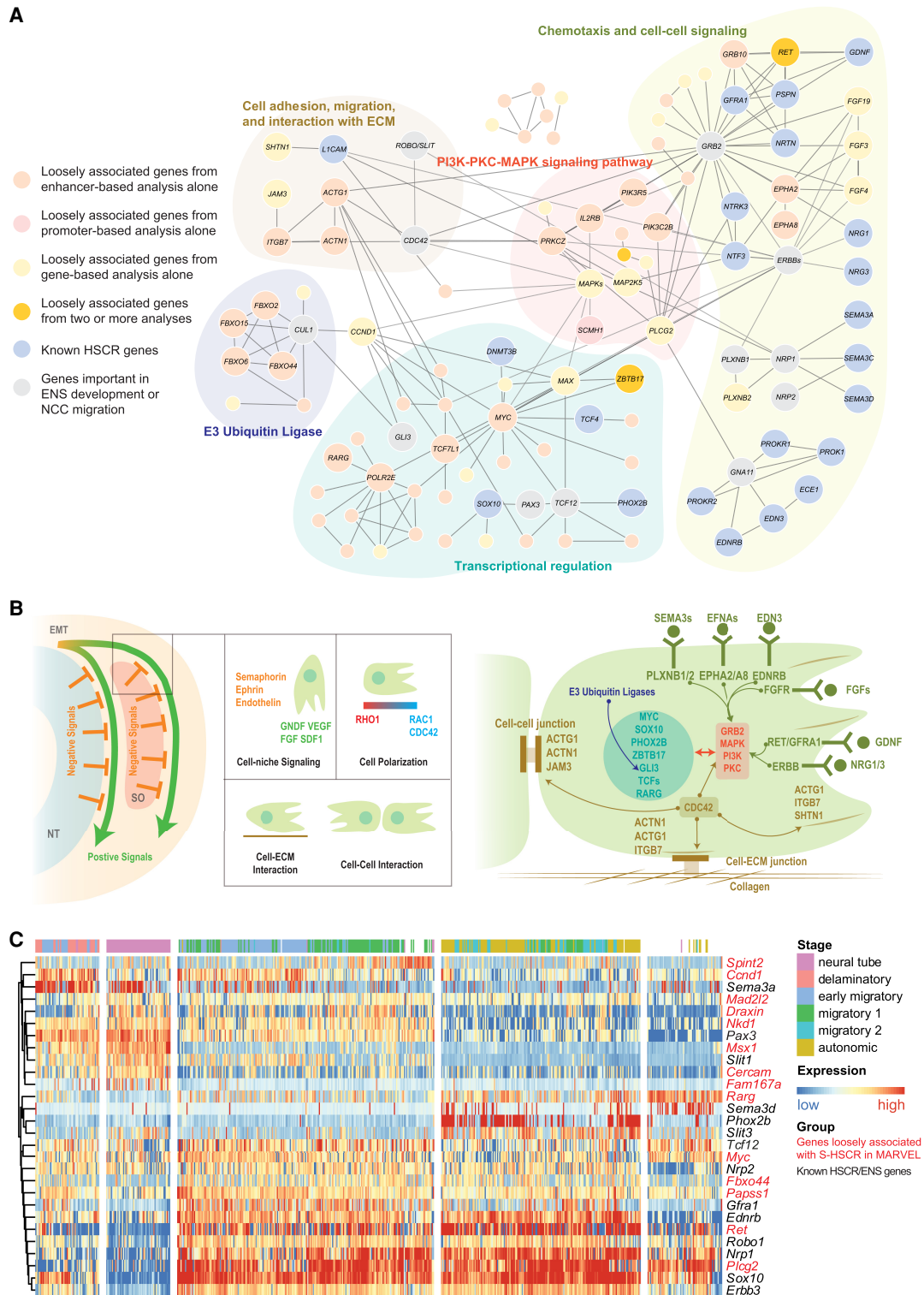


Figure 3. Functional landscape of S-HSCR-associated genes. (A) Functional interactions among genes loosely associated with S-HSCR, known HSCR genes, and genes important in ENS functions or NC migration. Each node corresponds to a gene and each edge corresponds to a functional interaction cataloged in Reactome. Genes of particular interest are shown in bigger nodes, labeled with their names. (B) Schematic illustration of some biological processes and genes involved in NC migration (Szabó and Mayor 2018). Colors of gene names follow their functional categories in panel A. (NT) Neural tube, (SO) somites. (C) Spatiotemporal expression profiles of mouse trunk NCs. Each row corresponds to the stage-specific expression pattern of a mouse homolog of a human gene shown in panel A. Genes identified by MARVEL as loosely associated with S-HSCR are shown in red. Each column corresponds to a single cell with the predicted stage label taken from the original publication (Soldatov et al. 2019). Both the rows and the columns were clustered using hierarchical clustering with Pearson's correlation as the similarity measure, and the columns are divided into five partitions according to the clustering results.

of the RAC1-RHO1 gradient, actin is selectively polymerized on one side of the cell, extending the cell along the migration direction (Cohen et al. 2018; Szabó and Mayor 2018). Similarly, in rat axonal growth cones, under the regulation of CDC42/RAC1, SHTN1 is involved in the connection between F-actin and the cell adhesion molecule L1CAM (Kubo et al. 2015). The extracellular matrix (ECM) serves as the road of NC migration (Szabó and Mayor 2018), and collagens are basic structures in the ECM required for successful NC migration (Nagy et al. 2018). Cell-ECM interactions such as focal adhesion through integrins and vinculins provide both structural support and signaling sensor for migrating NC. In a previous study, mutations of vinculin in HSCR patients were shown to affect its function in focal adhesion (Lai et al. 2017).

Signals from cell-ECM interactions, cell-cell interactions, and chemotaxis are integrated inside NC through several signaling pathways including the PI3K, MAPK/ERK, and PKC pathways (Dinsmore and Soriano 2018). Through these pathways, signals from the environment are transformed into regulatory signals, controlling the activation and deactivation of genes.

