An Association Test of the Spatial Distribution of Rare Missense Variants within Protein Structures Improves Statistical Power of Sequencing Studies

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Running Title: Spatial Test of Rare Variants within Proteins

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ABSTRACT

Over 90% of variants are rare, and 50% of them are singletons in the Alzheimer's Disease Sequencing Project Whole Exome Sequencing (ADSP WES) data. However, either single variant tests or unit-based tests are limited in the statistical power to detect the association between rare variants and phenotypes. To best utilize rare variants and investigate their biological effect, we exam their association with phenotypes in the context of protein. We developed a protein structure-based approach, POKEMON (Protein Optimized Kernel Evaluation of Missense Nucleotides), which evaluates rare missense variants based on their spatial distribution on the protein rather than allele frequency. The hypothesis behind this is that the three-dimensional spatial distribution of variants within a protein structure provides functional context and improves the power of association tests. POKEMON identified four candidate genes from the ADSP WES data, namely two known Alzheimer's disease (AD) genes (TREM2 and SORL) and two novel genes (DUSP18 and CSF1R). For known AD genes, the signal from the spatial cluster is stable even if we exclude known AD risk variants, indicating the presence of additional low frequency risk variants within these genes. DUSP18 has a cluster of variants primarily shared by case subjects around the ligand-binding domain, and this cluster is further validated in a replication dataset with a larger sample size. POKEMON is an open-source tool available at https://github.com/bushlab-genomics/POKEMON.

INTRODUCTION

High-throughput DNA sequencing of diverse human populations has identified millions of genetic variants, the vast majority of which are exceptionally rare. A survey of ~60,000 individuals from the Exome Aggregation Consortium (ExAC) found that out of ~7M variants, 99% have a frequency <1% and 54% are singletons (Karczewski et al., 2020). Similarly, in the Alzheimer's Disease Sequencing Project (ADSP) Whole Exome Sequencing (WES) of ~10k individuals, 97% of identified variants have a minor allele frequency <1% and 23% are singletons (Butkiewicz et al., 2018). However, the effect of most rare variants on diseases of interest remains unknown because of insufficient statistical power to detect the associations between these variants and phenotypes.

We hypothesized that rare variants contribute to common diseases by forming clustered or dispersed patterns within protein structures that reflect modest disruption of protein function. Based on this hypothesis, incorporating protein spatial context should improve rare variant association tests. Prior studies have shown missense variants exhibit non-random patterns in protein structures, such as cancer-associated hot spot regions with a high density of missense

somatic mutation (Tokheim et al., 2016). Our group (Sivley et al., 2018) also found that germline causal missense variants for Mendelian diseases exhibit non-random patterns in 3D space, including both clusters and depletion.

To test this hypothesis, we developed a kernel function to quantify genetic similarity among individuals by using protein structure information. Consider a scenario where two individuals have different missense variants distal in genomic coordinates but close in 3D protein structure; these individuals will be assigned a high genetic similarity through our kernel function, which when applied over an entire dataset captures the spatial patterns of rare missense variants. Using a statistical framework similar to SKAT (Wu et al., 2011), we test the association of rare variants with quantitative and dichotomous phenotypes using this structure-based kernel. We call this approach POKEMON (Protein Optimized Kernel Evaluation of Missense Nucleotides). We validated that POKEMON can identify trait associations with spatial patterns formed by missense variants both in simulation studies and real-world data.

POKEMON identified four candidate genes from the ADSP WES Discovery Dataset, namely two AD genes (*TREM2* and *SORL*1) and two novel genes (*DUSP18* and *CSF1R*). Both *DUSP18* and *CSF1R* have clusters of variants primarily shared by case subjects around ligand-binding sites. Specifically for the cluster identified in *DUSP18*, we examined it in the ADSP WES replication dataset with a larger sample size and found that the cluster is better formed by the additional variants included. In summary, the cluster we identified is populated with variants mostly from cases and likely has a functional association with AD risk.

By performing an association test in the context of protein structure, we have identified highly relevant gene-disease associations which are driven by specific clusters of variants. Such clusters imply functional domains in the protein structure susceptible to variation and related to disease risk. Analyzing missense variants from complex disease studies in this way provides a new structural aspect which can be leveraged for association tests.

