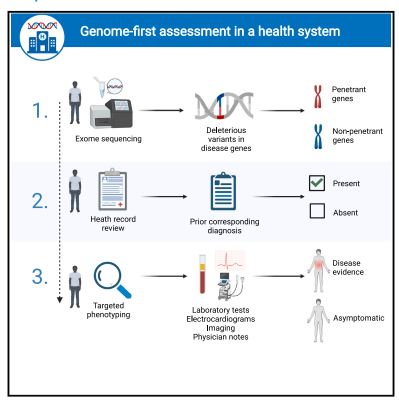
# Genome-first evaluation with exome sequence and clinical data uncovers underdiagnosed genetic disorders in a large healthcare system

### **Graphical abstract**



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### In brief

Forrest et al. pilot a genome-first strategy for assessing individuals genetically predisposed to disease who may be missing a diagnosis in their routine healthcare. Participants carrying diseaserisk variants are identified and undergo targeted phenotyping with laboratory measurements, electrophysiology, imaging, and physician notes to uncover new diagnoses in previously undiagnosed individuals.

### **Highlights**

- Persons with disease-risk variants and exome and clinical phenotype data are evaluated
- 3 in 4 observations of variants lack a corresponding diagnosis, 15% of which show symptoms
- Exome analysis enables targeted phenotyping and new diagnoses missing from clinical care







### **Article**

# Genome-first evaluation with exome sequence and clinical data uncovers underdiagnosed genetic disorders in a large healthcare system

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#### **SUMMARY**

Population-based genomic screening may help diagnose individuals with disease-risk variants. Here, we perform a genome-first evaluation for nine disorders in 29,039 participants with linked exome sequences and electronic health records (EHRs). We identify 614 individuals with 303 pathogenic/likely pathogenic or predicted loss-of-function (P/LP/LoF) variants, yielding 644 observations; 487 observations (76%) lack a corresponding clinical diagnosis in the EHR. Upon further investigation, 75 clinically undiagnosed observations (15%) have evidence of symptomatic untreated disease, including familial hypercholesterolemia (3 of 6 [50%] undiagnosed observations with disease evidence) and breast cancer (23 of 106 [22%]). These genetic findings enable targeted phenotyping that reveals new diagnoses in previously undiagnosed individuals. Disease yield is greater with variants in penetrant genes for which disease is observed in carriers in an independent cohort. The prevalence of P/LP/LoF variants exceeds that of clinical diagnoses, and some clinically undiagnosed carriers are discovered to have disease. These results highlight the potential of population-based genomic screening.

### **INTRODUCTION**

A major endeavor of precision medicine is to leverage genetic data to improve the diagnosis and risk stratification of genetic disorders. <sup>1,2</sup> Most genetic tests are applied to Mendelian disorders driven by a single gene mutation, such as familial breast cancer caused by a deleterious mutation in *BRCA1* or *BRCA2*. <sup>3,4</sup> The American College of Medical Genetics and Genomics has released guidelines for reporting secondary findings from clinical genomic sequencing. <sup>5</sup> These vetted disease-predisposition genes are clinically actionable, but their use in population screening has not yet been determined. <sup>6</sup> In addition, ClinVar <sup>7</sup> and ClinGen <sup>8</sup> advance the clinical interpretation of variants with annotations of pathogenicity, including pathogenic/likely pathogenic (P/LP), which is useful for genomic screening, since P/LP variants can be selected.

The increasing use of exome sequencing has permitted an assessment of the prevalence of P/LP variants in the population for numerous genetic diseases. <sup>9,10</sup> Notably, the allelic prevalence of P/LP variants has been shown to exceed the estimated

disease prevalence attributed to P/LP variants. <sup>11,12</sup> This indicates at least three possibilities: (1) incompletely penetrant variants, in which the presence of a variant does not always result in disease; <sup>13</sup> (2) underdiagnosed disease, where an individual with the variant expresses disease but is not clinically diagnosed; or (3) a combination of the two. In familial hypercholesterolemia, causative genetic variants may be highly penetrant, and the disorder is often underdiagnosed and undertreated. <sup>14</sup> Monogenic cardiac diseases have also been examined recently for underdiagnosis; a study of Noonan syndrome found that 67% of individuals with P/LP variants in *PTPN11* had a probable missed clinical diagnosis, while a study of hereditary transthyretin amyloidosis-induced heart failure showed that just 10 of 67 individuals with *TTR* V112I and disease had a clinical diagnosis. <sup>15,16</sup>

Population-based genomic screening can help fill the gaps in the clinical diagnosis of genetic disorders. The identification of individuals carrying clinical variants in known disease predisposition genes leads to targeted phenotyping and can improve diagnostic yield relative to a genetic-agnostic approach.<sup>17,18</sup> This genome-first strategy, defined as identifying variants first



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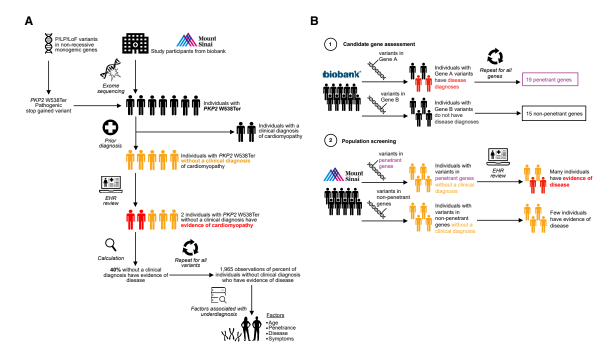


Figure 1. Schematic of genome-first assessment of pathogenic/likely pathogenic or predicted loss-of-function (P/LP/LoF) variants in the BioMe Biobank at Mount Sinai and UK Biobank

(A) Study design for genome-first evaluation in the BioMe Biobank at Mount Sinai using the plakophilin 2 (*PKP2*) W538Ter variant as an example. A total of P/LP/LoF variants in non-recessive monogenic genes for nine genetic disorders were curated: (1) variants reported as P/LP in the ClinVar repository with a minimum review status of two (multiple submitters) and previously unreported LoF variants identified by Variant Effect Predictor were obtained, and (2) non-recessive genes with a monogenic disease predisposition were identified from Online Mendelian Inheritance in Man and corroborated with literature review. A total of 29,039 participants with exome sequence and electronic health record (EHR) data were included in the study. This yielded 644 observations of 303 P/LP/LoF variants in 614 individuals. As an example, *PKP2* W538Ter was identified in seven individuals, of whom two had a prior clinical diagnosis of cardiomyopathy in the EHR. The remaining five individuals lacked a clinical diagnosis, of whom two (40%) were discovered to have EHR evidence of cardiomyopathy. This procedure was repeated for all variants to produce a dataset of the percentage of clinically undiagnosed observations that had evidence of disease, which was used to assess factors associated with the presence of disease in clinically undiagnosed individuals. Factors comprised the gene containing the variant, age of individuals, disease, and symptoms.

(B) Evaluation of clinically undiagnosed but symptomatic individuals with P/LP/LoF variants in BioMe Biobank by gene penetrance observed in UK Biobank. A total of 34 target genes were identified in an independent cohort from UK Biobank, for which disease was either observed in individuals with P/LP/LoF variants in the gene (19 penetrant genes) or not observed (15 non-penetrant genes). The proportion of clinically undiagnosed observations with disease evidence in BioMe Biobank was then compared between the penetrant genes and non-penetrant genes.

and then evaluating relevant phenotypes, <sup>19</sup> was adopted in recent studies to detect hereditary breast and ovarian cancer syndrome, Lynch syndrome, and familial hypercholesterolemia (tier 1 conditions designated by the Centers for Disease Control and Prevention [CDC]) in individuals with no prior diagnosis. <sup>20,21</sup> However, the value of genomic screening and extent of underdiagnosis across most genetic disorders are uncertain. Moreover, prior studies have not addressed the selection of genes and age of individuals carrying variants. Variants conventionally classified as P/LP do not always cause disease (incomplete penetrance), and penetrance estimates depend on individuals' age (i.e., age-dependent penetrance); <sup>13,22</sup> obtaining variants that are observed to occur with disease in real-world populations is important for conducting genome-first assessments but has not been studied.

Here, we asked whether exome sequencing in a large healthcare system has diagnostic utility in individuals lacking clinical diagnoses from routine care. We used a genome-first approach to evaluate individuals with clinical variants on a systematic level for monogenic disorders in a cohort of 29,039 participants from an electronic health record (EHR)-linked clinical care biobank. First, individuals harboring P/LP or predicted loss-of-function (LoF) variants in known disease predisposition genes from a prior study<sup>13</sup> were identified, including many who lacked a clinical diagnosis for the disease corresponding to their variant. Second, clinically undiagnosed individuals were evaluated for EHR evidence of disease symptoms and findings informed by their genotype. This permitted the discovery of individuals with P/LP/LoF variants who had EHR evidence of symptomatic disease but were not medicated or clinically diagnosed with the disease and motivated an investigation into the genetic and clinical factors driving this phenomenon.