An additional layer of regulation of these cellular signals was found at the protein level. For instance, E3 ubiquitin ligase is required by GLI3, a known HSCR-associated TF, to switch between its activator and repressor forms (Hsia et al. 2015), by which GLI regulates the differentiation and patterning of enteric NC (Liu et al. 2015).

Besides known HSCR TFs, we also found several novel TFs loosely associated with S-HSCR, such as MYC and ZBTB17 (Supplemental Table S2), which are known to regulate the size of the pool of premigratory NC (Kerosuo and Bronner 2016), and RARG, which is important in the retinoic acid signaling pathway, a pathway crucial for enteric NC migration (Uribe et al. 2018).

Overall, our results reveal an unprecedentedly broad linkage between noncoding genetic variants associated with S-HSCR and NC genes and clearly demonstrate that these genetic variants distribute across many different regulatory elements and genes that converge to key functional pathways.

As an additional way to explore the functions of the genes loosely associated with S-HSCR, we examined the spatiotemporal expression of their mouse homologs using a trunk NC single-cell RNA-seq (scRNA-seq) data set (Supplemental Materials; Soldatov et al. 2019). Some of the known HSCR genes and loosely associated genes display stage-specific expression (Fig. 3C; Supplemental Materials).

Upstream regulators of S-HSCR-associated genes

One major advantage of our framework is its ability to identify not only the disease-associated regulatory elements or genes but also their upstream transcriptional regulators whose binding sites are altered by the genetic variants. We developed a permutation test to identify TF binding motifs whose match scores are significantly more altered in the enhancers loosely associated with S-HSCR found in our enhancer-based analysis than the background set of all enhancers (Methods). This analysis identified 48 TFs that we will refer to as “recurrent TFs” (Supplemental Table S3). Among the top 100 promoters with the strongest association *P*-values with S-HSCR, 73 of them also had at least one of these recurrent TFs’ motifs selected by our procedure as a key motif, suggesting that, in NCs, these recurrent TFs regulate gene expression through both promoters and enhancers.

The recurrent TFs form an extensive network by way of physical protein-protein interactions (PPIs), with MYC and SMAD pro-

teins acting as interaction hubs (Fig. 4A). The premigratory NC pool size regulators ZBTB17 and MYC and the retinoic acid receptor RARG discussed above in the functional pathway analysis are also contained in this network, showing that both the expression of these genes themselves and their downstream targets could be perturbed in S-HSCR. Some other interactions in this network are also known to have important functional roles. For example, MYC interacts with SMAD to activate expression of *Snail*, which are essential TFs that regulate EMT of NC (Smith et al. 2009; Kerosuo and Bronner 2016). The E2F proteins have been shown to regulate neuronal migration (McClellan et al. 2007). There are also reported HSCR genes in the network, such as *GLI3* (Liu et al. 2015).

Using the same mouse trunk NC scRNA-seq data described above, we found that the recurrent TFs with the strongest across-cell expression variance are *Tfap2b* and *Plagl1* (Fig. 4B). *TFAP2B* (human homolog of *Tfap2b*) is a known migratory NC marker (Rabadán et al. 2013). *Plagl1* is up-regulated in neural tube and in the delaminatory stage, which potentially regulates the composition of the ECM (Varrault et al. 2017) and thus potentially affects the migration environment of NC. One gene that has a similar expression pattern as *Plagl1* is *Zic2*, which regulates the production of NC (Elms et al. 2003).

Enhancers significantly associated with S-HSCR confer cell stage-specific expression of *RET*

To delineate the potential biological roles of our newly identified S-HSCR-associated enhancers in disease susceptibility and pathogenesis, we employed hPSCs to model the development of the human ENS. We started with the most well-characterized HSCR-associated common SNP, rs2435357, where the risk allele *T* decreases the expression of *RET* and contributes to disease prevalence (Tang et al. 2016). As mentioned above, this SNP resides in an enhancer within intron 1 of *RET* that was found to be significantly associated with S-HSCR in our enhancer-based analysis (Table 1). It overlaps an hNC-specific ATAC-seq peak (Fig. 5A), suggesting that it may be implicated in the cell stage-specific expression of *RET*. To test this hypothesis, we introduced the risk allele *T* into a control hPSC line using a CRISPR-Cas9-mediated homology-directed repair (HDR) system with a specific single guide RNA (sgRNA) targeting the rs2435357(C/C) locus and single-stranded oligonucleotides (ssODNs) (Ran et al. 2013; Paquet et al. 2016; Kwart et al. 2017) containing the *T* allele (Fig. 5B). The control and mutant (UE-rs2435357) hPSC lines were then used to generate neural crest (hNC: SOX10⁺) and were subsequently directed to the neuronal lineage to make the early enteric neuronal progenitors (hNP: TUJ1⁺, ELAVL4⁺), recapitulating the two key developmental stages during the formation of the ENS as previously described (Fig. 5C; Fattahi et al. 2016; Lai et al. 2017). The single base (*C* > *T*) conversion alone did not affect hNC induction and the derivation of hNPs from hPSCs. We found that *RET* is highly expressed in the hNPs, while its expression remains at a low level in hNCs before committing to neuronal lineage, suggesting that the cell stage-specific enhancers are likely involved to mediate the dynamic expression of *RET* during the development of the ENS. In concordance with this observation, the *C* > *T* conversion in rs2435357 significantly reduced *RET* expression in hNPs derived from the UE-rs2435357 hPSC line, but it did not affect *RET* expression in hPSCs when compared to the isogenic control line (Fig. 5D). Our data suggest that the risk allele *T* in rs2435357 reduces *RET*