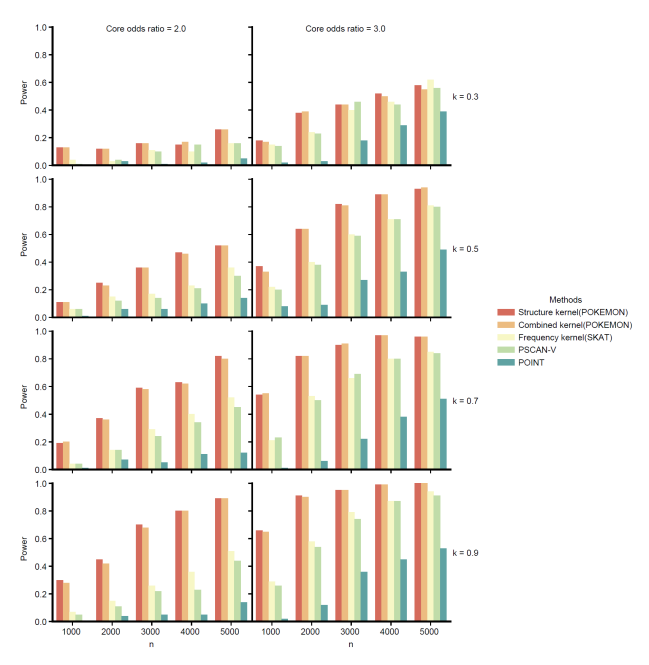


Figure 1. Empirical power to detect a pattern of association centered at the protein core among five methods with varying percentages of influential variants (k) and core variant odds ratios. Empirical power is calculated by counting the number of tests with a p-value below the significance level in 100 independent tests. The five methods include the structure kernel (POKEMON), combined kernel (POKEMON), frequency kernel (SKAT), PSCAN with variance test (PSCAN-V), and POINT. The core variant odds ratio is chosen as 2.0 or 3.0 (left to right). *k* is the percentage of pathological variants within the selected 50 variants, ranging from 0.3 to 0.9 (from top to bottom).

RESULTS

POKEMON can detect associations with spatially clustered or dispersed rare variants

As a proof of concept, we evaluated the performance of POKEMON using simulations that mimic real-world case/control studies. The simulation datasets varied in sample sizes (1000 - 5000), the odds ratio of the core variants (2.0 - 3.0), and the proportion of influential to neutral variants (0.3 - 0.9). For a cluster pattern, influential variants were simulated by establishing a maximum odds ratio decaying over a fixed distance of 14 Å. We limit the number of variants within the genotype profile to 50, which is the mean number of variants mapped per protein in the ADSP WES. For each simulation scenario, we generated 100 datasets and tested for spatial clustering using a structure kernel, estimating power as the proportion of significant tests. In general, POKEMON performed better than two other structure-based methods, PSCAN(Tang et al., 2020) and POINT(Marceau West et al., 2019) and a frequency-based kernel (SKAT) (see Figure 1). For scenarios with weaker effects and/or smaller sample sizes, POKEMON showed much better performance relative to other methods.

Additionally, to evaluate POKEMON's ability to identify a dispersed pattern, we simulated the scenario where variants are distributed on the protein's surface. When all influential variants on the surface had small odds ratios, none of the methods performed well. When increasing the odds ratio to 1.5, POKEMON outperformed other methods in most scenarios, except for cases where the percentage of influential variants was low (0.3) (see Figure S1).

We also assessed POKEMON's power at a higher resolution for different configurations of core odds ratios and influential variant proportions. Figure 2 illustrates the dynamics of statistical power for the POKEMON test under the assumption of a spatial effect. POKEMON achieved a power of 0.8 with study designs commonly found in sequencing studies of complex disease: a population of 3000 cases/3000 controls, the core odds ratio of 3.0, and 50% of the rare variants influential on the simulated phenotype with moderate effect. However, as expected, when the percentage of influential variants is low (<35%) and the core variant odds ratio is small (<1.8), POKEMON did not reach 80% power. A small core odds ratio and a low percentage of influential variants are more challenging for POKEMON to assess because more control subjects will carry variants located within the cluster region, making POKEMON less likely to identify associated patterns.

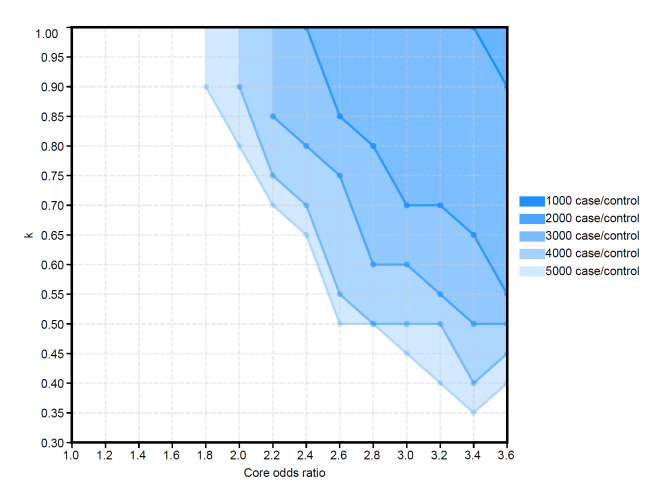


Figure 2. Power assessment for POKEMON at different configurations (for both structure kernel and combined kernel). Each line represents the minimum percentages of influential variants (k) and minimum core variant odds ratios to reach a power of 0.8 when the number of cases/controls is fixed.