### **RESULTS**

### Summary of the study population and variants

The study design is shown in Figure 1. The study population included 29,039 individuals from the BioMe Biobank (BioMe)

### **Article**



Table 1. Baseline demographic traits and clinical diagnoses in a clinical care cohort of 29,039 individuals

Trait	All participants $(n = 29,039)$	Individuals with P/LP/LoF variants ( <i>n</i> = 614)
Age, mean (SD) years	59 (16)	59 (16)
Female, n (%)	17,353 (60)	1,143 (62)
Ethnicity, n (%)		
African	7,190 (25)	114 (19)
Asian	1,349 (4.6)	28 (4.6)
European	9,376 (32)	242 (39)
Hispanic	8,528 (29)	182 (30)
Other	2,596 (8.9)	48 (7.8)
Clinical diagnoses, n (%)		
Amyotrophic lateral sclerosis	7 (0.024)	1/21 (4.8)
Breast cancer	1,488 (5.2)	62/168 (37)
Cardiomyopathy	823 (2.8)	78/213 (37)
Colorectal cancer	218 (0.75)	2/19 (11)
Endometrial cancer	115 (0.40)	1/13 (7.7)
Hypercholesterolemia	4,460 (15)	5/11 (45)
Prostate cancer	474 (1.6)	2/2 (100)
Retinitis pigmentosa	15 (0.052)	3/175 (1.7)
Type 2 diabetes	7,044 (24)	1/1 (100)

Demographic traits and prevalence of clinical diagnoses are shown for all study participants and in a subset of individuals carrying variants reported as pathogenic/likely pathogenic in ClinVar with a minimum review status of two or variants of predicted LoF molecular consequence (P/LP/LoF variants). The denominator of the proportion of clinical diagnoses in the second column refers to the number of individuals carrying variants corresponding to the disease for each row (e.g., for cardiomyopathy, 213 individuals carry cardiomyopathy-predisposition variants, of whom 78 have a clinical diagnosis of cardiomyopathy). An overview of baseline traits among individuals clinically diagnosed with each disease in the electronic health record (EHR) is provided (Table S2). Clinical diagnoses were identified by International Classification of Diseases 10 (ICD-10) codes in the EHR. Ethnicity, self-reported ethnicity; other ethnicity, miscellaneous ethnicities other than those listed; *n*, number; SD, standard deviation.

with exome sequence and EHR phenotype data who passed quality control (STAR Methods). The mean age was 59 years (standard deviation [SD], 16 years); 17,353 (60%) were female; 7,190 (25%), 1,349 (4.6%), 9,376 (32%), and 8,528 (29%) were of African, Asian, European, and Hispanic ethnicities, respectively; and 614 (2.1%) had at least one P/LP/LoF variant (Tables 1, S1, and S2). The prevalence of clinical diagnoses ranged from >20% for common conditions (e.g., 7,044 [24%] diagnosed with type 2 diabetes) to <1% for rare disorders (e.g., 15 [0.052%] diagnosed with retinitis pigmentosa).

P/LP/LoF variants from a previous study<sup>13</sup> were curated for a set of nine genetic disorders: 303 P/LP/LoF variants in 54 genes corresponding to the nine diseases were identified in the exomes of 614 individuals, yielding 644 observations of variants in individuals (Tables S1 and S3). Among these individuals harboring P/LP/LoF variants, the mean number of unique P/LP/LoF variants per person was 1.0 (SD, 0.16; range, 1–2). The diseases

with the largest number of observations of variants in individuals were cardiomyopathy (n = 234 observations [36%]), retinitis pigmentosa (n = 175 [27%]), and familial breast cancer (n = 168 [26%]) (Table 2). Demographic traits for individuals carrying P/LP/LoF variants are summarized in Table 1. The mean age of individuals with P/LP/LoF variants was 59 years (SD, 16 years); 371 (60%) were female; and 114 (19%), 28 (4.6%), 242 (39%), and 182 (30%) were of African, Asian, European, and Hispanic ethnicities, respectively.

# Assessment of the clinical diagnosis and phenotype of individuals with P/LP/LoF variants

We systematically evaluated the clinical diagnosis and phenotype of individuals with P/LP/LoF variants by examining their EHR for diagnoses and traits of diseases expected with each of the variants they carried. These observations of variants in individuals were categorized into three distinct groups based on the presence or absence of a clinical diagnosis and symptoms of disease corresponding to the variant (STAR Methods; Tables S4, S5, and S6): (1) 157 of 644 observations (24%) had a clinical diagnosis of the disease corresponding to the variant recorded in the EHR. Of the 487 observations without a clinical diagnosis, (2) 412 (85%) did not have EHR evidence of disease, and (3) 75 (15%) had EHR evidence of disease (Figure 2). Of the 75 individuals in the last group, 40 (53%) had P variants, 3 (4%) had LP variants, and 32 (43%) had LoF variants. None of the individuals in the last group had received a medication specifically indicated for treatment of the disease (Table S5), and the findings were consistent across different ancestries (Table S7).

We focused on the third group—individuals with P/LP/LoF variants who had EHR evidence of symptomatic disease but were not clinically diagnosed or medicated—as the primary outcome of interest to assess underdiagnosis for most of the subsequent analyses. We investigated factors contributing to smaller or larger proportions of these underdiagnosed observations, including the gene containing the variant, age of individuals, and disease and symptoms corresponding to the variant.

### Clinically undiagnosed individuals with evidence of disease stratified by genes

Population genomic screening would ideally use variants in penetrant genes with disease observed in real-world populations to increase clinical yield. We tested this hypothesis by comparing the proportion of clinically undiagnosed observations with disease evidence in BioMe for 19 penetrant genes and 15 non-penetrant genes observed in UK Biobank (STAR Methods). The median proportion of clinically undiagnosed observations that had disease evidence was 25% (interquartile range [IQR], 33) for penetrant genes compared to 0% (IQR, 29) for non-penetrant genes ( $p = 9.7 \times 10^{-5}$ ). Using linear regression adjusted for age, we observed a 10-percentage point increase (SE = 4.3) in clinically undiagnosed observations that had disease evidence for penetrant genes compared to non-penetrant genes (p = 0.02). A larger proportion of penetrant genes had ≥20% observations with disease evidence compared to non-penetrant genes (odds ratio = 1.09 adjusted for age; 95% confidence



Table 2. Clinical diagnosis and phenotypes for 644 observations of pathogenic/likely pathogenic or loss-of-function (P/LP/LoF) variants in 614 individuals

Disease	Genes	Variants	Observations, n	+Dx, n (% of observations)	-Dx -Sx, $n$ (% of $-Dx$ observations)	-Dx +Sx, n (% of −Dx observations)
Amyotrophic lateral sclerosis	4	19	21	1 (4.8)	15 (80)	5 (20)
Cardiomyopathy	21	151	234	80 (34)	124 (81)	30 (19)
Colorectal cancer	7	16	19	2 (11)	15 (88)	2 (12)
Endometrial cancer	1	1	13	1 (7.7)	8 (67)	4 (33)
Familial breast cancer	4	64	168	62 (37)	83 (78)	23 (22)
Familial hypercholesterolemia	1	7	11	5 (45)	3 (50)	3 (50)
Monogenic diabetes	1	1	1	1 (100)	-	-
Prostate cancer	1	2	2	2 (100)	_	-
Retinitis pigmentosa	15	42	175	3 (1.7)	164 (95)	8 (4.7)

A total of 644 observations of 303 pathogenic/likely pathogenic or predicted LoF (P/LP/LoF) variants carried by 614 individuals were identified for nine diseases. Observations were first categorized by whether the disease corresponding to the variant was clinically diagnosed (+Dx) or not clinically diagnosed (-Dx) in the individual with the variant. Electronic health records (EHRs) of clinically undiagnosed individuals were then evaluated and categorized by whether evidence of symptomatic disease was present (+Sx) or absent (-Sx). This resulted in three distinct groups: (1) clinically diagnosed (+Dx), (2) no clinical diagnosis and no EHR evidence of disease (-Dx -Sx), and (3) no clinical diagnosis but EHR evidence of disease (-Dx +Sx). n (% of observations), number and percentage of the observations of individuals with P/LP/LoF variants; n (% of -Dx observations), number and percentage of the observations of individuals with P/LP/LoF variants; n0 disease predisposition genes containing the P/LP/LoF variants; n1 variants; n2 variants, number of P/LP/LoF variants; n3 not applicable, as all observations for monogenic diabetes and prostate cancer were diagnosed with the corresponding disease.

interval [CI], 1.03-1.17; p=0.004). These trends were consistent across three age groups of individuals who were at least 20, 40, and 60 years old (Figure 3).