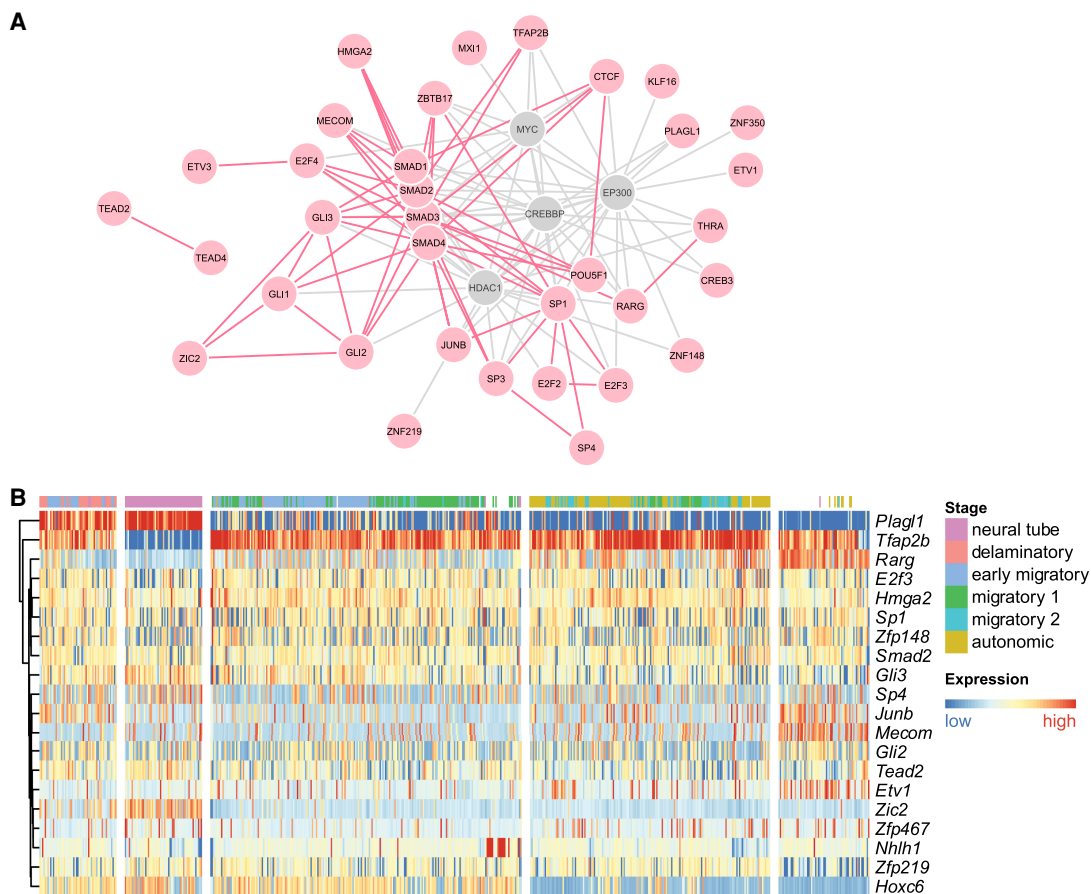


Figure 4. Analysis of the recurrent TFs. (A) PPIs among the recurrent TFs (red) and several other proteins frequently interacting with them (gray). Direct interactions among the recurrent TFs are shown in red, while direct interactions involving the other proteins are shown in gray. Recurrent TFs that have no interactions with other proteins in this figure are excluded. (B) Spatiotemporal expression of recurrent TFs with stage-specific expression profiles in mouse trunk NCs. The heat map was produced in the same way as in Figure 3C, with the same order of columns.

expression in the early enteric neural progenitors (hNPs), contributing to HSCR pathogenesis.

Next, we studied a novel enhancer (Chr 10: 43,253,960–43,254,961) significantly associated with S-HSCR within intron 1 of *RASGEF1A* that is located around 180 kbp downstream from the TSS of *RET* (Table 1). This putative enhancer is close to the TSS of *RET* in the three-dimensional (3D) genome architecture in neural progenitor cells based on previously published Hi-C data (Fig. 5E; Dixon et al. 2015). ATAC-seq data also revealed that there was an hNC-specific peak in this region, implying that this enhancer may also be implicated in ENS development (Fig. 5F). To investigate the potential effect of this novel enhancer in controlling *RET* expression, we generated a knockout (KO) mutant hPSC line (*RASGEF1A-int1-KO*) using a CRISPR-Cas9 system (Ran et al. 2013) with a pair of specific sgRNAs flanking the target region of intron 1 of *RASGEF1A*. Single hPSC colonies were isolated and genotyped to confirm the deletion of the target region (Fig. 5G). After neural induction and neuronal differentiation, we found that the differentiation potential of *RASGEF1A-int1-KO* hPSCs to make hNCs and hNPs was highly comparable to that of the control. Comparable numbers of SOX10-, TUJ1-, and HU-expressing cells were found in the control and the *RASGEF1A-int1-KO* groups (Fig. 5H). However, the deletion significantly reduced *RET* expression in hNPs, but not in hPSCs (Fig. 5I), implying that intron 1 of

RASGEF1A contains a long-range regulator of *RET* in the early enteric progenitors.

Disruption of NFIA binding in a novel S-HSCR-associated regulatory element in intron 10 of *PIK3C2B* interferes with the expression of *PIK3C2B*, *PPP1R15B*, and *SOX13*

We further experimentally studied the functional potential of another S-HSCR-associated regulatory element located in intron 10 of *PIK3C2B* (Chr 1: 204,456,576–204,457,577). This regulatory element was of particular interest because (1) it contained a strong hNC-associated ATAC-seq peak (Fig. 6A), (2) an *A > T* variant in this locus (rs551359143) was identified in six of our S-HSCR patients but not in any control subjects, and (3) the *A > T* conversion was predicted to disrupt the binding motif of NFIA (Table 1), which is a TF with a key role in regulating the differentiation of neural stem cells (Fig. 6A; Sanosaka et al. 2009; Piper et al. 2010; Kang et al. 2012; Tchieu et al. 2019). We generated another mutant hPSC line in which 171 bp of the intron 10 of *PIK3C2B* flanking the putative NFIA binding motif were deleted using a CRISPR-Cas9 system. The deletion of the target region was confirmed by genotyping in the *PIK3C2B-int10-KO* hPSC clone (Fig. 6B). The mutant hPSC clone was used to make hNCs and hNPs as described above. Similar to the other enhancers discussed above, the deletion

