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1	Overlapping Genetic Architecture between Parkinson Disease and Melanoma
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22 Abstract

Epidemiologic studies have reported inconsistent results regarding an association between 23 24 Parkinson disease (PD) and cutaneous melanoma (melanoma). Identifying shared genetic architecture between these diseases can support epidemiologic findings and identify common 25 risk genes and biological pathways. Here we apply polygenic, linkage disequilibrium-informed 26 27 methods to the largest available case-control, genome-wide association study summary statistic 28 data for melanoma and PD. We identify positive and significant genetic correlation (correlation: 0.17, 95% CI 0.10 to 0.24; $P = 4.09 \times 10^{-06}$) between melanoma and PD. We further 29 demonstrate melanoma and PD-inferred gene expression to overlap across tissues (correlation: 30 0.14, 95% CI 0.06 to 0.22; $P = 7.87 \times 10^{-04}$), and highlight seven genes including *PIEZO1*, 31 32 TRAPPC2L, and SOX6 as potential mediators of the genetic correlation between melanoma and PD. These findings demonstrate specific, shared genetic architecture between PD and 33 melanoma that manifests at the level of gene expression. 34 **Keywords:** Parkinson disease; melanoma; genetic correlation; polygenic; TWAS; shared 35

36 genetic architecture

38 Introduction

39 An association between idiopathic Parkinson disease (PD), neuropathologically characterized by the degeneration of pigmented dopaminergic neurons, and cutaneous 40 melanoma (melanoma), a cancer of pigment-producing melanocytes, was first reported in 1972 41 [80]. This association was hypothesized to result from the chronic systemic administration of 42 43 levodopa (L-DOPA) – an intermediate in the dopamine synthesis pathway [23] – for the treatment of PD [4, 80] as L-DOPA is also a biosynthetic intermediate in the production of 44 45 melanin [23]. Since that time, several epidemiologic studies have examined the association between PD and melanoma as well as other cancers [5, 17, 21, 27, 29, 36, 42, 53, 67, 68, 81, 46 87, 91]. The majority of studies have found that individuals with PD appear to have a lower 47 incidence of most cancers, with the exception of melanoma [21, 27, 36, 67, 68, 81, 91]. Both 48 49 prospective and retrospective studies have also found an increased risk of melanoma in PD that appears to be independent of L-DOPA treatment [5, 29, 42, 67, 91]. For example, 92 out of 50 51 2,106 (4.4%) individuals with neurologist-confirmed PD had either a personal history or current dermatologist-diagnosed melanoma in a 2010 study [5]. The increased risk of melanoma in PD 52 has been observed to extend to family members and be reciprocal in nature with individuals 53 54 being at greater risk for PD if their relatives have a melanoma diagnosis and vice versa [29, 42]. 55 For example, 40 of 1,544 (2.6%) of individuals with pathologically-confirmed melanoma had a 56 neurologist-confirmed diagnosis of PD in a 2017 study [17]. However, not all studies have identified an association between melanoma and PD in affected individuals [19, 27] or their 57 relatives [91]. An epidemiologic association between lighter hair color and PD, a potentially 58 59 shared risk factor with melanoma [6], has also been inconsistently reported [19, 30]. Epidemiologic association studies are not without biases. PD is known to have an extended 60 prodromal period and a melanoma diagnosis necessitates longitudinal follow up, both of which 61 increase medical surveillance and thus the chance for spurious epidemiologic associations [27, 62

33]. In contrast, studies of genetic variants associated with disease or cross-disease risk are not
expected to be influenced by usage of medical care, though they may be subject to similar
misclassification [75] and ascertainment biases.

The first investigations of a genetic relationship between melanoma and PD focused on 66 67 variants in *MC1R*, a gene strongly associated with pigmentation and melanoma risk [45]. While early reports identified an association between PD and MC1R variants [30, 83] other studies 68 failed to replicate these findings [24, 26, 28, 55]. Analyses focused on single variants in other 69 70 melanoma risk genes have also failed to yield consistent associations with PD [19, 28, 56]. 71 Multi-variant analyses have thus far reported a lack of genetic association as well. For example, 72 a melanoma genetic risk score – calculated by aggregating the effect of melanoma genome-73 wide association study (GWAS)-significant ($P < 5 \times 10^{-8}$) loci included in the GWAS catalog [89] 74 as of 2012 – was not significantly associated with PD [65]. Similarly, no evidence for an association between GWAS-significant melanoma loci and PD is observed in a more recent 75 76 multi-variant, Mendelian randomization study [66]. In contrast, genes associated with Mendelian forms of PD have been identified to be somatically mutated in melanoma lesions [37, 40, 48]. 77 There may also exist an enrichment of Mendelian PD gene germline variants in individuals with 78 79 melanoma [37], though this requires replication. Nevertheless, over 90% of individuals with PD 80 do not have mutations in any known Mendelian PD genes [1] and thus variants in Mendelian PD 81 genes are unlikely to fully explain any genetic correlation between melanoma and PD.