Of medical importance, the American College of Medical Genetics and Genomics secondary findings (ACMG SF) v.3.1 genes are vetted disease predisposition genes with clinical actionability, but the clinical utility of population screening for individuals with variants in these genes has not been established.<sup>6</sup> We therefore assessed the diagnoses and phenotypes of individuals with P/LP/LoF variants in ACMG SF v.3.1 genes comprising 22 of 54 genes (41%) (Figure S1). A greater proportion of observations of ACMG SF v.3.1 gene variants in individuals had a corresponding clinical diagnosis for the expected disease than non-ACMG SF v.3.1 gene variants (134 of 396 [34%] versus 23 of 248 [9.3%],  $p = 1.6 \times 10^{-13}$ ). There was also a greater proportion of clinically undiagnosed observations with evidence of disease for ACMG SF v.3.1 gene variants (57 of 262 [22%]) compared to variants in non-ACMG SF v.3.1 genes (18 of 225 [8.0%]) ( $p = 2.6 \times 10^{-5}$ ).

### Underdiagnosis stratified by disease and symptoms

Next, we evaluated the phenotype of individuals without a clinical diagnosis carrying P/LP/LoF variants for each of the nine genetic disorders (Table 2; Figure 4). Genetic disorders are diverse, with different affected systems and clinical manifestations, which may be differentially detected and reported in a healthcare setting. The proportion of clinically undiagnosed observations with evidence of disease varied by genetic disorder, with a median of 22% (IQR, 14) across all nine disorders. The greatest proportion of clinically undiagnosed observations with evidence of disease was noted for familial hypercholesterolemia (3 of 6 [50%]), endometrial cancer (4 of 12 [33%]), amyotrophic lateral sclerosis (5 of 20 [25%]), breast cancer (23 of 106 [22%]), and

cardiomyopathy (30 of 154 [19%]). All observations of monogenic diabetes and prostate cancer were diagnosed. In terms of absolute numbers, cardiomyopathy had the most clinically undiagnosed observations, with evidence of disease with 30 such observations.

We reasoned that the symptoms associated with greater risk of hospitalization may be better detected and recorded in a healthcare setting and, therefore, more prevalent in our analysis of disease evidence. Hence, we investigated the association of symptom hospitalization risk and prevalence of symptoms detected as evidence of disease in clinically undiagnosed observations of variants in individuals, using a score<sup>23</sup> for adverse phenotypes ranging from 0 (lowest risk of hospitalization and death) to 1 (highest risk of hospitalization and death) (Table S8). There was a positive association between symptom score and prevalence of symptoms detected in observations lacking a clinical diagnosis with a 4.6-percentage point increase (95% CI, 2.6-6.6) in the prevalence of symptoms per 0.1 increase in symptom score ( $p = 6.7 \times 10^{-5}$ ) (Figure S2). The symptom with the lowest score (breast enlargement) was found in just 1.9% of observations without a clinical diagnosis for breast cancer, while the symptom with the highest score (acute cardiac failure) was found in 58% of observations without a clinical diagnosis for cardiomyopathy.

### Genome-first identification of new diagnoses in clinically undiagnosed individuals

We undertook a detailed case series analysis of clinically undiagnosed individuals carrying P/LP/LoF variants in plakophilin 2 (*PKP2*) and low-density lipoprotein receptor (*LDLR*) genes who had EHR evidence of symptomatic disease as examples for demonstrating how a genome-first evaluation can guide a targeted phenotypic evaluation and reveal new diagnoses.

### **Article**



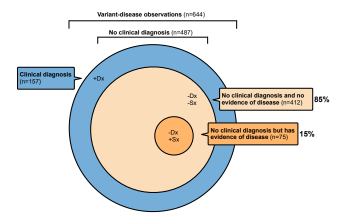


Figure 2. Assessment of the clinical diagnosis and phenotypes of 644 observations of 303 pathogenic/likely pathogenic or predicted LoF (P/LP/LoF) variants in 614 individuals

Observations were categorized into three distinct groups: (1) clinically diagnosed (+Dx), where the individual with a P/LP/LoF variant had a prior clinical diagnosis for the genetic disorder corresponding to the variant; (2) no clinical diagnosis and no evidence of disease (-Dx -Sx), where the individual with a P/LP/LoF variant lacked a prior clinical diagnosis for the corresponding genetic disorder and had no electronic health record (EHR) evidence of disease symptoms or findings; and (3) no clinical diagnosis but evidence of disease (-Dx +Sx), where the individual with a P/LP/LoF variant lacked a clinical diagnosis for the corresponding genetic disorder but had EHR evidence of disease symptoms or findings. The latter group was the primary outcome of interest for downstream analyses, comprising 75 of 487 (15%) clinically undiagnosed observations that had EHR evidence of symptomatic untreated disease.

Seven individuals harbored the P/LoF PKP2 W538Ter variant (NM\_004572.3:c.1613G>A), of whom five (71%) lacked a clinical diagnosis of cardiomyopathy (Figure 1). Of these five individuals, two (40%) were found to have EHR evidence of cardiomyopathy. Physician notes, including for electrocardiograms and echocardiography, for these two individuals were reviewed and summarized. Both individuals presented with multiple syncopal episodes, palpitations, and chest pain along with marked tachycardia and prolonged QRS duration on the electrocardiogram. These findings in two individuals with PKP2 W538Ter suggested a diagnosis of arrhythmogenic right ventricular cardiomyopathy according to the 2010 Task Force Criteria.<sup>24</sup>

Of 11 individuals with P/LP/LoF variants in LDLR, six (55%) were missing a corresponding clinical diagnosis, of whom three (50%) had EHR evidence of familial hypercholesterolemia and carried three different P/LP/LoF variants (NM\_000527. 4:c.772G>T, p.E258Ter; NM\_000527.4:c.1860G>T, p.W620C; and NM\_000527.4:c.590G>A, p.C197Y). Physician notes and lipid panel results for the three individuals were obtained and reviewed. All individuals presented on at least three occasions with chest pain, at least one occasion with claudication, and in one individual, one occasion with ischemic stroke due to an occluded carotid artery. Elevated total cholesterol and low-density lipoprotein cholesterol levels were noted in all individuals on multiple occasions, and one individual had a family history of myocardial infarction and coronary artery bypass grafting. These findings in three individuals with P/LP/LoF variants in LDLR indicated a diagnosis of familial hypercholesterolemia based on the Simon-Broome criteria.<sup>25</sup>

### **DISCUSSION**

Here, we performed a population-based genome-first evaluation of individuals carrying P/LP/LoF variants for nine genetic disorders in a large clinical care cohort, using a rich set of phenotype data to characterize individuals lacking a clinical diagnosis. The diagnostic utility of exome sequencing was assessed comprehensively for a wide array of diseases and revealed a preponderance of individuals with P/LP/LoF variants missing a relevant diagnosis in the EHR: 487 of 644 observations (76%) of variants in individuals lacked a corresponding clinical diagnosis. Individuals with evidence of symptomatic but clinically undiagnosed disease were identified by targeted phenotype assessment tailored to each person's genetic risk profile, such as a cardiovascular-specific evaluation for those with variants in cardiomyopathy-predisposition genes, demonstrating how precision medicine can be achieved with the integration of genomic and clinical data. We also interrogated key factors driving the detection of underdiagnosed disease. For example, not all clinically undiagnosed individuals had manifestations of disease, attributed in part to some P variants not associated with disease; 13 we found a greater prevalence of disease evidence in individuals with variants in genes observed to be penetrant in an independent cohort, emphasizing the importance of prioritizing genetic variation with demonstrable disease occurrence (e.g., penetrance) in the population rather than presumed pathogenicity.