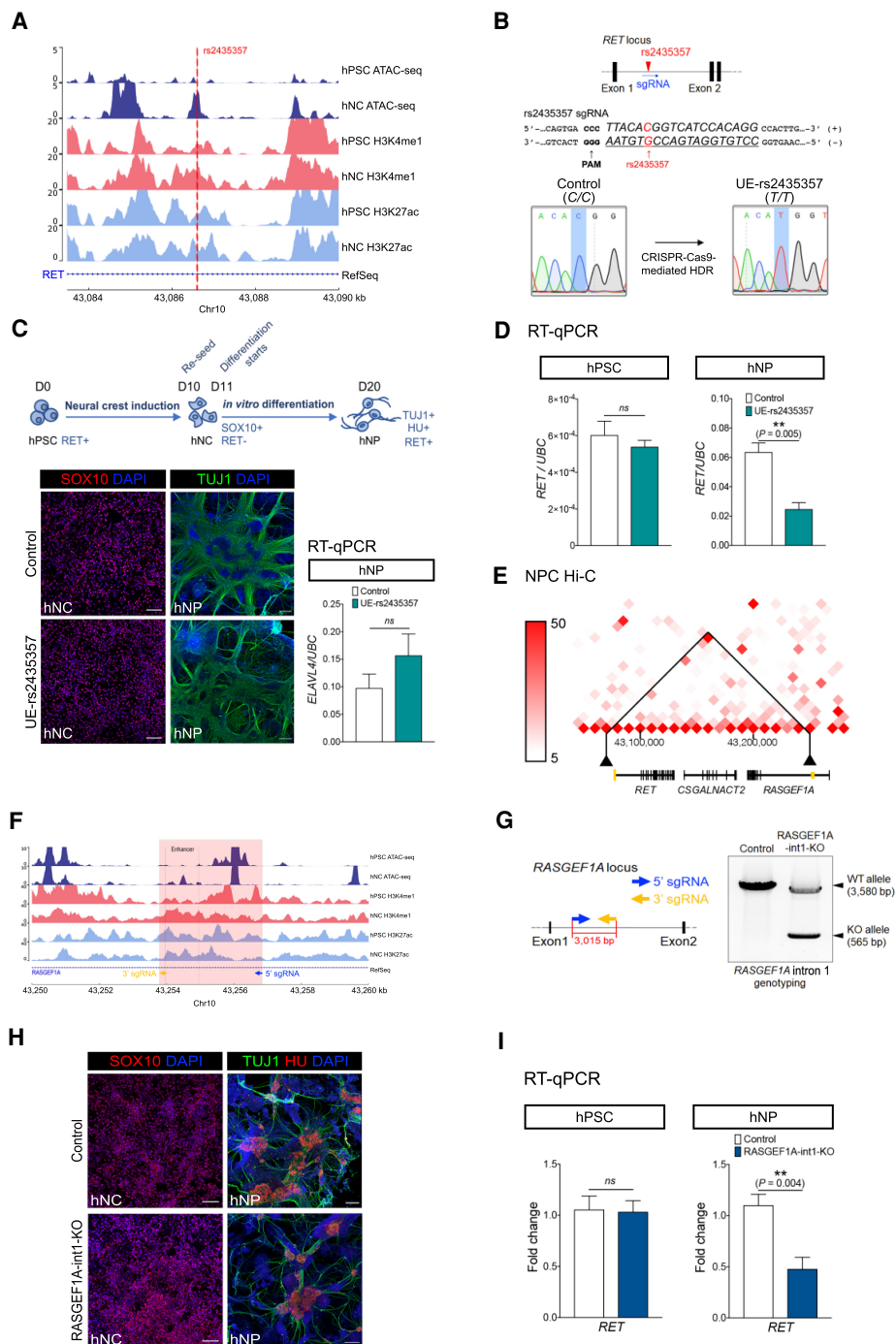


Figure 5. Functional impacts of a HSCR-associated SNP (rs2435357) and the deletion of a novel S-HSCR enhancer on *RET* expression. (A) ATAC-seq and ChIP-seq profiles of hPSC and hNC in intron 1 of *RET* show that rs2435357 is residing in a hNC-specific ATAC-seq peak. (B) Location of rs2435357 in the *RET* gene locus and in the sgRNA used for CRISPR-Cas9-mediated HDR for editing the C allele to the HSCR-associated risk allele T. The electropherograms of Sanger sequencing show the successful introduction of the risk allele at rs2435357 in the UE-rs2435357 hPSC line. (C) Differentiation strategy to generate human neural crest (hNC) and neuronal progenitor (hNP). HU is encoded by the *ELAVL4* gene. Immunostaining of SOX10 and TUJ1 in hNC and hNP of the control and the mutant (UE-rs2435357) lines. Scale bars: (hNC) 100 μ m; (hNP) 200 μ m. RT-qPCR analysis showing the comparable *ELAVL4* expression level in hNP in the control ($n=5$) and the mutant (UE-rs2435357) ($n=3$) lines. *t*-test, (ns) not significant. (D) RT-qPCR analysis showing *RET* expression in the hPSC and hNP stages of the control ($n=5$) and the mutant (UE-rs2435357) ($n=3$) lines. *t*-test, (ns) not significant. (E) Hi-C data from neural progenitor cells show that the enhancer in intron 1 of *RASGEF1A* (marked in yellow on the right) has physical interaction with the promoter of *RET* (marked in yellow on the left) at 10-kbp bin size. (F) ATAC-seq and ChIP-seq data from hPSC and hNC at the *RASGEF1A* intron 1 locus. (G) The design of sgRNAs used for the CRISPR-Cas9 system for deleting the DNA fragment in *RASGEF1A* intron 1. Genotyping reveals the specific deletion of *RASGEF1A* intron 1 in the UE-RASGEF1A-int1-KO hPSC line. (WT) Wild type, (KO) knockout. (H) Immunostaining of SOX10, TUJ1, and HU in hNC and hNP of the control and the mutant (RASGEF1A-int1-KO) lines, respectively. Scale bars: (hNC) 100 μ m; (hNP) 200 μ m. (I) RT-qPCR reveals the expression level of *RET* in the hPSC and hNP stages of the control ($n=4-5$) and the mutant (RASGEF1A-int1-KO) ($n=6-7$) lines. *t*-test, (ns) not significant.