POKEMON replicates the cancer-related spatial clusters from the TCGA dataset

To demonstrate POKEMON's ability to identify spatial patterns from real-world data, we analyzed germline variants from The Cancer Genome Atlas (TCGA), which has previously been evaluated for spatial clusters associated with cancer risk and metastasis (Mashl et al., 2018). We constructed a case/control dataset by combining 10389 subjects from TCGA across 33 cancer types with 4919 presumably cancer-free controls from the ADSP WES Discovery Dataset. We restricted our POKEMON analysis to rare variants with unknown effects and 31 proteins with functional assessement in the literature. This analysis directly tested the hypothesis that cancer-related variants tend to cluster in a protein hotspot while rare variants from cancer-free subjects are randomly distributed. We observed significant enrichment of statistical associations within the 31 proteins evaluated (27 with p < 0.05, see Supplementary Table 1).

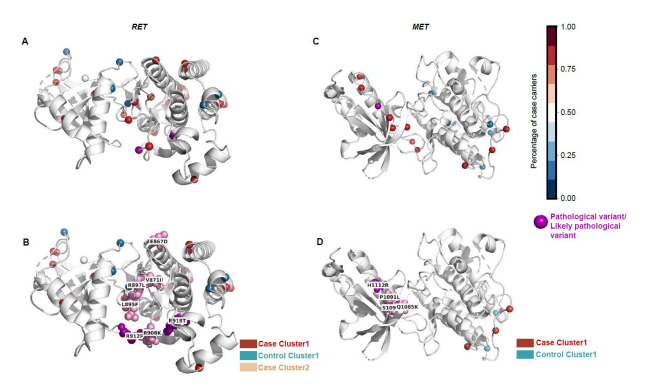


Figure 3. Spatial distribution of variants from TCGA dataset within MET (PDB:1R0P) and RET (PDB:2IVT). A and C show the rare missense variants with unknown effects mapped to the structure. The color scale indicates the percentage of case subjects that carry the variants out of the overall sample. Pathological and likely pathological variants are highlighted in purple. B, D show clusters identified by POKEMON. Clusters that are consistent with the original literature are highlighted with pink sphere models. Pathological and likely pathological variants are highlighted with purple sphere models.

From these results, we focus specifically on two genes highlighted in the literature that have been experimentally validated, namely *RET* and *MET* (Table 1). We found similar patterns of variant clustering for *RET* and *MET*, formed by somatic variants and pathological/likely pathological germline variants (Mashl et al., 2018). For *MET*, POKEMON identified a cluster formed by P1091L, S1092G, and Q1085K, which surrounds the pathological variant H1112R (Figure 3A and B). For *RET*, POKEMON identified a cluster formed by E867D, V871I, L895F, R897L, and R908K, which surrounds the pathological variants R912P and R918T (Figure 3C and D). Notably, POKEMON identified these two clusters via case/control analysis of rare germline variants while excluding known pathological variants. Thus our significant association statistic is driven by additional rare variants within *MET* and *RET* surrounding those with known pathological effects.

Table 1: Results for MET and RET from TCGA dataset

Gene	PDB entry	Phenotype	# SNPs	#SNPs mapped	p-value
MET	PDB:1R0P	cancer	757	17	5.454e-04
RET	PDB:2IVT	cancer	765	33	1.606e-04

POKEMON identifies known AD risk genes (*TREM2* and *SORL1*) and novel candidate genes (*CSF1R* and *DUSP18*)

To discover any spatial rare variant patterns associated with AD, we applied POKEMON with a structure kernel to the ADSP WES Discovery Dataset with 5,522 AD cases and 4,919 controls. We perform the POKEMON test on 4,173 genes with available protein structures and \geq 5 rare missense variants (MAF<0.05). *APOE* £2 and £4 dosages were included as covariates.

We used two significance thresholds to identify candidate genes: a typical Bonferroni correction threshold and an empirical threshold we derived from the *MET* and *RET* results based on our TCGA analysis which reflects the order of magnitude of significance of a true-positive signal obtained from variants with unknown effects and extremely low allele frequency (<0.001).

Overall, there are four genes out of 4,173 identified as candidate genes. *TREM2* was identified with the Bonferroni correction, while *SORL1*, *CSF1R*, and *DUSP18* were identified with the empirical threshold (<1E-03).

Table 2: genes associated with AD based on structure kernel

Gene	# SNP	# SNP mapped	PDB Entry	p-value	log(p-value)
TREM2	40	33	PDB:6XDS	1.261E-08	7.899
SORL1	203	56	PDB:3WSY	8.167E-05	4.088
CSF1R	74	38	PDB:4LIQ	4.428E-04	3.354
DUSP18	25	18	PDB:2ESB	6.717E-04	3.509

The spatial cluster is stable for SORL1, while TREM2 is driven primarily by a single variant.

As with our TCGA analyses of *RET* and *MET*, both *SORL1* and *TREM2* harbor known AD-assocated variants. To determine if the cluster pattern we detected is stable even in the absence of these known effects, we excluded any AD-related variants previously identified in GWAS studies leaving only rare genetic variants with unknown effects within *SORL1* and *TREM2*. A significant result from this analysis indicates that rare additional variants within these genes contribute to AD risk.