Using a combination of phenotypes captured in the EHR, we observed that most individuals carrying P/LP/LoF variants lacked a relevant clinical diagnosis and, notably, some also had evidence of untreated symptomatic disease. This suggests a need to improve diagnostic approaches for individuals carrying P/LP/LoF variants. One possibility explored in the current study is to use a genome-first approach, screening for deleterious variants in well-known and empirically supported disease predisposition genes, the presence of which triggers a targeted disease evaluation in individuals with the variant. Such information gained from exome sequencing facilitated new genomeinformed diagnoses for several individuals who were missing a clinical diagnosis from routine care. For example, the finding of a P/LP/LoF PKP2 variant in two undiagnosed individuals informed a targeted evaluation of cardiac phenotypes in the EHR that supported a diagnosis of cardiomyopathy, while the identification of P/LP/LoF LDLR variants in three undiagnosed individuals guided an assessment of their cardiovascular and lipidic EHR profile that indicated a diagnosis of familial hypercholesterolemia. Guidelines for genetic testing of cardiomyopathy, familial hypercholesterolemia, and other inherited cardiovascular diseases hinge on a clinical diagnosis of the condition in the individual and/or the individual's family to trigger genetic testing<sup>10</sup>; similarly, genetic testing for hereditary cancers is recommended for individuals with a personal and/or family history of clinically diagnosed disease. 26-28 Instead, we implemented a genome-first testing<sup>29</sup> strategy to diagnose genetic disorders in the population that may otherwise have been missed with the standard of care.

Calls have been growing to use genomic sequencing to obtain a genetic diagnosis, <sup>9–11</sup> and nascent efforts have increased the diagnostic yield of several genetic disorders. <sup>14,15,30</sup> These have



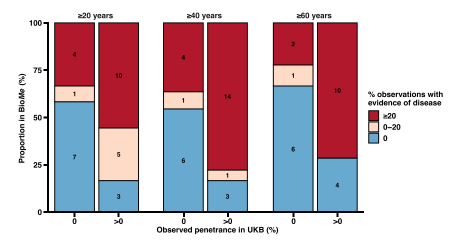


Figure 3. Rates of disease evidence found in clinically undiagnosed individuals with pathogenic/likely pathogenic or predicted LoF (P/LP/LoF) variants in BioMe Biobank (BioMe) for genes with versus without disease occurrence in UK Biobank (UKB)

A subset of 34 genes and six diseases that had clinically undiagnosed observations in BioMe were identified in UKB. Disease was observed in individuals from UKB with P/LP/LoF variants in 19 genes (penetrant genes) and not observed for 15 genes (non-penetrant genes) (STAR Methods). Yield of disease evidence in BioMe was then compared for P/LP/LoF variants found in penetrant genes versus non-penetrant genes. For each gene, clinically undiagnosed observations were evaluated for electronic health record evidence of symptomatic disease, and the proportion of observations with disease evidence was categorized as 0% (blue),

between 0% and 20% (tan), or  $\geq$ 20% (red). This analysis was completed three times using individuals who were  $\geq$ 20,  $\geq$ 40, and  $\geq$ 60 years of age. For example, for individuals  $\geq$ 40 years of age, 14 of 18 (78%) genes with disease observed in UKB had  $\geq$ 20% observations with disease evidence compared to 4 of 11 (36%) genes with no disease observed in UKB.

predominantly targeted three conditions designated tier 1 by the CDC.31 The present study adds to this literature a systematic evaluation of nine disorders beyond solely tier 1 conditions. Whereas previous studies have focused on small clinical cohorts of individuals with a family or personal history of disease, we used a large sample of unrelated individuals from a non-disease-ascertained cohort. Furthermore, we examined the clinical yield of variants in many ACMG SF v.3.1 genes; the ACMG has called for population screening studies of these clinically actionable genes, 5,6 and we found that 1 in 4 clinically undiagnosed observations of variants in ACMG SF v.3.1 genes had disease evidence in the population. We also accounted for variable disease risk of P/LP/LoF variants and the age of individuals. 13 It was previously unclear whether the lack of clinical diagnosis in genetically predisposed individuals is due to absence of disease, as in the case of incomplete penetrance and/or younger age, or presence of disease that is underdiagnosed. We assessed both of these factors; we evaluated the prevalence of disease evidence in clinically undiagnosed individuals who had variants in penetrant genes in an independent cohort from UK Biobank and in three age groups of individuals. Individuals with variants in penetrant genes had a greater prevalence of disease evidence than those with variants in non-penetrant genes, and age-stratified analyses showed consistent trends across the three age groups. Together, these results indicate that clinically undiagnosed individuals who have positive disease expression in the EHR are underdiagnosed, rather than simply lacking disease as a product of incompletely penetrant variants or younger age.

Clinical actionability of genetic findings is crucial to consider. Many genes in the study are in the CDC tier 1 genomic applications and the ACMG SF v.3.1, both of which comprise conditions for which early detection and intervention can reduce morbidity and mortality. Among the 75 individuals with clinical manifestations but no diagnosis, 57 (76%) had variants in ACMG SF v.3.1 genes, and 25 (33%) had variants in CDC tier 1 genes. This includes 3 individuals with LDLR variants who would be candidates for targeted lipid-lowering therapy and 22 individuals

with *BRCA1/BRCA2* variants who would be candidates for earlier mammography and breast cancer screening. Furthermore, a higher prevalence of clinical manifestations was observed among individuals carrying variants in penetrant genes; this suggests that penetrant genetic variation resulting in clinical manifestations that can be targeted by clinical interventions and treatment should be prioritized for genomic screening. Notably, most individuals carrying P/LP/LoF variants were of European ancestry, owing in part to the predominance of clinically interpreted variants identified in Europeans and underrepresentation of non-European ancestries in genetic databases, populations, and biobanks.<sup>32,33</sup> Further inclusion of diverse ancestries in clinical and population genetic studies will help identify and characterize more clinically relevant variants in non-European ancestries.

In conclusion, the findings of this population-based genome-first study demonstrate the untapped potential of using exome sequencing in healthcare systems to assess individuals carrying P/LP/LoF variants. We used exome data not yet part of routine clinical care in an EHR setting to inform a targeted strategy of phenotyping individuals at genetic risk for nine diseases. These data motivate the development and implementation of population-based genomic screening programs, which should be tested prospectively to diagnose individuals in the population who carry demonstrably deleterious genetic variation in well-known disease predisposition genes.

### **Limitations of the study**

There were several study limitations. First, disease symptoms were identified in part by International Classification of Diseases 10 (ICD-10) diagnosis codes. ICD-10 codes are commonly used to ascertain phenotypes in biobank studies;<sup>34,35</sup> however, we cannot exclude the possibility of some misclassification. Importantly, we supplemented ICD-10-based phenotypes with a thorough review of physician notes in the EHR (Table S6). This provided a check for accuracy of ICD-10 diagnosis codes and captured additional symptoms that may have otherwise been

### **Article**



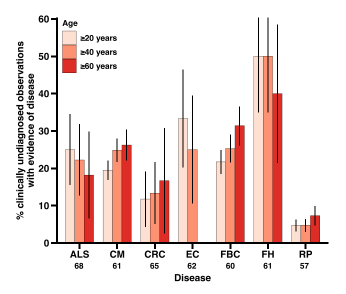


Figure 4. Evidence of symptomatic disease in clinically undiagnosed observations of variants in individuals from BioMe

The proportion of clinically undiagnosed observations that had electronic health record evidence of disease is depicted along with the 95% CIs (error bars) for three age groups and each of the seven disorders: amyotrophic lateral sclerosis (ALS), cardiomyopathy (CM), colorectal cancer (CRC), endometrial cancer (EC), familial breast cancer (FBC), familial hypercholesterolemia (FH), and retinitis pigmentosa (RP). The mean age of diagnosis is listed below each disease on the x axis. Monogenic diabetes (MD) and prostate cancer (PC) are not shown, as they both had zero observations lacking a clinical diagnosis (i.e., all individuals with MD or PC variants were diagnosed with the disease).

missed with an exclusively ICD-10-based phenotyping strategy. Nonetheless, not all phenotypes will be captured in the EHR, as individuals may have received care elsewhere, provider ICD-10 coding may not be standardized, and known biases exist when using ICD-10 diagnosis codes. Second, subgroups stratified by age had smaller sample sizes than in the primary analysis, particularly for the older age groups, which may account for more variability in the proportions of disease evidence in these subgroups. Furthermore, variants have heterogeneous effects on disease onset, including some instances where large-effect size variants are associated with earlier onset of disease; hence, age should be considered as a risk factor for disease. Third, a few of the disorders were associated with little to no detection of symptoms in the EHR, such as retinitis pigmentosa. It is possible that these individuals were seen by specialists in ophthalmology. Specialized tests and evaluations documented outside the standard physician note and diagnosis code system in the EHR would not have been detected in our analysis, possibly decreasing the number of clinically undiagnosed individuals with disease evidence. Another possibility is that some diseases and their manifestations are more evident to patients; symptoms associated with greater risk of hospitalization were more prevalent among observations of disease evidence. Fourth, while this study illuminated several important factors contributing to missing clinical diagnoses for genetic disorders, other reasons remain unknown. These include socioeconomic determinants, insurance status, access to healthcare systems, health literacy, environmental and lifestyle factors, demographics, and biological factors. Future studies are needed to further scrutinize the differences between individuals with genetic predisposition for disease who have and have not received a clinical diagnosis.