MARVEL identifies functional noncoding variants

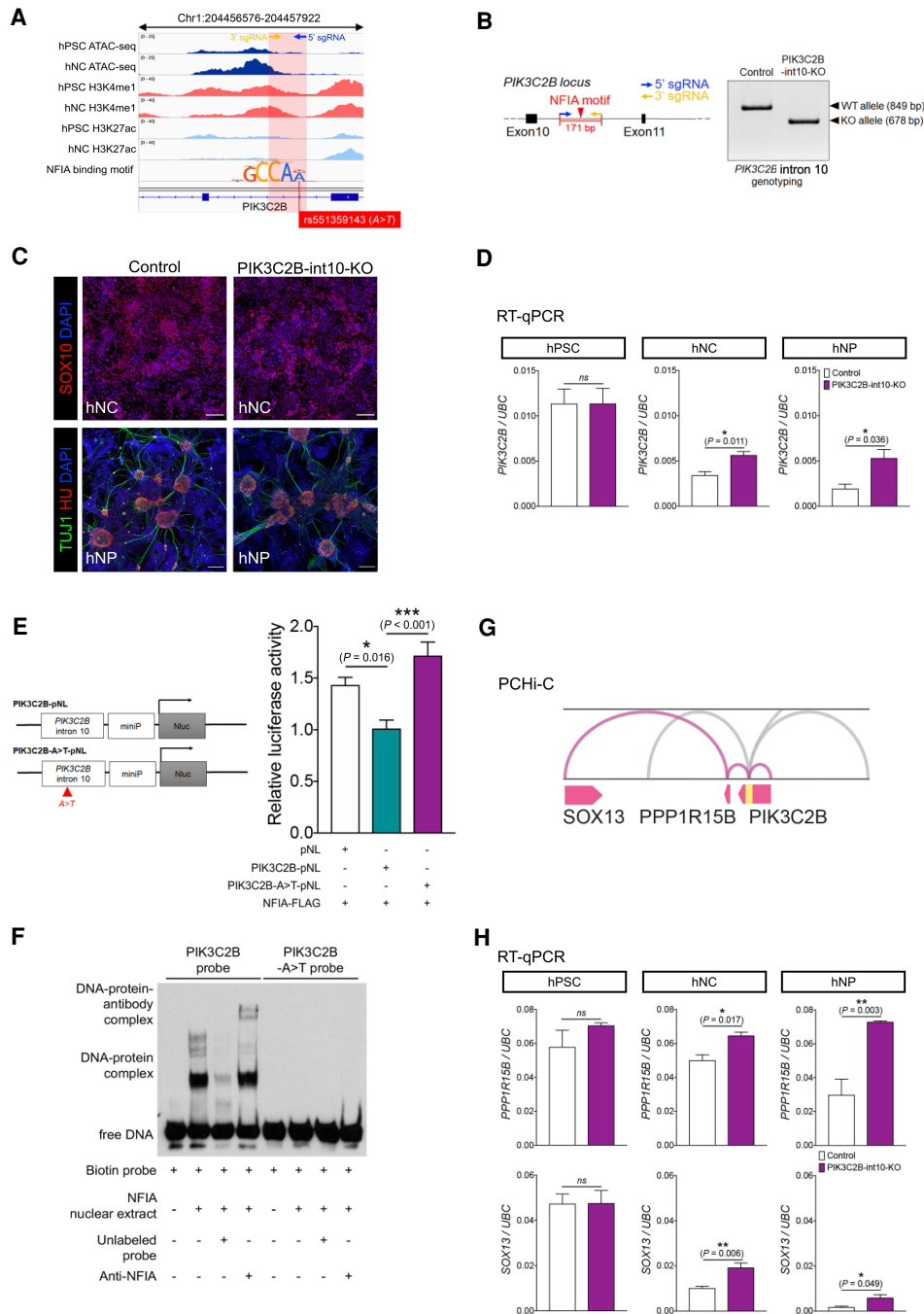


Figure 6. Characterization of a novel S-HSCR-associated regulatory element in intron 10 of *PIK3C2B*. (A) Overview of ATAC-seq and ChIP-seq profiles showing the putative hNC-specific regulatory element in *PIK3C2B* intron 10. The red shaded region indicates the location of the regulatory element and the line shows the *A > T* variant (rs551359143) found exclusively in the S-HSCR cases that disrupts the NFIA binding motif. The motif is not drawn to the same scale as the genomic signal tracks, with magnified characters. (B) Design of sgRNAs used for the CRISPR-Cas9 system for deleting the regulatory element. Genotyping reveals the specific deletion of the 171-bp fragment in intron 10 of *PIK3C2B* in the *PIK3C2B*-int10-KO hPSC line. (WT) Wild type, (KO) knockout. (C) Immunostaining shows that both the control and mutant (*PIK3C2B*-int10-KO) lines have comparable capability to make hNCs and hNPs. Scale bars: (hNC) 100 μ m; (hNP) 200 μ m. (D) RT-qPCR shows the changes in the expression of *PIK3CB* in different cell stages in the control and mutant lines. *t*-test, (ns) not significant. $n = 3-4$ per group. (E) Design of the constructs used for the luciferase assay. The bar chart shows the relative luciferase activities when the cells were transfected with different sets of constructs as indicated. Three independent assays were performed, each in triplicate. One-way ANOVA. (F) Gel mobility shift assays were performed with biotin-labeled probes containing the *PIK3C2B* intron 10 regulatory element with or without the *A > T* conversion and the nuclear extract from NFIA-overexpressing cells, in the presence of unlabeled probes or anti-NFIA antibody (0.1 μ g). (G) Significant contacts (FDR < 0.05) in the promoter capture Hi-C data from GM12878 cells at the *PIK3C2B* locus. The putative regulatory element in intron 10 of *PIK3C2B* is marked in yellow. Contacts between the regulatory element and the TSSs of *SOX13*, *PPP1R15B*, and *PIK3C2B* are shown in purple curves, while contacts between the regulatory element and other promoters are shown in gray curves. Contacts that extend too far are trimmed. (H) RT-qPCR analysis shows the changes in the expression of *PPP1R15B* and *SOX13* in the control and the mutant lines at different cell stages. *t*-test, (ns) not significant. $n = 3-4$ per group.