The genetic risk architecture underlying complex diseases like PD and melanoma is mediated by many common genetic variants of small effect size, most of which do not demonstrate GWAS-significant associations given current study sample sizes [8]. Analyses which only include GWAS-significant loci are not expected to fully represent the genetic architecture of these complex diseases and thus may lead to false negative genetic overlap results. Recently, statistical methods that aggregate all loci from disease-specific GWAS

summary statistic datasets in a linkage disequilibrium (LD)-informed manner have been
developed to better model these polygenic architectures [11]. These aggregated signals can be
leveraged to estimate the genetic correlation between different diseases [11, 54], even at the
level of gene expression in specific tissues [35, 57] or across tissues [38]. Here, we apply these
novel methods to GWAS summary statistics derived from the largest currently available studies
of melanoma [45], PD [13, 63, 64], and other neurodegenerative diseases [25, 44] to investigate
whether there exists specific genetic architecture overlap between melanoma and PD.

95

96 Methods

97 **GWAS Summary Statistics**:

We obtained the largest available, European genetic ancestry, case-control, GWAS 98 99 summary statistic data for melanoma (Law2015 [45]) and three independent studies of PD 100 (Nalls2014 [64]; Chang2017 [13]; Nalls2019 [63]) as well as two negative control comparator neurodegenerative diseases: Alzheimer disease (Kunkle2019 [44]) and frontotemporal dementia 101 (Ferrari2014 [25]). The summary statistics for these datasets included p-value, effect allele, 102 103 number of individuals or studies, and standard error for every genetic variant reported in each study. All individual studies contributing to the GWAS summary statistic datasets used in the 104 105 current analysis received approval from the pertinent institutional review boards or ethics committees, and all participants gave informed consent. Additional details for each dataset are 106 107 included below and in the individual study articles [13, 25, 44, 45, 63, 64].

108

109 Melanoma – Law2015

We obtained meta-analysis Melanoma risk summary statistic data from the Melanoma
meta-analysis consortium (<u>https://genomel.org/</u>). This data was published in Law et al., Nature
Genetics, 2015 [45]. This dataset includes melanoma-association results for 9,469,417

genotyped and imputed variants derived from 12,814 pathologically-confirmed melanoma casesand 23,203 controls of European ancestry.

115

116 Parkinson disease – Nalls2014

117 We obtained PD risk summary statistic data from PDGENE (http://www.pdgene.org/). This dataset was published in Nalls et al., Nature Genetics, 2014 [64] and Lill et al, PLoS 118 119 Genetics 2012 [50]. The summary statistic data we obtained did not include any 23andMe participants and thus the dataset includes PD-association results for 7,799,580 genotyped and 120 imputed variants derived from 9,581 PD cases - mostly diagnosed, but some self-reported -121 and 33,245 controls of European ancestry. This dataset only included the number of studies, 122 and not the number of individuals, supporting the association results for each variant. 123 124 Consequently, we only included variants supported by at least 12 of 13 studies in downstream 125 analyses. 126 Parkinson disease – Chang2017 127 We obtained Parkinson disease (PD) risk summary statistic data from 23andMe, Inc., a 128 129 personal genetics company (https://research.23andme.com/dataset-access/). This data was published in Chang et al., Nature Genetics, 2017 [13]. This dataset includes PD-association 130 results for 12,896,220 genotyped and imputed variants derived from 6,476 self-reported PD 131 cases and 302,042 controls of European ancestry. This dataset excludes any 23andMe 132 133 participants included in the Nalls2014 study. 134 Parkinson disease – Nalls2019 135 136 We obtained PD risk summary statistic data from the IPDGC (https://pdgenetics.org/). 137 This dataset was published in Nalls et al., The Lancet Neurology, 2019 [63]. The summary