#### **STAR**\*METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- RESOURCE AVAILABILITY
  - Lead contact
  - Materials availability
  - Data and code availability
- EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS
  - O Study design
  - Study participants
- METHOD DETAILS
  - Variant sequencing and selection
  - Analysis of genes, age, diseases, and symptoms in clinically undiagnosed individuals
- QUANTIFICATION AND STATISTICAL ANALYSIS
  - Statistical analysis

### SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.xcrm.2024.101518.

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### **AUTHOR CONTRIBUTIONS**

I.S.F., A.D., and R.D. conceived and designed the study. All authors aided in the acquisition, analysis, and interpretation of data. I.S.F. and R.D. drafted the manuscript, and all authors provided critical revisions of the manuscript. I.S.F., A.D., J.K.P., and H.M.T.V. performed statistical analyses. G.N.N., J.H.C., and R.D. provided administrative, technical, and material support. R.D. supervised the study. I.S.F. and R.D. had access to and verified all of the data in the study.

### **DECLARATION OF INTERESTS**

R.D. reported receiving grants from AstraZeneca and grants and nonfinancial support from Goldfinch Bio and being a scientific co-founder, consultant, and equity holder for Pensieve Health (pending) and a consultant for Variant Bio, all not related to this work. G.N.N. reported being a scientific co-founder, consultant, advisory board member, and equity owner of Renalytix Al; a scientific



co-founder and equity holder for Pensieve Health (pending); and a consultant for Variant Bio and receiving grants from Goldfinch Bio and personal fees from Renalytix AI, BioVie, Reata, AstraZeneca, and GLG Consulting. L.R.P. is a consultant for Eyenovia, Twenty Twenty, and Skye Bioscience.

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### **STAR**\*METHODS

#### **KEY RESOURCES TABLE**

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Deposited data		
Summary table of clinical variants analyzed in Bio <i>Me</i> Biobank	Mendeley	Mendeley: https://doi.org/10.17632/c2g66gycvx.1
Penetrance of clinical variants in UK Biobank	Publication and Mendeley	https://doi.org/10.1001/jama.2021.23686 https://data.mendeley.com/datasets/v37dmjkbfj/ draft?a=a2f58e92-da19-461d-991a46dc70128860
ClinVar database	NCBI	https://www.ncbi.nlm.nih.gov/clinvar/
Institute for Personalized Medicine summary and access to BioMe Biobank	NCBI	https://www.ncbi.nlm.nih.gov/projects/gap/cgi-bin/study.cgi?study_id=phs000925.v1.p1
Human reference genome NCBI build 38, GRCh38	Genome Reference Consortium	http://www.ncbi.nlm.nih.gov/projects/genome/assembly/grc/human/
Software and algorithms		
R 3.5.3	R	https://cran.r-project.org/
PLINK 2.0	PLINK	https://www.cog-genomics.org/plink2/
Variant Effect Predictor	Github	https://github.com/Ensembl/ensembl-vep

#### **RESOURCE AVAILABILITY**

### Lead contact

Further information and requests for resources should be directed to and will be fulfilled by the lead contact, Ron Do (ron.do@ mssm.edu).

### **Materials availability**

This study did not generate new unique reagents.

### Data and code availability

- All data associated with this study are present in the paper or Supplementary Material and use existing data from a previous study. <sup>13</sup> A tabulated summary of all variants analyzed in the study is found in Table S3. This comprehensive list is annotated with information regarding genomic location, gene, associated disease, variant effect, amino acid change, ClinVar clinical significance and review status, minor allele frequencies in gnomAD, and minor allele frequencies in BioMe. In addition, this complete list has been deposited in Mendeley and its DOI is listed in the key resources table. Individual-level data, including sequencing, EHR phenotypes, and physician notes analyzed in this study are not publicly available due to Institutional Review Board (IRB) restrictions and research participant privacy concerns; however, requests from accredited researchers for access to data relevant to this manuscript can be made by contacting the lead contact.
- Any additional information required to reanalyze the data reported in this work paper is available from the lead contact upon request.
- No custom code was generated in this study.

### **EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS**

### Study design

A genome-first approach was applied to a population-based cohort whereby individuals carrying P/LP/LoF variants were identified and then phenotyped for the relevant genetic disorder. A schematic of the study's assessment of clinically undiagnosed individuals with P/LP/LoF variants is shown (Figure 1A). A diverse set of nine genetic disorders was analyzed: amyotrophic lateral sclerosis, cardiomyopathy, colorectal cancer, endometrial cancer, familial breast cancer, familial hypercholesterolemia, monogenic diabetes, prostate cancer, and retinitis pigmentosa. These were selected from a previous study<sup>13</sup> to ensure representation of a range of different systems (cardiac, neoplastic, metabolic, ocular, etc.) and prevalence (rare and common), and diseases that are analyzable in the EHR. Clinical diagnosis of disease was defined by the presence of a corresponding ICD-10 diagnosis code used in a previous study<sup>13</sup> (Table S4).

### **Article**



Individuals with P/LP/LoF variants were categorized phenotypically on the basis of their clinical diagnosis and disease symptoms extracted from the EHR. Evidence of symptomatic disease comprised ICD-10 codes and physician notes for symptoms captured by a set of curated clinical criteria (Table S6). This genome-first approach of first identifying individuals at high genetic risk for disease and then examining their ICD-10 codes, physician notes, and medications has been used previously. <sup>15,16,36</sup> This produced three distinct groups of individuals: clinically diagnosed (with a clinical diagnosis of the relevant disease), no clinical diagnosis and no evidence of disease (no clinical diagnosis and no EHR evidence of symptomatic disease), or no clinical diagnosis but has evidence of disease (EHR evidence of symptomatic disease but no clinical diagnosis). This latter group was the outcome of interest for most analyses unless otherwise stated. We reviewed the EHR pharmacy record and confirmed that all clinically undiagnosed individuals with evidence of disease were not treated with a disease-specific medication, defined as a medication with an indication specific to the disease of interest (Table S5). Medication indications were obtained from Drugbank.<sup>37</sup>

The study protocols were approved by the IRB at Mount Sinai (GCO#07–0529; STUDY-11-01139) and written informed consent was obtained for all participants. Use of data from the UK Biobank was approved under application number 16218 in the UK Biobank Resource. The study used de-identified genetic and EHR data for research purposes only.

### Study participants

A cohort of participants from an EHR-linked population-based biobank (BioMe) was included in the study. BioMe consists of over 60,000 individuals of African, Hispanic, European, Asian, and other self-reported ethnicities who were recruited from outpatient centers in the Mount Sinai Health System from 2007 onwards. All individuals in BioMe consented to providing biological and DNA samples linked to de-identified EHRs. Exome sequencing and quality control were performed for the first 31,250 participants. Samples with discordance between genetic sex and recorded sex, low coverage, contamination, low call rate, or duplications were excluded, leaving 30,813 samples. Those without complete demographic data (n = 345), younger than 20 years of age (n = 610), or without ICD-10 diagnosis data (n = 819) were excluded, leaving a set of 29,039 samples for analysis.

### **METHOD DETAILS**

### Variant sequencing and selection

Exome sequence data and variant call files (VCFs) were generated by the Regeneron Genetics Center (preparation and quality control described extensively elsewhere  $^{13}$ ). The average 20X coverage was 95% and greater than 99% of the samples had more than 85% of the targeted bases covered at 20X or more. Briefly, 9,202,884 variants were called in the samples and the Goldilocks Filter (GF) was applied to the VCFs. To single nucleotide polymorphisms (SNPs), cells with depth-normalized quality scores <3 or depth of coverage <7 were set to missing. For insertions and deletions (indels), cells with depth-normalized quality scores <5 or depth of coverage <10 were set to missing. Variant sites were then filtered, whereby sites of heterozygous variation that failed the Allele Balance (AB) cutoff were removed. SNP sites required  $\geq 1$  sample to carry an alternate AB  $\geq 15\%$  and indel sites required  $\geq 1$  sample to carry an alternate AB  $\geq 20\%$ . These site filters left 8,761,478 variants after GF. Next, sites with missing genotypes for >2% of individuals in the dataset (267,955 sites) were removed. AB was calculated for biallelic SNPs and 320,877 sites with AB <0.3 or >0.8 were removed, leaving 8,172,646 sites. Lastly, the dataset was filtered to regions within the target regions of the exome capture platform (IDT xGen capture platform; 4,256,827 sites) and separated into two file sets for biallelic and multiallelic sites (3,948,623 and 308,204, respectively). All variants were ascertained from VCFs using PLINK (version 2.0).