Indeed, for *SORL1*(Ensembl: ENST00000260197; PDB: 3WSY), even though AD-related variants A528T (Overall MAC:439; MAF:0.0210), E270K (Overall MAC:990; MAF:0.0474), and T947M (Overall MAC:2; MAF: 9.578e-05) were excluded respectively (Vardarajan et al., 2015), the

signals persist. While the results for the structure kernel with these loci excluded are still significant, *SORL1* is not significantly associated with AD even with all variants included for the frequency kernel analysis (Table 3a). The result indicates that the spatial pattern of variants within the 3WSY structure of *SORL1* is associated with AD.

For *TREM2* (Ensembl: ENST00000373113; PDB: 5ELI), The signal is likely driven primarily by the variant R47H. The results for *TREM2* from the structure kernel test are comparable to that from the frequency kernel test before and after R47H being excluded. Excluding R47H (Korvatska et al., 2015) changes the p-value drastically for both the structure kernel and the frequency kernel (Table 3b).

Table 3a: Results for SORL1 w/ and w/o known loci

Method	Gene	PDB Entry	p-value				
	SORL1		8.167E-05				
Structure kernel	SORL1(exclude A528T)		2.841E-02				
Fraguency kernel (SKAT)	SORL1	PDB:3WSY	0.0923				
Frequency kernel (SKAT)	SORL1(exclude A528T)		8.578E-01				
Structure kernel	SORL1		8.167E-05				
Structure kerner	SORL1(exclude E270K)		7.373E-05				
Frequency kernel (SKAT)	SORL1	PDB:3WSY	0.0923				
Frequency kerner (SKAT)	SORL1(exclude E270K)		3.881E-02				
Structure kernel	SORL1		8.167E-05				
Structure kerner	SORL1(exclude T947M)		8.166E-05				
Frequency kernel (SKAT)	SORL1	PDB:3WSY	0.0923				
Frequency kerner (SKAT)	SORL1(exclude T947M)		9.214E-02				

Table 3b: Results for TREM2 w/ and w/o known loci

Method	Gene	PDB Entry	p-value
Structure kernel	TREM2		1.261E-08
	TREM2(exclude R47H)		8.982E-04
	TREM2	PDB: 6XDS	5.218E-09
Frequency kernel (SKAT)	TREM2(exclude R47H)		3.501E-03

Gene associations passing the significance threshold(0.05) are highlighted in bold

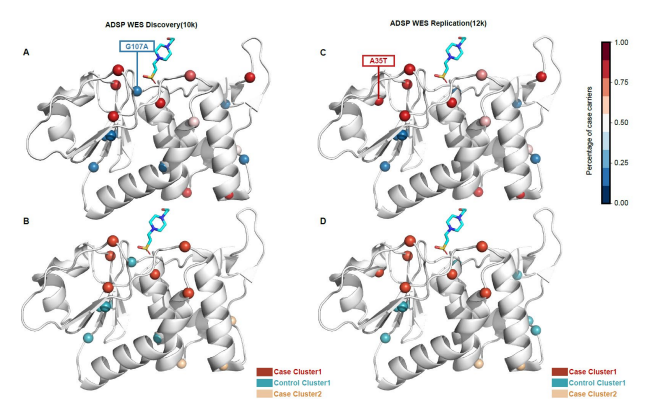


Figure 4. Spatial distribution of variants from ADSP WES discovery and replication dataset within DUSP18 (PDB:2ESB.A). A and C show the rare missense variants with unknown effects mapped to the structure. The color scale indicates the percentage of case subjects that carry the variants out of the overall sample. Variants that show a different percentage of case carriers and are being classified into other clusters between ADSP WES discovery and replication dataset are highlighted and labeled. B, D show clusters identified by POKEMON. Clusters are colored differently.

<u>DUSP18</u> has a cluster of variants primarily shared by case subjects around the ligand-binding domain

The signal identified within *DUSP18* (Ensembl: ENST00000334679; PDB: 2ESB) is driven by a cluster of variants primarily shared by case subjects in the catalytic domain for ligand binding, highlighted as Case Cluster 1 in Figure 4. This association is strengthened in the replication dataset (Table 4), where the cluster is better formed by an additional case variant A35T and the deletion of a control variant G107A (Figure 4). Overall, Case Cluster 1 is formed by missense

variants A35T, N51S, E55G, S74A, R78C, and R110H, which surround the ligand-binding site (Figure 5). Indeed, Variant R110H, a catalytic triad residue, interacts with E55G (Jeong et al., 2006), indicating a functional interaction related to AD risk. *DUSP18* has been previously reported to inhibit SUMOylation and reduce *ATXN1* aggregation (Ryu and Lee, 2018), and *ATXN1* loss of function is associated with an increased risk for AD (Suh et al., 2019).