of the fragment did not affect the differentiation potential of the hPSCs to make hNCs and hNPs (Fig. 6C). In terms of gene expression, our quantitative RT-PCR (RT-qPCR) revealed a sequential down-regulation of *PIK3C2B* along the enteric neuronal progenitor differentiation (hPSC-to-hNC-hNP). When the 171-bp fragment was deleted, the expression of *PIK3C2B* in hNCs and hNPs was significantly increased as compared to the isogenic control, implying that this 171-bp fragment probably serves as a negative regulatory element to mediate the sequential down-regulation of *PIK3C2B* along the differentiation path and the loss of it led to aberrant expression of *PIK3C2B* gene in hNCs and hNPs (Fig. 6D). We then further demonstrated the involvement of NFIA in mediating *PIK3C2B* expression using a luciferase assay. Our data showed that NFIA mainly acts as a repressor and the luciferase activity was significantly up-regulated when the NFIA binding motif was disrupted by introducing the *A > T* substitution (Fig. 6E). A direct binding of NFIA onto the intron 10 of *PIK3C2B* was demonstrated using a gel shift and supershift assay, in which NFIA binding was consistently abolished by the *A > T* conversion (Fig. 6F). All these results indicate that *PIK3C2B* was negatively regulated by NFIA and the HSCR-associated *A > T* variation would disrupt NFIA binding, leading to the increased expression of *PIK3C2B*.

Long-range chromatin interactions form topologically associating domains (TADs) and the genes within a TAD are likely to be regulated by common enhancers (Chatterjee et al. 2016; Laugsch et al. 2019). A previous promoter capture Hi-C experiment suggests that chromatin loops can be formed between *PIK3C2B* intron 10 and the TSSs of *PPP1R15B* and *SOX13* in GM12878 cells (Fig. 6G; Mifsud et al. 2015). This suggests that the gene expression of *PPP1R15B* and *SOX13* may also be influenced by the S-HSCR-associated *A > T* variant in intron 10 of *PIK3C2B*. RT-qPCR revealed that the expression levels of *PPP1R15B* and *SOX13*, similar to that of *PIK3C2B*, were not affected in the hPSC stage but up-regulated in the hNC and hNP stages after the deletion of the 171-bp fragment (Fig. 6H). This result suggests that *PIK3C2B* intron 10 contains an element that can regulate the expression of genes within the same TAD and the usage of this regulatory element may be dynamic during the development of ENS. During neuronal differentiation, this element likely exerts negative control over gene expression, as the deletion of it consistently increased gene expression. In comparison, in the control line, all the genes regulated by this element were down-regulated when the cells differentiated into neuronal lineages, suggesting that up-regulation of these genes associated with the *A > T* variant may impose disease-causing risks during the later stage of enteric NC cell development.

Discussion

In this work, we have proposed a novel framework, MARVEL, for identifying noncoding genetic variants of potential functional significance. It was designed to overcome various issues in the study of noncoding variants. Our application of MARVEL to the analysis of WGS data of S-HSCR has clearly demonstrated the advantages of these designs. Starting from 37 million noncoding variants observed in the cases and controls, the epigenomic data reduced the number of genomic regions to study to the scale of tens of thousands (of genes and promoters) to hundreds of thousands (of enhancers), and the statistical procedures further selected no more than 10 key motifs from each of these regions among 771 of them. As a result, we were able to obtain some results other than those in the dominating *RET* locus at fairly strong signifi-

cance levels, which would be impossible in the traditional approach of testing the disease association of each genetic variant separately since most noncoding variants are rare or have small effect sizes.

Comparing the enhancer-based and promoter-based analyses, we obtained more significant results from the former, showing that it is important to move beyond the immediate TSS-proximal regions when analyzing the noncoding variants. The designs of MARVEL made it possible to study genetic variants in distal enhancers in a computationally and statistically feasible manner.

The gene-based analysis led to findings in Chromosomes 9 and 15 not obtained from the enhancer-based or promoter-based analysis. We have found that around these gene loci there were more enhancers with marginally significant associations than other gene loci, suggesting that it was the aggregation of weak signals from multiple regulatory elements that helped identify these gene loci as significantly associated with S-HSCR. The importance of this aggregation idea was further demonstrated in the functional pathway analysis, in which some of the 417 genes loosely associated with S-HSCR were grouped into functional clusters that are highly related to NC migration.

Our analysis of upstream regulators has identified 48 recurrent TFs, and they form an extensive network through PPIs. Notably, some of these TFs also have their corresponding genes identified in the association tests, showing that the disruptions of both the expression of these TFs themselves and their binding at other genes' regulatory regions are associated with S-HSCR, illustrating two main mechanisms by which genetic variants can exert their functional effects.

Making use of the CRISPR-Cas9 genome editing platform, we were able to generate various hPSC lines with isogenic background, such that any phenotypical differences observed should be solely due to the effects of the risk variants. This platform allowed us to study the functional impacts for the risk variants in the human genome, in contrast to previous studies that made use of other animal systems for enhancer studies, which usually focused on the highly conserved regions of the genome instead of the disease-associated variants.

The motif analysis in our new model is also capable of identifying the upstream transcription factors that regulate the HSCR-associated regulatory elements. For instance, we have found that NFIA may act as a negative regulator when binding to the regulatory element of intron 10 of *PIK3C2B* during neural differentiation. This finding is consistent with a previous report that suggests NFIA represses *Hes1* during neural progenitor differentiation in mouse forebrain (Piper et al. 2010). A recent study from Qi et al. (2020) has found that the conditional knockout of *Nfia* in mouse neural crest cells (i.e., *Wnt1-Cre; Nfia^{fl/fl}*) would only disrupt the development of NPY2R-expressing neurons but not other neurons such as TH-expressing neurons in vivo (Qi et al. 2020). This suggests that NFIA is responsible for controlling the development of a specific neuronal subtype and may be involved in the later stage of neuronal development. Inhibition of *PIK3C2B* has been shown to improve function and prolong survival in myotubular myopathy through regulating the phosphoinositide (PIP) metabolism (Sabha et al. 2016). As the dysregulation of PIP metabolism may lead to neurodegenerative disease (Burke 2018), the up-regulation of *PIK3C2B* caused by the *A > T* SNP in intron 10 of *PIK3C2B* may affect the survival of enteric neurons.