Variants in the analysis were derived from a set of P/LP/LoF exonic variants (LoF variants in genes mediating disease via LoF mechanism) without recessive inheritance (excluded due to insufficient sample size) that were previously characterized in BioMe. <sup>13</sup> The previous study assessed population-based penetrance of a wide array of variants, while the present study investigated clinical underdiagnosis of individuals carrying rare P/LP/LoF variants, factors associated with underdiagnosis, and optimization of genome-first approaches for clinical utility. P/LP variants reported in ClinVar and previously unreported variants with a LoF molecular consequence (splice acceptor/donor, stop gained/lost, frameshift, or start lost) annotated by Variant Effect Predictor <sup>40</sup> were identified. Variants in the last exon or last 50 base pairs of the penultimate exon were considered not LoF due to a predicted lack of efficient nonsense-mediated decay, with exception for variants predicted to delete over 20% of the gene. Only LoF variants present in MANE Select or MANE Plus Clinical transcripts were retained. <sup>41</sup> LoF variants in a gene were then mapped to disease based on prior P/LP variant submissions in ClinVar linking genes to diseases (e.g., *BRCA1* LoF variants mapped to breast cancer). Variants with benign, uncertain, or conflicting clinical significance in ClinVar and predicted synonymous consequence were removed, as were variants in genes with recessive inheritance reported in Online Mendelian Inheritance in Man. <sup>42</sup> ClinVar variants were further restricted to those reported by multiple clinical testing labs or reviewed by an expert panel such as ClinGen. <sup>8</sup> Rare LoF variants with ancestry-specific allele frequency <0.001 in BioMe and ancestry-specific allele frequency <0.001 in or absent from gnomAD v3.1.2<sup>43</sup> were included. This yielded 303 P/LP/LoF variants in 54 disease-predisposition genes corresponding to nine genetic disorders (Table S1).

### Analysis of genes, age, diseases, and symptoms in clinically undiagnosed individuals

Importantly, the evaluation of disease in clinically undiagnosed individuals with P/LP/LoF variants accounted for different genes, ages, diseases, and symptoms. First, we hypothesized that a genome-first approach using variants in genes for which disease is observed in real-world non-disease ascertained populations would result in higher diagnostic yield. Although all variants in the target



genes had evidence of pathogenicity, expected pathogenicity does not always translate to occurrence of disease in the population. <sup>13</sup> Population genomic screening would ideally use variants in genes with disease observed in real-world populations to increase clinical yield. We tested this hypothesis by first identifying 19 target genes for which disease was observed in individuals with any P/LP/LoF variants in the gene in an independent cohort from UK Biobank<sup>13</sup> (penetrant) and 15 target genes for which disease was not observed (non-penetrant) in UK Biobank, and then comparing the proportion of clinically undiagnosed observations with disease evidence in BioMe for penetrant genes versus non-penetrant genes (Figure 1B). The exome sequence and EHR data used to characterize disease-associated genes from the UK Biobank have been previously described. <sup>13</sup> The proportion of observations of individuals with variants who had symptomatic evidence of disease in BioMe was then compared for variants in the penetrant genes versus non-penetrant genes.

Second, the presence or absence of disease evidence may be explained by age; for example, younger individuals may not have manifested the disease yet. To address this, we separately evaluated the proportion of clinically undiagnosed observations that had evidence of disease in three age groups of individuals  $\geq 20$  years,  $\geq 40$  years, and  $\geq 60$  years.

Third, genetic disorders are heterogeneous in terms of affected systems and clinical presentation, such that certain diseases and symptoms may be better detected in the healthcare setting. We therefore examined the proportion of clinically undiagnosed observations of variants in individuals with evidence of disease for each genetic disorder and the prevalence of each symptom detected. We tested the hypothesis that symptoms associated with hospitalization would be better detected and have a higher prevalence in our analysis using a previously published score<sup>23</sup> on an ordinal scale ranging from 0 (smallest hospitalization and mortality risk) to 1 (greatest hospitalization and mortality risk) assigned to each of the 39 disease symptoms (Table S8).

#### **QUANTIFICATION AND STATISTICAL ANALYSIS**

### Statistical analysis

Differences in categorical and continuous variables were assessed with a two-sided unpaired Fisher's exact test and t-test, respectively. Individuals with at least one allele for a P/LP/LoF variant were identified and the proportions of observations of variants in individuals with 1) a clinical diagnosis; 2) no clinical diagnosis or evidence of disease; and 3) no clinical diagnosis but have evidence of disease were determined. Analyses of disease in clinically undiagnosed observations were stratified by gene, age groups ( $\geq$ 20 years,  $\geq$ 40 years,  $\geq$ 60 years), and disease. Multivariable linear regression adjusted for age was used to model the proportion of clinically undiagnosed observations with evidence of disease as a function of gene category (penetrant gene versus gene without penetrant in UK Biobank) and to model the prevalence of symptoms in clinically undiagnosed observations as a function of symptom score. All statistical tests and plots were generated with R (version 3.5.3).

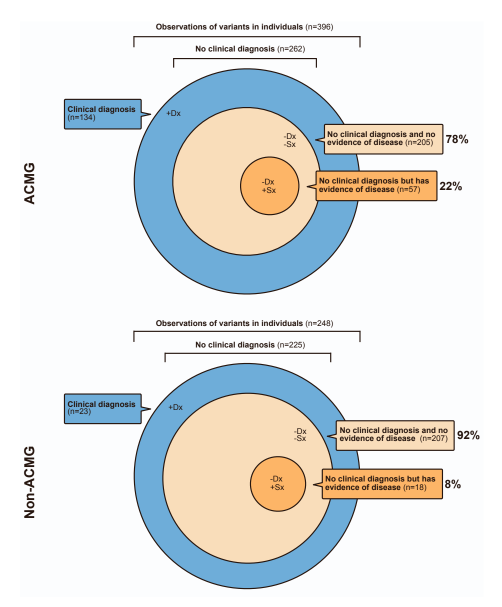
Cell Reports Medicine, Volume 5

### **Supplemental information**

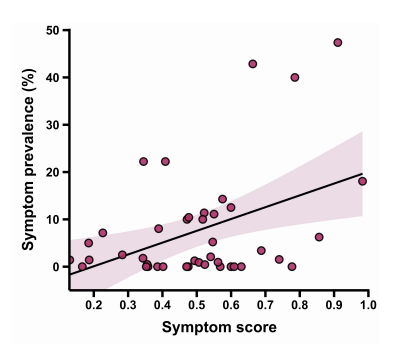
Genome-first evaluation with exome sequence and clinical data uncovers underdiagnosed genetic disorders in a large healthcare system

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### SUPPLEMENTARY FIGURES



Supplementary Figure 1. Clinical diagnosis and phenotypes of observations in individuals of pathogenic/likely pathogenic or predicted loss-of-function (P/LP/LoF) variants in genes from the American College of Medical Genetics and Genomics' Recommendations for Reporting of Secondary Findings (ACMG SF v3.1). Observations of variants in individuals were categorized into three distinct groups: 1) clinically diagnosed (+Dx), the individual with a P/LP/LoF variant had a prior clinical diagnosis for the genetic disorder corresponding to the variant; 2) no clinical diagnosis and no evidence of disease (-Dx -Sx), the individual with a P/LP/LoF variant lacked a prior clinical diagnosis for the corresponding genetic disorder and had no electronic health record (EHR) evidence of disease symptoms or findings detected; and 3) no clinical diagnosis but has evidence of disease (-Dx +Sx), the individual with a P/LP/LoF variant lacked a clinical diagnosis for the corresponding genetic disorder, but has EHR evidence of disease symptoms or findings. Related to Figure 2.



Supplementary Figure 2. Prevalence of symptoms detected in individuals without a clinical diagnosis carrying pathogenic/likely pathogenic or loss-of-function (P/LP/LoF) variants. P/LP variants, reported in ClinVar with a minimum review status of two; LoF variants, variants with predicted LoF consequence annotated by Variant Effect Predictor; clinical diagnosis, clinical diagnosis of disease corresponding to a variant; symptom prevalence, prevalence of symptom detected in observations of variants in individuals without a clinical diagnosis; symptom score, on an ordinal scale ranging from 0 (lowest risk of hospitalization and death) to 1 (highest risk of hospitalization and death) from a previously published study<sup>1</sup>. Related to Figure 4.