Table 4: Results for candidate genes from the replication dataset

		ADSP WES Discovery(10k)			ADSP WES replication (12k)		
		Overall: 10441		Overall: 11828			
		Case/Control: 5522/4919		Case/Control: 6238/5590			
Gene	PDB entry	# SNPs	#SNPs mapped	p-value	# SNPs	#SNPs mapped	p-value
TREM2	PDB:6XDS	40	33	1.261E-08	61	29	5.049E- 11
SORL1	PDB:3WSY	203	56	8.167E-05	446	59	9.754E- 06
CSF1R	PDB:4LIQ	74	38	4.428E-04	196	38	6.128E- 03
DUSP18	PDB:2ESB	25	18	6.717E-04	37	19	1.037E- 05

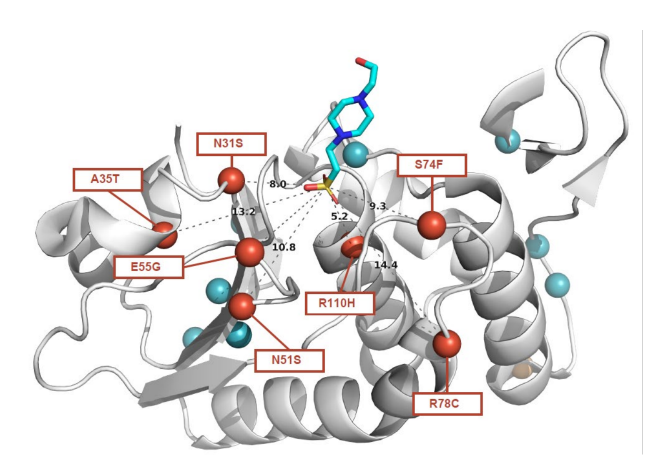


Figure 5. Spatial distribution of variants and their distance to the EPE within DUSP18 (PDB:2ESB.A). The distance is calculated from $C\alpha$ to S. All the variants that belong to the cluster of interest are highlighted and labeled with the AA change.

The spatial pattern within CSF1R is unstable through the replication dataset

For *CSF1R* (Ensembl: ENST00000286301; PDB: 4LIQ), the structure kernel result is more significant than the frequency kernel (Table 5), indicating that spatial patterns within the protein improve detection power. A case cluster identified by POKEMON in the Discovery Dataset contains T163A, N153H, A123G, L111I, R106Q, R106W, Q77E, and P53L (Figure 6.B: Case Cluster 1). The distances between those variants range from 5.3 to 14.3 Å, while their average sequence distance is 18 amino acids. This cluster is within the extracellular region (AA 1-498), where CSF-1 or IL-34 binds (Stanley and Chitu, 2014).

Table 5: Results for *CSF1R* with different kernels

Method	Gene	PDB Entry	p-value
Structure kernel	CSE1D	PDB:4LIQ	4.428E-04
Frequency kernel (SKAT)	CSFTK		7.731E-03

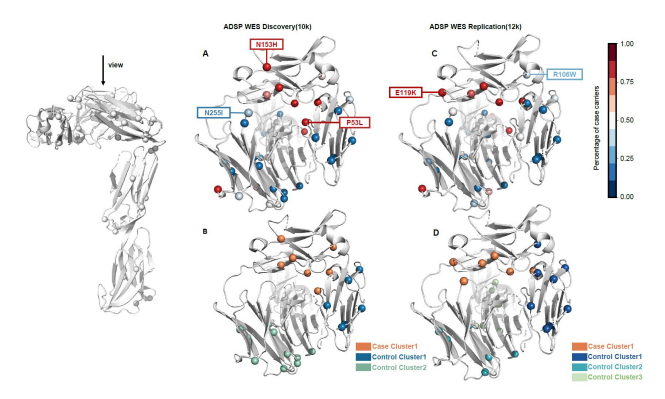


Figure 6. Spatial distribution of variants from ADSP WES discovery and replication dataset within CSF1R (PDB:4LIQ.A+E). A and C show the rare missense variants with unknown effects mapped to the structure.

The color scale indicates the percentage of case subjects that carry variants from the overall sample. Variants that show a different percentage of case carriers and are classified into other clusters (illustrated by color) between ADSP WES discovery and replication dataset are highlighted and labeled. B, D show clusters identified by POKEMON.

However, when we examined *CSF1R* with the replication dataset, we did not observe an improved signal compared to the Discovery Dataset (Table 4). Reviewing the PDB:4LIQ structure, we found that Case Cluster 1 persists in the replication dataset while other clusters become sporadic. Specifically, Case Cluster 1 has two additional case variants E119K and N255I, while two case variants P53L, N153H drop out of the dataset (Figure 6.A, C). The complete comparison of variants between the Discovery Dataset and the Replication dataset can be found in the Supplementary Tables.

Discussion

We have shown that POKEMON improves the power to detect rare variant associations in the context of protein structure. We found POKEMON outperforms other structure-based methods through simulation studies except in a small number of cases where all existing methods have insufficient power. Specifically, POKEMON achieves a power of 0.8 for what we presume to be a common scenario in sequencing studies: a study population of 3,000 cases/3,000 controls, a core odds ratio of 3, and 50% of the variants associated with the phenotype. In contrast, all other methods tested for this scenario have power below 0.6. We applied POKEMON to the ADSP Discovery WES dataset and identified spatial patterns of rare variants related to AD risk. The spatial patterns of variants within *SORL1*, is consistent with a previously reported association using PSCAN(Tang et al., 2020). We also identify two potentially AD-associated clusters of variants within *CSF1R* and *DUSP18*, located around ligand-binding sites. Specifically, the cluster within *DUSP18* is validated with the replication dataset with a larger sample size.