138 statistic data we obtained did not include any 23andMe data nor Nalls2014 data and thus

- includes PD-association results for 17,510,617 genotyped and imputed variants derived from
 33,674 PD cases diagnosed and UKB proxy-cases, that is individuals with a first-degree
 relative with PD and 449,056 controls of European ancestry.
- 142

143 Alzheimer disease – Kunkle2019

We downloaded stage 1 meta-analysis Alzheimer Disease (AD) risk GWAS summary statistic data from NIAGADS (National Institute on Aging Genetics of Alzheimer Disease Data Storage Site) website: <u>https://www.niagads.org/datasets/ng00075</u> (#NG00075). This data was generated by the International Genomics of Alzheimer Project and published in Kunkle et al., Nature Genetics, 2019 [44]. The stage 1 meta-analysis dataset includes AD-association results for 11,480,632 genotyped and imputed variants derived from 21,982 AD cases and 41,944 cognitively normal controls of European ancestry.

151

152 Frontotemporal Dementia – Ferrari2014

We obtained discovery phase Frontotemporal Dementia (FTD) risk GWAS summary statistic data from the International Frontotemporal Dementia Genomics Consortium (IFGC, <u>https://ifgcsite.wordpress.com/data-access/</u>). This data was generated by the IFGC and published in Ferrari et al., Lancet Neurology, 2014 [25]. The discovery phase dataset includes FTD-association results for 6,026,385 variants derived from 2,154 individuals with FTD and 4,308 control of European ancestry.

159

160 Meta-analyzing PD GWAS datasets

We used METAL software [90] to perform an inverse-variance weighted meta-analysis of the three independent PD GWAS summary statistics. We refer to this meta-analyzed PD dataset in the text, tables, and figures as METAPD (49,731 cases and 784,343 controls).

165 Standardization and Filtering of GWAS Summary Statistics

We standardized all summary statistics prior to polygenic analyses. We first confirmed 166 the genome build to be GRCh37, and then annotated variants with dbSNP v151 rs-identifiers 167 and gnomAD [41] non-Finnish European (NFE) allele frequencies using ANNOVAR software 168 169 (2018Apr16) [88]. We only included bi-allelic variants with rs-identifiers and in instances where 170 multiple variants shared the same rs-identifiers, we selected the variant that was supported by 171 the largest number of studies and/or the greatest sample size. Finally, we processed and filtered 172 summary statistics using the munge sumstats.py tool provided with Linkage Disequilibrium Score Regression Software (LDSC) [11]. This processing and filtering removed variants with an 173 effect allele frequency of less than 0.05 in the gnomAD NFE population, variants with strand-174 ambiguous alleles, variants supported by a low sample size or effective sample (N_{eff} = 175 176 $4/(1/N_{cases}+1/N_{controls}))$ for the meta-analysis [90], and variants that were not reported in the 177 HapMap3 study [31]. The number of variants overlapping across all processed GWAS summary statistic datasets analyzed in the present study are presented in Table 1. 178

179

180 Estimating Genetic Overlap by GNOVA

181 We calculated genetic overlap using GNOVA software [54]. GNOVA estimates genetic covariance based on all the genetic variants shared between two GWAS summary statistic 182 datasets. In brief, the summary statistic z-scores observed for each variant are multiplied and 183 their product is regressed against the LD score for that variant, with the LD score being 184 185 calculated based on the external 1000 genomes project CEU population [84]. Genetic covariance is then estimated based on all shared variants using the method of moments and a 186 block-wise jackknife approach as described in the GNOVA manuscript [54]. GNOVA further 187 188 provides an estimate of genetic correlation based on this calculated genetic covariance and the 189 estimated GWAS variant-based heritabilities. As with LD score regression [11], GNOVA is able to statistically correct for any sample overlap between two different sets of GWAS summary 190

191 statistics. In addition, GNOVA produces unbounded genetic correlation estimates which may be 192 greater than one for traits which are highly genetically correlated. GNOVA provides greater statistical power and higher estimation accuracy for genetic correlations than LD score 193 regression, especially when the correlations are moderate [54], as is expected for melanoma 194 195 and PD. We ran GNOVA software on the processed GWAS summary statistics using default 196 parameters and the 1000 Genomes [84] European population-derived reference data provided 197 with the software. Given we test the genetic correlation of melanoma against PD, AD, and FTD we use a Bonferroni corrected significance threshold of $P < 1.67 \times 10^{-02} (0.05 / 3)$ for our 198 199 primary analysis. We also ran annotation-stratified analyses using the minor allele frequency guartile and chromosome annotations provided with GNOVA software as well as the 200 aforementioned reference data and parameters. In the text we present genetic correlations, 201 202 95% confidence intervals, and p-values that have been corrected for sample overlap by 203 GNOVA.