1. Gottlieb A, Hoehndorf R, Dumontier M, Altman RB. Ranking adverse drug reactions with crowdsourcing. *J Med Internet Res* 2015; **17**. DOI:10.2196/jmir.3962.

### **SUPPLEMENTARY TABLES**

Supplementary Table 1. Summary of 54 monogenic disease predisposition genes containing pathogenic/likely pathogenic or predicted loss-of-function (P/LP/LoF) variants including a subset in the American College of Medical Genetics and Genomics' Recommendations for Reporting of Secondary Findings list of 78 genes (ACMG SF v3.1). Related to Table 2.

Disease	Gene	ACMG SF v3.1
Amyotrophic lateral sclerosis	FIG4	FALSE
Amyotrophic lateral sclerosis	FUS	FALSE
Amyotrophic lateral sclerosis	OPTN	FALSE
Amyotrophic lateral sclerosis	SETX	FALSE
Cardiomyopathy	ABCC9	FALSE
Cardiomyopathy	ACTN2	FALSE
Cardiomyopathy	CSRP3	FALSE
Cardiomyopathy	DES	TRUE
Cardiomyopathy	DSG2	TRUE
Cardiomyopathy	DSP	TRUE
Cardiomyopathy	EYA4	FALSE
Cardiomyopathy	MYBPC3	TRUE
Cardiomyopathy	МҮН6	FALSE
Cardiomyopathy	МҮН7	TRUE
Cardiomyopathy	NEXN	FALSE
Cardiomyopathy	PKP2	TRUE
Cardiomyopathy	PRKAG2	TRUE
Cardiomyopathy	RBM20	TRUE
Cardiomyopathy	RYR2	TRUE
Cardiomyopathy	SCN5A	TRUE
Cardiomyopathy	SDHA	FALSE
Cardiomyopathy	TNNI3	TRUE
Cardiomyopathy	TNNT2	TRUE
Cardiomyopathy	TTN	TRUE
Cardiomyopathy	VCL	FALSE
Colorectal cancer	APC	TRUE
Colorectal cancer	BUB1B	FALSE
Colorectal cancer	FLCN	FALSE
Colorectal cancer	MLH1	TRUE
Colorectal cancer	MLH3	FALSE
Colorectal cancer	MSH2	TRUE
Colorectal cancer	TGFBR2	FALSE
Endometrial cancer	MSH6	TRUE
Familial breast cancer	BRCA1	TRUE
Familial breast cancer	BRCA2	TRUE
Familial breast cancer	PALB2	TRUE
Familial breast cancer	PPM1D	FALSE
Familial hypercholesterolemia	LDLR	TRUE

Monogenic diabetes	KCNJ11	FALSE
Prostate cancer	BRCA2	FALSE
Retinitis pigmentosa	AIPL1	FALSE
Retinitis pigmentosa	CRB1	FALSE
Retinitis pigmentosa	DHDDS	FALSE
Retinitis pigmentosa	KLHL7	FALSE
Retinitis pigmentosa	NR2E3	FALSE
Retinitis pigmentosa	NRL	FALSE
Retinitis pigmentosa	PDE6B	FALSE
Retinitis pigmentosa	PROM1	FALSE
Retinitis pigmentosa	PRPF8	FALSE
Retinitis pigmentosa	PRPH2	FALSE
Retinitis pigmentosa	RHO	FALSE
Retinitis pigmentosa	RLBP1	FALSE
Retinitis pigmentosa	RP1	FALSE
Retinitis pigmentosa	RPE65	TRUE
Retinitis pigmentosa	SEMA4A	FALSE

### Supplementary Table 2. Summary of baseline demographic traits by genetic disorder. Related to Figure 4 and Table 1.

Disease	Age, mean (SD) years	Female, n (%)	European, n (%)	African, n (%)	Hispanic, n (%)	Asian, n (%)	Other, n (%)
Alzheimer disease	75 (13)	464 (63)	161 (22)	185 (25)	317 (43)	11 (1.5)	65 (8.8)
Amyotrophic lateral sclerosis	72 (12)	5 (71)	4 (57)	0 (0)	3 (43)	0 (0)	0 (0)
Breast cancer	64 (14)	1488 (100)	580 (39)	335 (23)	409 (27)	43 (2.9)	121 (8.1)
Cardiomyopathy	67 (14)	344 (42)	226 (27)	248 (30)	250 (30)	21 (2.6)	78 (9.5)
Colorectal cancer	72 (11)	122 (56)	59 (27)	58 (27)	78 (36)	3 (1.4)	20 (9.2)
Endometrial cancer	69 (11)	115 (100)	25 (22)	30 (26)	51 (44)	2 (1.7)	7 (6.1)
Hypercholesterolemia	67 (13)	2777 (62)	1437 (32)	1114 (25)	1391 (31)	158 (3.5)	360 (8.1)
Prostate cancer	75 (9.7)	0 (100)	177 (37)	154 (32)	105 (22)	6 (1.3)	32 (6.8)
Retinitis pigmentosa	63 (15)	7 (47)	4 (27)	6 (40)	5 (33)	0 (0)	0 (0)
Type 2 diabetes	67 (13)	4113 (58)	1087 (15)	2208 (31)	2850 (40)	250 (3.5)	649 (9.2)

Clinical diagnoses were extracted as International Classification of Diseases 10 (ICD-10) codes from the EHR and ancestries were self-reported. Other ancestry, miscellaneous ancestries other than those listed; n, number; SD, standard deviation.

Supplementary Table 3. Study set of 303 pathogenic/likely pathogenic or predicted loss-of-function (P/LP/LoF) variants in 54 genes corresponding to nine genetic conditions. Related to Figures 1-4 and Tables 1-2.

Dataset is available at: <a href="http://dx.doi.org/10.17632/c2g66gycvx.1">http://dx.doi.org/10.17632/c2g66gycvx.1</a>.

# **Supplementary Table 4. ICD-10 diagnosis codes for nine genetic disorders.** Related to Figures 1, 2, and 4, and Tables 1 and 2.

Disease	ICD-10 diagnosis code
Amyotrophic lateral sclerosis	Amyotrophic lateral sclerosis ('G12.21')
Cardiomyopathy	Cardiomyopathies, including dilated and hypertrophic types ('I42*')
Colorectal cancer	Malignant neoplasm of the colon and/or rectum ('C18*', 'C19')
Endometrial cancer	Malignant neoplasm of the endometrium ('C54.1')
Breast cancer	Malignant neoplasm of the breast ('C50*')
Familial hypercholesterolemia	Hypercholesterolemia, unspecified or familial ('E78.0*')
Prostate cancer	Malignant neoplasm of the prostate, including personal history of prostate cancer and in situ carcinoma of the prostate ('C61', 'Z85.46', 'D07.5')
Retinitis pigmentosa	Retinitis pigmentosa ('H35.52')
Monogenic diabetes	Type 2 diabetes ('E11*')

<sup>\*,</sup> any ICD-10 diagnosis code below the stated parent level (e.g., 'F90\*' includes 'F90.1').

# Supplementary Table 5. Disease-specific medications for nine genetic disorders with prescriptions identified in the electronic health record. Related to Figures 1 and 4, and Table 2.

Disease	Medications
Amyotrophic lateral sclerosis	riluzole
Cardiomyopathy	ivabradine, dexrazoxane
Colorectal cancer	oxaliplatin, panitumumab, tipiracil
Endometrial cancer	
Familial breast cancer	abemaciclib, adriamycin, anastrozole, arimidex, aromasin, capecitabine, carboplatin, cisplatin, cyclophosphamide, docetaxel, doxorubicin, exemestane, femara, femring, fl uorouracil, gemcitabine, goserelin, herceptin, ibandronate, ibrance, ixabepilone, lapat inib, letrozole, lynparza, megace, methotrexate, methyltestosterone, mitomycin, navel bine, neratinib, olaparib, otrexup, paclitaxel, palbociclib, pertuzumab, raloxifene, ras uvo, tamoxifen, taxotere, toremifene, trastuzumab, tucatinib, verzenio, vinblastine, vi norelbine, xeloda, zoladex
Familial hypercholesterolemia	atorvastatin, simvastatin, pravastatin, lovastatin, fluvastatin, pitavastatin, rosuvastatin, cholestyramine, colestipol, colesevelam, niacin, ezetimibe, alirocumab, evolocumab
Monogenic diabetes	acarbose, actos, admelog, albiglutide, alogliptin, amaryl, avandia, canagliflozin, chlor propamide, cycloset, dapagliflozin, dulaglutide, empagliflozin, ertugliflozin, exenatid e, glimepiride, glinide, glipizide, glucophage, glucotrol, glumetza, glyburide, glyset, humalog, insulin, insulin degludec, insulin detemir, insulin glargine, insulin lispro, in vokana, januvia, jardiance, junior kwikpen, levemir, linagliptin, liraglutide, lixisenati de, metformin, miglitol, nateglinide, onglyza, ozempic, pioglitazone, pramlintide, pre cose, repaglinide, riomet, rosiglitazone, rybelsus, saxagliptin, saxenda, semaglutide, s itagliptin, starlix, steglatro, symlinpen, tanzeum, tradjenta, tresiba, trulicity, victoza
Prostate cancer	degarelix, triptorelin, bicalutamide, flutamide, abiraterone, enalautamide, nilutamide, apalutamide, darolutamide, sipuleucel
Retinitis pigmentosa	

Disease-specific medication, medication with an indication specific to the disease (e.g., anastrozole is specifically indicated for breast cancer, whereas cyclophosphamide's indication is not specific to breast cancer); --, no disease-specific medication.