Notably, an advantage for POKEMON over other rare-variant analysis methods is that statistical power increases with the observation of any new variant including singletons assuming the existence of divergent spatial patterns between cases and controls. In most rare variant association tests, increasing sample size only increases the power for non-singleton variants in the resulting data. Even for those non-singleton variants, the improvement in power is not necessarily proportional to the increase in sample size. Moreover, additional neutral variants will be introduced and negatively impacting the statistical power when the sample size increases. In

contrast, POKEMON can utilize rare variants and even singletons with the structure kernel, regardless of their low allele frequency. The increasing number of rare variants helps form a spatial pattern, which can be identified by POKEMON with higher power (Figure S4D).

Based on our analyses of *MET/RET* in the TCGA dataset, and SORL1 in the ADSP WES Discovery dataset, we also demonstrated that the association detected by our spatial kernel is not driven by a single variant but rather a collection of variants with modest effects. Additionally, even though we didn't include any population-related covariates in the structure kernel test for ADSP WES Discovery Dataset, the overall results didn't show large genomic inflation (GC=1.23). This confirms our assumption that the structure kernel in POKEMON is less susceptible to population stratification than frequency-based tests; any constraint to the positions of rare variants within protein structures is likely independent of the variants' population origin and therefore does not confound analyses as is typical of a frequency-based test.

POKEMON is designed to leverage pre-existing biological information for sequencing datasets where typically only variant counts or frequencies are considered. Even though protein structure information of variants has been incorporated into association tests like POINT and PSCAN (Marceau West et al., 2019; Tang et al., 2020), they serve as guiding information for more traditional association tests ultimately based on allele frequency. Therefore, these approaches are still potentially subject to the limitations in unit-based or single variant tests. With the structure kernel, POKEMON uses the spatial information of a missense variant, which is independent of allele frequency. Assuming the rare variants form spatial patterns, POKEMON ameliorates the power dilemma induced by increasing numbers of singleton variants as the sample size of sequencing studies increases.

Importantly, POKEMON does not have sufficient power to detect patterns for some scenarios. As shown in the simulations, POKEMON lacks the power even with 5,000/5,000 cases/controls when the percentage of influential variants is low (<30%) and the core variant odds ratio is small (<1.5). While we expected lower power in this scenario, this result may also reflect a limitation of our simulation strategy; we limit the number of variants simulated to 50. If more variants are included in the test, the underlying spatial pattern is better formed, and thus the power of POKEMON will improve.

We anticipate POKEMON will be helpful as a large-scale screening method to detect potentially disease-associated proteins in a proteome-wide fashion. Currently, the PDB structures deposited in the PDBank only cover ~70% of the identified molecular functions in the human genome (Somody et al., 2017). We expect that the improvement in Cryo-EM and prediction methods like

AlphaFold2 (Senior et al., 2020) will massively increase the availability of structural information for proteins and complexes. As POKEMON is a unit-based test, it only provides a single association statistic for the influence of all missense variants within the protein for a phenotype. Follow-up analyses to assess specific SNVs or refine SNV subsets may provide more detailed quantitative assessments of specific variant spatial patterns.

MATERIAL AND METHODS

Derivation of the POKEMON method

We briefly review the linear mixed model used in association tests and then introduce the construction of a structure kernel for POKEMON. Assume we have n individuals for whom we have p non-genetic covariates, genotypes for m SNPs, and the phenotype p as an p and p vector. Genotype p is a p matrix. Covariate p is a p matrix.

A linear mixed model contains a fixed effect from covariates $X\beta$, a random effect annotated by Zu, and an error term ϵ . The y is fit with a high-dimensional normal distribution (2). The random effect can be further divided into two parts, an environmental effect $\sigma_e^2 I$ and a genetic effect $\sigma_1^2 K_g$. K_g is the kernel containing the genetic similarity between individuals. σ_1^2 is the amount of variance of y explained by K_g .

$$y = X\beta + Zu + \epsilon (1)$$
$$y \sim N(X\beta, \sigma_1^2 K_a + \sigma_e^2 I) (2)$$

The null hypothesis $\sigma_1=0$ indicates that K_g does not explain any variance of y. The score statistic Q is defined as the partial differential for the log-likelihood on σ_1^2 . Under the null hypothesis, Q follows a mixed chi-squared distribution(3), where S projects y into a space orthogonal to covariates and λ_i are the eigenvalues of SK_gS .

$$\frac{Q}{\sigma_e^2} = y^T \mathbf{S} \mathbf{K}_g \mathbf{S} y \sim \sum_{i=1}^n \lambda_i \chi_1^2(3)$$

For POKEMON, we construct the $n \times n$ kernel K_g in the context of protein as follows. For K_g , each entry is the genetic similarity between individuals based on the variants they carry, which is weighted by variants' distance in the protein structure(4). d_{kl} is the distance of pair-wise single nucleotide variants (SNVs) in angstroms (Å) within the protein. k and l represent the kth variant from individual i and the lth variants from individual j.