Similar to *PIK3C2B*, up-regulation of *PPP1R15B* may induce cellular stress and affect cell survival through inducing protein misfolding conditions as seen in Huntington's disease

(Krzyszosiak et al. 2018). On the other hand, it has been demonstrated that SOX13 can functionally replace SOX5 and SOX6 in governing oligodendrocyte development in mouse spinal cord (Baroti et al. 2016), suggesting that it modulates cell differentiation into glial lineages instead of neuronal lineages. Therefore, ectopic SOX13 expression may affect the cell fate determination of the enteric NC cells. Given that many of these genes may affect various stages of the ENS development, further studies are therefore required to unveil the functions of these differentially regulated genes in the development of ENS.

Overall, our work has produced a large number of novel regulatory elements, genes, and upstream regulators associated with S-HSCR for follow-up investigations. Due to the buffering of different regulatory elements and different genes, single perturbations may have little observable effects on cell differentiation, and as a result their roles in disease susceptibility and pathogenesis are subtle. In order to have a more complete understanding of the functional importance of the candidate regulatory elements, genes, and TFs identified in this study, it would be useful to perform large-scale genome editing assays to systematically perturb them individually and in combination and determine how they may interfere in various steps of ENS development.

Methods

Statistical procedures of the MARVEL framework

Here, we describe the statistical procedures used by the MARVEL framework. Additional details of the framework can be found in [Supplemental Materials](#).

Identification of important motifs and phenotype-associated target regions

Suppose there are n subjects with their phenotype information recorded in the binary vector \mathbf{y} , where $y_i = 1$ if subject i is a case and $y_i = 0$ if subject i is a control. For each target region, the standardized score profiles are represented by an $n \times m$ matrix \mathbf{X} , where m is the number of motifs. Covariates of the n subjects are given in an $n \times v$ matrix \mathbf{C} , where v is the number of covariates.

Based on \mathbf{X} and \mathbf{y} , generalized linear model-based least angle regression (GLM-LARS) (Tibshirani et al. 2004) is performed to select a subset of motifs whose match scores in this target region can best explain the phenotypes. We use `glmLasso` (Park and Hastie 2007) (<https://cran.r-project.org/web/packages/glmLasso/index.html>) to perform GLM-LARS with parameter settings `min.lambda = 1e-2`, `max.steps = 10`, and `max.vars = 10` such that up to 10 motifs can be selected and the computations are efficient.

To quantify how well the target region's genetic variations are associated with the phenotype based on the scores of the selected motifs, we construct two logistic regression models

$$\begin{aligned} H_0: g(\mathbf{y}) &= \mathbf{C}\boldsymbol{\beta}_0, \\ H_1: g(\mathbf{y}) &= (\mathbf{X}'|\mathbf{C})\boldsymbol{\beta}_1, \end{aligned}$$

where $g(u) = \log \frac{u}{1-u}$ is the logit function, \mathbf{X}' is a reduced version of \mathbf{X} retaining only the columns corresponding to the selected motifs, $(\mathbf{X}'|\mathbf{C})$ is a matrix that concatenates the columns of \mathbf{X}' and \mathbf{C} , and $\boldsymbol{\beta}_0$ and $\boldsymbol{\beta}_1$ are coefficients. The likelihood ratio between these two models is then used as a test statistic

$$\text{LR} = -2 \ln \frac{\text{Likelihood}(H_1)}{\text{Likelihood}(H_0)}.$$

The statistical significance of a target region is evaluated by comparing its LR with those from a permutation set, which in-

volves all the target regions with 100 random permutations of \mathbf{y} . For example, when we applied this procedure to perform association tests for the enhancers in the S-HSCR study, since there were 150,828 enhancers in total, each of these LR values were compared against the 15,082,800 values from the permutations to compute a P -value. The P -values of all the target regions are then corrected for multiple hypothesis testing using the Benjamini-Hochberg method (Benjamini and Hochberg 1995).

Estimation of effect size

We use AUROC to estimate the effect size of each target region. Specifically, after constructing the model $g(\mathbf{y}) = (\mathbf{X}'|\mathbf{C})\boldsymbol{\beta}_1$ for a target region, for each subject, this model is applied to compute a score that indicates how likely this subject has the phenotype. These phenotype scores together with the actual phenotype labels of the subjects are then used to produce the receiver operating characteristic, and the area under the curve is used as an estimate of the effect size.

Analysis of recurrent TFs

To identify the recurrent TFs, all motifs selected by GLM-LARS from the full set of 200 enhancers loosely associated with S-HSCR were collected. Ten thousand random sets of 200 enhancers were then formed by sampling from all enhancers, and GLM-LARS was applied to each of these sets. For each motif, its total model coefficients across all enhancers in a set were used as a test statistic, and a P -value was computed by comparing the test statistic obtained from the loosely associated enhancers with those from the random enhancer sets, to see if the former is significantly larger than the latter. These raw P -values were then corrected by the Benjamini-Hochberg method to control the FDR at 0.05.

The PPIs among the recurrent TFs were obtained from STRING (v11) (Szklarczyk et al. 2017). We included only the interactions based on manual curation or experimental evidence with a combined score >0.4 .

Functional studies

Cell culture

A control hPSC line (UE02302) was established and maintained as previously described (Lau et al. 2019). Neural crest induction was performed according to a previously described protocol (Lai et al. 2017). In brief, hPSCs were plated on Matrigel-coated plates in a density of 5×10^4 cells cm^{-2} in ES cell medium containing 10 ng/mL fibroblast growth factor 2 (FGF2, Peprotech). The differentiation was started by replacing ES cell medium with KSR medium and gradually switched to N2 medium from day 4 to day 10. For hNC differentiation, the cells were treated with LDN193189, SB431542, CHIR99021, and retinoic acid.