Supplementary Table 7. Clinical diagnosis and phenotype of individuals with pathogenic/likely pathogenic or predicted loss-of-function (P/LP/LoF) variants stratified by ancestry. Related to Figure 2, and Tables 1 and 2.

	01 41	· D (0/)	D (0/)	D C (0/)	D (0/)
Ancestry	Observations, n	+Dx, n (%)	-Dx, n (%)	-Dx -Sx, n (%)	-Dx +Sx, n (%)
African	121	30 (25)	89 (74)	72 (81)	17 (19)
Asian	29	9 (31)	20 (69)	17 (85)	3 (15)
European	259	67 (26)	191 (74)	164 (86)	27 (14)
Hispanic	182	39 (21)	145 (80)	123 (85)	22 (15)
Other	53	12 (23)	42 (79)	36 (86)	6 (14)

Assessment of the clinical diagnosis and phenotypes of 644 observations of 303 pathogenic/likely pathogenic or predicted loss-of-function (P/LP/LoF) variants in 614 individuals stratified by self-reported ancestry. Observations were categorized into three distinct groups: 1) clinically diagnosed (+Dx), the individual with a P/LP/LoF variant had a prior clinical diagnosis for the genetic disorder corresponding to the variant; 2) no clinical diagnosis and no evidence of disease (-Dx -Sx), the individual with a P/LP/LoF variant lacked a prior clinical diagnosis for the corresponding genetic disorder and had no electronic health record (EHR) evidence of disease symptoms or findings; and 3) no clinical diagnosis but has evidence of disease (-Dx +Sx), the individual with a P/LP/LoF variant lacked a clinical diagnosis for the corresponding genetic disorder, but had EHR evidence of disease symptoms or findings.

Supplementary Table 8. Symptom scores for 39 symptoms and their prevalence among clinically undiagnosed observations of variants and disease in individuals with pathogenic/likely pathogenic or predicted loss-of-function (P/LP/LoF) variants. Related to Figure 4 and Table 2

Disease	Sign/symptom	Matched phenotype	Symptom	No clinical	Evidence of	
		from Gotlieb et al.1	score	diagnosis, n	disease, n (% of	
					no clinical diagnosis)	
Prostate cancer	Metastatic disease	Bone cancer metastatic	0.9829212	0		
Cardiomyopathy	Heart failure	Cardiac failure acute	0.9108916	154	90 (58)	
Amyotrophic					(00)	
lateral sclerosis	Respiratory failure	Respiratory failure	0.856541	20	4 (20)	
Familial		•			, ,	
hypercholesterole						
mia	Coronary heart disease	Coronary artery disease	0.7857806	6	3 (50)	
Cardiomyopathy	Valvular dysfunction	Cardiac valve disease	0.7401669	154	23 (15)	
Cardiomyopathy	Cardiac remodeling	Cardiac hypertrophy	0.6879898	154	16 (10)	
Endometrial	Abnormal uterine bleedin					
cancer	g	Uterine haemorrhage	0.6631796	12	2 (17)	
Familial						
hypercholesterole		Peripheral vascular diso				
mia	Peripheral artery disease	rder	0.6298026	6	0 (0)	
Prostate cancer	Anomalous DRE findings	Prostatic disorder	0.6086371	0		
	Endometrial hyperplasia a					
Endometrial	nd disorganization on bio	Endometrial hyperplasi				
cancer	psy	a	0.5750928	12	1 (8.3)	
Endometrial	Thickened endometrium o	Endometrial hypertroph				
cancer	n imaging	у	0.5683344	12	0 (0)	
Familial breast	Axillary lymphadenopath					
cancer	у	Lymphadenopathy	0.5619201	106	20 (19)	
Prostate cancer	Urethral compression	Urethral obstruction	0.5502927	0		
Familial breast						
cancer	Mass	Breast mass	0.5465219	106	6 (5.7)	
		Prostatic specific antige				
Prostate cancer	Elevated PSA	n increased	0.5405282	0		
Colorectal cancer	Exophytic colonic mass	Colonic polyp	0.5235142	17	1 (5.9)	
	Abnormal cardiac exam fi					
Cardiomyopathy	ndings	Heart sounds abnormal	0.5215908	154	17 (11)	
Familial						
hypercholesterole						
mia	Hypercholesterolemia	Hypercholesterolaemia	0.5175085		1 (17)	
Colorectal cancer	Obstructed lumen	Colonic obstruction	0.5063074	17	0 (0)	
	Bulbar symptoms (dysarth					
Amyotrophic	ria, dysphagia, facial wea					
lateral sclerosis	kness, tongue atrophy)	Facial paresis	0.493729	20	0 (0)	
	Infiltration to genitourinar					
Prostate cancer	y system	Haematuria	0.4768623	0		
Monogenic						
diabetes	Insulin perturbations	Insulin resistance	0.4752398	0		
Familial						
hypercholesterole						
mia	Corneal arcus	Corneal deposits	0.4716786	6	1 (17)	

Endometrial	Enlarged uterus/abdomina				
cancer	l mass	Uterine enlargement	0.470359	12	0 (0)
Monogenic	Dysregulated glycemic co	Blood glucose increase			
diabetes	ntrol	d	0.4085338	0	
Familial					
hypercholesterole					
mia	Xanthoma	Xanthoma	0.4019195	6	0 (0)
		Faecal occult blood posi			
Colorectal cancer	Occult bleeding or melena	tive	0.3889438	17	3 (18)
Amyotrophic					
lateral sclerosis	Labile affect	Emotional disorder	0.3856494	20	0 (0)
Monogenic					
diabetes	Acanthosis nigricans	Acanthosis nigricans	0.358083	0	
Familial breast		Breast microcalcificatio			
cancer	Imaging anomalies	n	0.3556099	106	5 (4.7)
Retinitis	Nyctalopia and glare sensi				
pigmentosa	tivity	Night blindness	0.3528767	168	0 (0)
Monogenic	Polyuria, polydipsia, poly				
diabetes	phagia	Polyuria	0.3449517	0	
Retinitis	Progressive, peripheral vis				
pigmentosa	ion loss	Tunnel vision	0.3432615	168	3 (1.8)
Amyotrophic	Asymmetric limb weakne				
lateral sclerosis	SS	Muscular weakness	0.2832084	20	1 (5)
Endometrial					
cancer	Pelvic pain	Pelvic pain	0.2263445	12	1 (8.3)
Familial breast					
cancer	Dermatological changes	Breast tenderness	0.1866282	106	0 (0)
Amyotrophic					
lateral sclerosis	Muscle fasciculations	Muscle twitching	0.185048	20	0 (0)
Familial breast					
cancer	Nipple abnormalities	Breast discharge	0.1665234	106	0 (0)
Familial breast					
cancer	Histopathological findings	Breast enlargement	0.1300308	106	2 (1.9)

No clinical diagnosis, observations of variants in individuals lacking a corresponding clinical diagnosis; evidence of disease, clinically undiagnosed observations that had electronic health record evidence of symptomatic disease; symptom score, on an ordinal scale ranging from 0 (lowest risk of hospitalization or death) to 1 (highest risk of hospitalization or death) from a previously published study<sup>1</sup>; --, not applicable as all observations for monogenic diabetes and prostate cancer were diagnosed with the corresponding disease.

1. Gottlieb A, Hoehndorf R, Dumontier M, Altman RB. Ranking adverse drug reactions with crowdsourcing. *J Med Internet Res* 2015; **17**. DOI:10.2196/jmir.3962.