$$K_{ij} = \sum_{i_k j_l} A_k A_l min\{f(d_{kl})\}$$
(4)

Some protein structures are formed by identical subunits (homo-multimer), which introduces redundancy in the variant-to-amino acid projection (i.e., one variant can map to multiple amino acids located in different subunits). To eliminate the spatial similarity induced by multiple mapping locations of a single variant in a homo-multimer, we took d_{kl} to be the minimum distance over all pair-wise distances. Function f(d) converts a Euclidean distance to the similarity score for a pair of variants.

$$f(d_{kl}) = e^{-\frac{d_{kl}^2}{2t^2}} (5)$$

As a default, the exponential function for f in (5) with t set to a value of 7 Å, with which the effect decays below 0.1 after 2t (14 Å). 14 Å was chosen as it is a commonly adopted short-range non-bonded cut-off in molecular dynamic simulation (Monticelli et al., 2008).

Apart from spatial patterns, we also account for the magnitude of the protein change due to the different amino acid substitutions. We scaled the pair-wise variants by their amino acid substitution, which is defined as A_k and A_l . A_k and A_l are the weights for amino acid substitution for variant k and variant k according to the BLOSUM62 matrix (Henikoff and Henikoff, 1992), respectively. For a less conservative amino acid substitution, the score s_k in BLOSUM62 matrix will be negative and consequently a_k will be greater than 1. In contrast, for a neutral or conservative amino acid substitution, s_k will be positive and a_k will be less than 1.

$$A_k = -\sqrt{e^{s_k}}$$
 (6)

The structure kernel is nonlinear in contrast to the SKAT tests (Wu et al., 2011), which uses a linear kernel (e.g., K = GWW'G') to calculate the genetic similarity between individuals. The genetic similarity in a linear kernel between individuals is the sum of weighted SNVs being shared. However, singletons are carried by only a single individual and thus fail to be included in calculating genetic similarity. With the structure kernel, a pair of singleton variants will be assigned non-zero weights if they are spatially proximate in the protein structure. The interpretation of the structure kernel is that case individuals are genetically similar because they share more spatially clustered or dispersed rare variants than the control individuals.

We also allow for incorporating allele frequency in the POKEMON test and develop a combined kernel function. Variants clustered in protein structure already contribute to a high genetic similarity based on our structure kernel. With a *combined* kernel, those variants will be further upweighted if they are rare in allele frequency and vice versa. The combined kernel function is based on K_q , extended by further scaling variants by weights derived from the allele frequency. $w_k =$

Beta(MAF_k ; a, b) is the weight for the k^{th} variant characterized by beta density with a=1 and b=25 as default.

$$K_{ij} = \sum_{i_k j_l, k \neq l} A_k A_l min\{f(d_{kl})\} + \sum_{i_k j_k} w_k A_k^2$$
 (7)

Workflow of POKEMON

POKEMON requires a genotype matrix and consequence profile containing variant-to-amino acid mapping information as inputs (see Figure S2). Additional covariate files are optional to adjust for covariates. POKEMON first maps the variants using their coordinates into the 3D protein, which is accomplished with the consequence profile generated by Ensembl Variant Effect Predictor (VEP v95) and the reference from SIFTS function mapping a PDB entry to a UniProt residue level (Dana et al., 2019). A single variant may be mapped to multiple amino acids for multimers with identical subunits. The protein structures are fetched from the PDB during the analysis. If multiple protein structures are available for a single gene, the structure with the most variants mapped will be selected. The user may also specify a specific PDB entry. After mapping, the score between a pair of variants is calculated based on the minimum distance between them, which is further scaled by the amino acid substitution weight from the BLOSUM62 matrix by default. The pair-wise genetic similarity between individuals is the summation of all pair-wise scores of variants. The genetic similarity kernel K_g is then evaluated in the variance component test.

Data Simulation

We conducted simulation studies to assess POKEMON's power in detecting disease-associated protein variant patterns. We hypothesized that variants with moderate effects on a phenotype form spatial patterns within a protein structure and induce subtle alterations to the protein's function. To test the hypothesis, we established two patterns. The first pattern entails an embedded core within the protein disrupted by rare variants (i.e., variant clustering), while the other represents the localization of influential variants to the protein's surface (i.e., variant dispersion). Both patterns are shown in Fig S4.A & Fig S4.B. We randomly selected a protein structure, Human c-Fms Kinase Domain (PDB:2OGV), to carry out simulations. Structural information for PDB:2OGV is available for both PSCAN and SKAT.

We simulated a clustering pattern by distributing influential variants within the core of the protein structure and scaling the variant odds ratios proportionally to their distance from the core. We then randomly sampled 50 variants from the protein. The minor allele frequencies for all the variants were randomly sampled from a log-transformed uniform distribution within an interval (-4, -2.3). This variant sampling strategy restricted the selected minor allele frequencies within the range (0.0001, 0.005) and generates singletons, which is consistent with ADSP WES studies (Figure S3). To investigate how neutral variants influence the power, we varied the percentage of influential variants out of all variants being sampled (Figure S4). For each set of parameters (e.g., sample size, core variant odds ratio, etc.), the empirical power was estimated by the percentage of successful tests out of 100 independent tests with a significance level of 0.05. We compared the empirical power of POKEMON with three other methods: SKAT, PSCAN-V, and POINT. The number of case and control subjects sampled is from 1000 to 5000. Additional details for the simulation can be found in Figure S4 and supplementary materials.