For neuronal differentiation of hNCs to hNPs, hNC cells (2.5×10^4) were seeded as droplets on polyornithine/laminin/fibronectin-coated surface in N2 medium containing 10 ng/mL FGF2 and 3 μM CHIR99021. For the *RET*-associated study, p75^{NTR} and HNK-1 double-positive hNC cells were sorted by a BD FACSAria III Cell Sorter; 5×10^4 sorted cells were seeded as droplets on a polyornithine/laminin/fibronectin-coated surface. Neuronal differentiation was initiated by replacing the medium with N2 medium containing BDNF, GDNF, NT-3, NGF, dibutyryl cAMP, and ascorbic acid for 9 d. Detailed procedures are described in [Supplemental Materials](#).

Plasmid constructions

Human codon-optimized high fidelity Cas9 nuclease construct (Ran et al. 2013) was obtained from Addgene (48138). Oligos for sgRNA cloning are listed in Supplemental Table S5. For the luciferase assay, *PIK3C2B* intron 10 fragment was cloned into a NanoLuc luciferase reporter construct (pNL3.2[*NlucP/minP*]) (Promega N1041) to generate the PIK3C2B-pNL construct, while *NFIA* ORF was cloned into the pFLAG-CMV plasmid to generate an NFIA-FLAG expression construct. The A>T variant was introduced to the PIK3C2B-pNL construct by site-directed mutagenesis using a QuikChange Lightning Site-Directed Mutagenesis kit (Agilent). The cloning primers and mutagenesis primers are listed in Supplemental Table S5, and detailed procedures are described in Supplemental Materials.

Generation of new hPSC lines using the CRISPR-Cas9 system

For the generation of the mutant (UE-rs2435357) hPSC line, UE control hPSCs were transfected with sgRNA construct, ssODNs and pSpCas9(BB)-2A-GFP construct by the Human Stem Cell Nucleofector kit 2 (Lonza). For the generation of UE-RASGEF1A-int1-KO and PIK3C2B-int10-KO hPSC lines, UE control hPSCs were transfected with a pair of pSpCas9(BB)-2A-GFP constructs containing the specific sgRNAs using the P3 Primary Cell 4D-Nucleofector X kit (Lonza). hPSCs expressing GFP were sorted as single cells and expanded for 2 wk. The single colonies were genotyped to confirm the site-specific conversion or the deletion of the target regions. Detailed methods are described in Supplemental Materials.

RT-qPCR

Total RNA from hPSCs, hNCs, and hNPs was extracted by an RNeasy Mini kit (Qiagen) and reverse-transcribed to cDNA using HiScript II Q RT SuperMix (Vazyme). The expression levels of the target genes were quantitated using real-time quantitative RT-PCR or droplet digital PCR (ddPCR). For real-time quantitative RT-PCR, target genes were amplified with Luna Universal Probe qPCR Master Mix (New England Biolabs) using a specific TaqMan Gene Expression Assay by ViiA 7 Real-Time PCR System (Thermo Fisher Scientific). ddPCR was used to measure *RET* expression in hNPs. In brief, cDNA samples were mixed with ddPCR Supermix for Probes (Bio-Rad 186-3010) and specific TaqMan probes and then subjected to droplet generation following the manufacturer's protocol. The reaction droplets were then subjected to thermal cycling, and the end-point fluorescence signals were measured by a QX200 Droplet Reader (Bio-Rad). Detailed procedures are described in Supplemental Materials.

Gel shift assay

The NFIA-FLAG expression construct was transfected to HeLa cells by FuGENE HD Transfection Reagent (Promega), and the nuclear extracts containing NFIA protein were then extracted using a nuclear and cytoplasmic extraction kit (Thermo Fisher Scientific). ssODNs derived from the intron 10 of *PIK3C2B* were biotin-labeled using a Biotin 3'-End DNA Labeling kit (Thermo Fisher Scientific) and then annealed with reverse complementary ssODNs to generate biotin-labeled probes. A gel shift assay was performed by mixing the nuclear extracts with biotin-labeled probes according to the manufacturer's protocol (LightShift Chemiluminescent EMSA kit; Thermo Fisher Scientific). For supershift assays, 0.1 µg anti-NFIA (Sigma-Aldrich HPA006111) was added to the mixture in the final step before incubation. Detailed methods are described in Supplemental Materials.

Luciferase assay

Control firefly luciferase construct (pGL3-control), NanoLuc luciferase constructs (pNL3.2[*NlucP/minP*] or PIK3C2B-pNL or PIK3C2B-A>T-pNL), and NFIA expression construct (NFIA-FLAG) were cotransfected into SH-SY5Y cells using jetPRIME transfection reagent (Polyplus Transfection) according to the manufacturer's protocol. Luciferase activities were detected with a Nano-Glo Dual-Luciferase Reporter Assay System (Promega) by the VICTOR Nivo Microplate Reader (PerkinElmer). Detailed procedures are described in Supplemental Materials.

Immunostaining

hNC cells and hNP-D9 cells were fixed in 4% PFA for 20 min at room temperature. Fixed cells were blocked with blocking solution (1% BSA, 0.1% Triton X-100 in PBS) at room temperature for 1 h. The blocked cells were then incubated with primary antibodies (mouse anti-SOX10 [1:500, R&D Systems MAB2864], rabbit anti-TUJ1 [1:1000, Abcam ab18207] and mouse anti-HU [1:1000, Thermo Fisher Scientific A-21271]) at 4°C overnight. After washing, the sections were incubated with respective secondary antibodies conjugated with Alexa Fluor 488/594 (Thermo Fisher Scientific 1:500) at room temperature for 2 h. The stained cells were mounted with ProLong Diamond Antifade Mountant with DAPI (Thermo Fisher Scientific). Fluorescence images were acquired by a Carl Zeiss LSM780 or LSM800 confocal microscope.

Chromatin interaction data

Hi-C data of NPC were obtained from the NCBI Gene Expression Omnibus (GEO; <https://www.ncbi.nlm.nih.gov/geo/>) with accession number GSE52457.

Data access

All raw and processed sequencing data generated in this study have been submitted to the NCBI Gene Expression Omnibus (GEO; <https://www.ncbi.nlm.nih.gov/geo/>) under accession number GSE154972. The source code of MARVEL is provided as Supplemental Code S1 and on GitHub (<https://github.com/fuxialexander/marvel>).

Competing interest statement

The authors declare no competing interests.

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