204

205 Disease-Inferred Gene Expression Overlap Analyses

We investigated whether the genetic overlap between PD and melanoma was mediated 206 207 by shared regulation of gene expression. To do this we generated tissue-specific, diseaseinferred gene expression profiles from the processed GWAS summary statistics using 208 FUSION/TWAS software with the default parameters [35]. FUSION/TWAS imputes gene 209 210 expression using *cis* expression quantitative trait loci (eQTL) data derived from reference panels 211 of paired genotype and tissue-specific gene expression data. As gene expression is imputed based on disease-specific GWAS summary statistics, FUSION/TWAS identifies disease-inferred 212 gene expression profiles with tissue-level resolution. For this study, we used eQTL weights 213 214 based on the 48 tissue Genotype-Tissue Expression (GTEx) [34] version 7 (v7) reference panel 215 provided with FUSION/TWAS to generate all disease-inferred gene expression profiles. We tested for overlap or correlation between the disease-inferred gene expression using RHOGE 216

217 software [57], providing the effective sample size [90] for each dataset and only including those 218 FUSION/TWAS results that were at least nominally (p < 0.05) associated with each disease as per the default RHOGE parameters. RHOGE provides an estimate of the genetic correlation 219 220 between two traits that can be attributed to eQTLs as represented by the different trait-inferred 221 gene expression profiles. We exclude the major histocompatibility complex (MHC) region from 222 disease-inferred gene expression overlap analyses due to its complex LD structure [35, 57]. To consider an overlap as significant we used a Bonferroni corrected threshold: $P < 1.04 \times 10^{-03}$ 223 (0.05 / 48 tissues) and present uncorrected p-values and 95% confidence intervals in the text. 224

225

Highlighting Genes Underlying Disease-Inferred Gene Expression Overlap

227 We used UTMOST software [38] to generate single-tissue, disease-inferred gene 228 expression, and then aggregated them into a summary metric representing cross-tissue, 229 eGene-disease associations. eGenes are those genes whose expression are influenced by a least one *cis* disease-associated genetic variant [93]. For this analysis, we generated the single 230 tissue disease-inferred results based on the processed GWAS summary statistics and the 44 231 tissue GTEx v6 reference panel provided with UTMOST, using default parameters. We similarly 232 233 generated the cross-tissue summary metric using default parameters. The UTMOST cross-234 tissue test summary metric represents the maximum one-sided likelihood ratio test statistic for an eGene being associated with the disease, with larger test statistics indicating greater support 235 236 for an association. This summary metric does not include any indicator of uncertainty. We 237 identified transcriptome-wide significant, cross-tissue, eGene-disease associations using a false discovery rate (FDR) threshold of 0.05, that is five expected false discoveries per 100 reported. 238 We compared PD and melanoma UTMOST summary metric eGene results for the disease-239 240 specific GWAS summary statistics to identify eGenes that were independently associated with 241 both diseases.

242

243 Investigating for Differential Expression of Highlighted eGenes in PD Brain Tissues

244 To investigate whether the eGenes we identified as being independently associated with both melanoma and PD demonstrated differential expression in PD, we downloaded publicly 245 available, normalized microarray gene expression data derived from substantia nigra brain 246 247 tissues donated by individuals with and without PD. These datasets were deposited in the Gene 248 Expression Omnibus (GEO) under the accession codes: GDS2821 [47] and GDS3129 [22, 62]. 249 The GDS2821 dataset includes Affymetrix Human Genome U133 Plus 2.0 array data collected 250 from 16 individuals with neuropathologically-confirmed PD and nine aged individuals with no history or pathological diagnosis of neurologic or psychiatric disease [47]. The GDS3129 251 dataset includes Affymetrix Human Genome U133B array data derived from 15 samples of 252 medial substantia nigra and nine samples of lateral substantia nigra from individuals with 253 254 neuropathologically-confirmed PD as well as eight samples of medial substantia nigra and 255 seven samples of lateral substantia nigra from control individuals without neurodegenerative disease pathology [22, 62]. We extracted the normalized expression levels of GPATCH8, 256 MYO9A, PIEZO1, SOX6, TRAPPC2L, ZNF341, and ZNF778 genes and compared the 257 expression between controls using a Mann-Whitney test using Graphpad Prism 8.0. 258