We also simulated a dispersion pattern by distributing influential variants on the protein's surface. Considering the selected protein PDB:2OGV is about 40 Å in diameter, we defined the surface variants as those more than 21 Å away from the core, which yielded 33 variants. All the surface variants were assigned with the same odds ratio (e.g., 1.1), while the rest were considered neutral with an odds ratio of 1. The simulation settings were similar to the clustering pattern, with the only difference that we sampled 30 variants from the protein, which allowed us to tune the percentage of influential variants to as large as 90%.

Applying POKEMON to ADSP WES data

Discovery Dataset

We used the whole-exome sequencing (WES) data from the Discovery Case-Control study under the Alzheimer's Disease Sequencing Project (ADSP). ADSP WES data contains 5,740 late-onset AD cases and 5,096 cognitively normal controls primarily of European ancestry, with 218 cases and 177 controls of Caribbean Hispanic ancestry. Cases were determined based on diagnosis using cognitive testing data and medical records, while controls were determined on their low risk of developing AD by age 85 years (Bis et al., 2018; Beecham et al., 2017). ADSP WES data is available by applying to the NIAGADS Data Sharing Service.

We selected 10,441 subjects of European ancestry from the ADSP as the study group (5,522 late-onset AD cases and 4,919 cognitively normal controls). The whole-exome sequencing for these 10,441 subjects provided \sim 850,000 variants, of which 97.5% have a minor allele frequency < 0.01. We retained the missense variants with minor allele frequency < 0.05 for our assessment. Overall, we selected 4,173 genes with experimentally determined protein structures and \geq 5 rare

missense variants mapped to the structure. The mean number of rare missense variants mapped per gene was ~50. To exclude signals induced by the well-known *APOE* association, we included *APOE* £2 and £4 dosages as covariates.

Replication Dataset

The ADSP FUS contains the 9389 subjects from the Discovery Phase Case-Control study plus additional 1044 cases and 1395 controls for a total of 11828 non-Hispanic white subjects. The WES data in FUS was reprocessed using joint genotype calling approaches implemented in the VCPA pipeline (Leung et al., 2019). The genotype calling approaches for the replication dataset were updated from the ATLAS genotype calling process implemented for the Discovery Dataset. Therefore, we consider that this replication dataset is a validation by expanding the sample size and accounting for variability in the variant calling process. If the spatial pattern POKEMON identified is stable, the additional variants identified in the replication dataset should also contribute to the association signal.

Our primary purpose for using the replication dataset was to validate the spatial patterns identified from the discovery analysis, there for we examined only *SORL1*, *TREM2*, *DUSP18*, and *CSF1R* genes. Tests on the replication dataset were conducted similarly to the discovery analyses, with *APOE* £2 and £4 dosages included as covariates to regress out the *APOE* association.

Applying POKEMON to TCGA data

The TCGA data is a real-world, true-positive example of spatial patterns of missense variants associated with phenotypes (Kamburov et al., 2015). To create a dataset in the form of a case-control study, we combined 4919 control subjects from the ADSP WES Discovery Dataset and 10389 subjects from TCGA data diagnosed with 33 cancer types (Mashl et al., 2018). We assumed that 4919 control subjects from the ADSP WES Discovery Dataset are cancer-free controls. While this is not an ideal study design, any violation of this assumption would reduce statistical power rather than identifying spurious associations. The combined case/control dataset provided a real-world assessment of our hypothesis that rare variants from cancer tissues would form spatial patterns, while the rare variants from control subjects would be randomly distributed within the protein.

All previously identified pathological or likely pathological variants were excluded from the TCGA data. Moreover, we set a stringent MAF threshold as <0.001 to retain rare variants. In summary, the entire test was carried out to examine rare variants with unknown effects. We carried out POKEMON tests on 31 genes with potential hotspots (Mashl et al., 2018) and available protein structures. We only applied structure kernel tests with no covariate included.

CODE AVAILABILITY

The code for this study is available at GitHub: https://github.com/bushlab-genomics/POKEMON with open access.

DATA ACCESS

Whole Exome Sequencing Data from the Alzheimer's Disease Sequencing Project are available via NIAGADS (NG00067 - ADSP Umbrella). https://dss.niagads.org/datasets/ng00067/NIAGADS Data Sharing Service is needed to access the data.

The Cancer Genome Atlas data are available via dbGaP Study Accession (phs000178.v1.p1) and accessible via the National Cancer Institute Genomic Data Commons (https://gdc.cancer.gov/access-data/obtaining-access-controlled-data)An application to the

COMPETING INTEREST STATEMENT

The authors declare no competing interests.

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