259 **Results**

260 Polygenic Analysis Reveals Specific Genetic Overlap between Melanoma and PD

Prior to cross-disease analyses, we first confirmed that the three independent PD datasets demonstrated positive and significant genetic correlation with each other (genetic correlation range: 0.94 to 1.07, Table 2) using GNOVA software. Following this confirmation and method validation, we proceeded to analyze for potential genetic correlations between melanoma, PD, and the comparator neurodegenerative disease datasets. We identified a significant and positive genetic correlation between melanoma and the meta-analyzed PD dataset (genetic correlation: 0.17, 95% CI 0.10 to 0.24; $P = 4.09 \times 10^{-06}$,

268 Table 3). This result was not driven by any specific PD dataset, but all three independent datasets contributed to the association (P < 0.05; genetic correlation range: 0.14 to 0.25, Figure 269 1 and Table 4). We further investigated the genetic correlation between melanoma and the 270 meta-analyzed PD dataset by stratifying it to the level of minor allele frequency and 271 272 chromosome annotations. Consistent with the polygenic nature of these diseases, we found 273 their genetic correlation to be most highly enriched in those genetic variants annotated as being 274 in the top quartile of minor allele frequency (Supplementary Table 1, online resource). We also 275 found the genetic correlation between melanoma and the meta-analyzed PD dataset to be 276 enriched in chromosomes 1, 2, 8, 11, 16, and 17 (Supplementary Table 2, online resource). We found no shared genetic architecture between melanoma and Alzheimer disease (genetic 277 correlation: -0.02, 95% CI -0.11 to 0.07; P = 0.73, Table 3) nor between melanoma and 278 279 Frontotemporal dementia (genetic correlation: -0.13, 95% CI -0.37 to 0.12; P = 0.32, Table 3). 280 We similarly did not observe any significant correlation between the meta-analyzed PD dataset and AD (Table 3), although one of the individual PD studies showed nominal correlation with AD 281 (Nalls2014: genetic correlation: -0.22, 95% CI -0.22 to 0.00, $P = 4.94 \times 10^{-02}$; Table 4). We did 282 identify a positive and significant genetic correlation between the meta-analyzed PD dataset and 283 284 FTD (genetic correlation: 0.27, 95% CI 0.07 to 0.47; $P = 8.43 \times 10^{-03}$, Table 3), but this appeared to be primarily driven by one of the individual PD studies (Table 4). Together these 285 results demonstrate a consistent, positive and significant genetic correlation between melanoma 286 287 and PD but not between melanoma and FTD or AD.

288

289 PD and Melanoma Disease-Inferred Gene Expression Overlaps Across Tissues

To investigate whether melanoma and PD-associated risk variants regulated the
 expression of the same genes, we generated disease-inferred, tissue-specific gene expression
 profiles from the processed melanoma and METAPD GWAS summary statistic datasets via

FUSION/TWAS software [35]. We further investigated for overlap between the different diseaseinferred gene expression profiles using RHOGE software [57].

We identified a positive and significant overlap between the PD- and melanoma-inferred 295 296 gene expression profiles in a joint analysis of the 48 tissues included in the GTEx v7 reference 297 panel provided with the FUSION/TWAS software (disease-inferred gene expression correlation: 0.14, 95% CI 0.06 to 0.22; P: 7.87 \times 10⁻⁰⁴). Analyzing the PD- and melanoma-inferred gene 298 299 expression correlation in each of the reference panel tissues individually, we observed positive overlap in 44 tissues (disease-inferred gene expression correlation median: 0.25, IQR: 0.13, 300 Figure 2 and Table 5), but only a statistically significant overlap in the suprapubic, non-sun-301 exposed, skin tissue (disease-inferred gene expression correlation: 0.37, 95% CI 0.17 to 0.57; 302 $P: 7.58 \times 10^{-04}$). Eleven additional tissues demonstrated positive and nominal (Figure 2 and 303 304 Table 5) the PD- and melanoma-inferred gene expression overlap including spleen (diseaseinferred gene expression correlation: 0.40, 95% CI 0.13 to 0.66; $P: 5.49 \times 10^{-03}$), minor salivary 305 aland (disease-inferred gene expression correlation: 0.45, 95% CI 0.15 to 0.75; P: 7.49 \times 10⁻⁰³), 306 307 heart atrial appendage (disease-inferred gene expression correlation: 0.31, 95% CI 0.09 to 0.54; $P: 8.27 \times 10^{-03}$) brain substantia nigra (disease-inferred gene expression correlation: 0.42, 95%) 308 309 CI 0.14 to 0.71; P: 9.02×10^{-03}), and brain caudate nucleus (disease-inferred gene expression correlation: 0.29, 95% CI 0.01 to 0.58; P: 4.89 × 10⁻⁰²). 310

To highlight genes whose expression was commonly regulated by PD and melanoma 311 risk variants, we generated cross-tissue, summary metric eGene-disease associations using 312 313 UTMOST [38] software. Applying UTMOST to the METAPD GWAS summary statistics, we identified 606 eGenes significantly associated with PD (Supplementary Table 3, online 314 resource), including genes in previously reported PD-associated loci [50, 64], such as MAPT (P: 315 316 1.28×10^{-04}). In the melanoma dataset, we identified 168 significantly associated eGenes 317 (Supplementary Table 4, online resource) including those reported in a previous TWAS study [92], such as MAFF (P: 1.28×10^{-12}). Comparing the two sets of cross-tissue summary metric 318

319 results, we identify seven eGene-disease associations that passed the FDR threshold for both PD and melanoma: GPATCH8, MYO9A, PIEZO1, SOX6, TRAPPC2L, ZNF341, and ZNF778 320 (Figure 3 and Table 6). In addition, we found evidence for differential expression between 321 individuals with and without neuropathologically-confirmed PD for five of these seven eGenes in 322 323 publicly available substantia nigra microarray datasets (Supplementary Figure 1A-O, online resource). Together, these results suggest that some component of the genetic correlation 324 325 between melanoma and PD may be mediated by the shared regulation of gene expression 326 across tissues.

327

328 Discussion

In this study, we have identified a positive and significant genetic correlation between melanoma and PD by leveraging the largest available GWAS summary statistic datasets and recent advances in polygenic complex trait modeling [11, 54] (Tables 3-4). Our results support the findings of several epidemiologic studies of shared – individual and familial – risk [5, 17, 21, 27, 29, 36, 42, 53, 67, 68, 81, 87, 91] between the two diseases. We also demonstrate no evidence for shared genetic overlap between melanoma and two negative comparison neurodegenerative diseases: AD and FTD (Table 3), suggesting specificity.

Our results of positive genetic correlations between melanoma and PD stand in contrast to negative results from several other genetic studies including single-variant analyses [24, 26, 28, 55, 65, 66] and multi-variant analyses [65, 66]. Both melanoma and PD are complex diseases with inherently polygenic risk architectures. Consequently, efforts to identify shared genetic architecture at the single-variant level are likely underpowered, especially given the moderate epidemiologic and genetic, correlation between melanoma and PD. This is especially true given the fact that the GWAS results analyzed for such single-variant level investigations

343 are themselves currently underpowered. For example, a power analysis reported in the largest 344 PD GWAS to date (Nalls2019), suggests that an adequately powered PD GWAS would require the inclusion of approximately 99,000 PD cases – more than double their current PD case 345 346 sample size [63]. Consequently, our current knowledge regarding the genetic architectures of 347 PD and melanoma is hardly comprehensive and larger GWAS may reveal shared individual risk loci between these diseases in the future. Similarly, previous multi-variant genetic analyses 348 349 investigating melanoma and PD have focused specifically on GWAS-significant loci and thus 350 can be expected to have missed a substantial proportion of the genetic architecture [8] 351 underlying these complex diseases. Genetic correlation methods that consider linkage disequilibrium structure and incorporate all common variants are better powered to detect 352 genetic overlap, especially given current GWAS sample sizes, as we demonstrate here for 353 354 melanoma and PD.

The classification and ascertainment of participants was different between the three 355 356 independent PD datasets included in the present study; however, they all demonstrate positive 357 and significant genetic overlap with each other (Table 2). While this overlap does not guarantee specificity of the represented genetic architecture [12], the fact we observe all three independent 358 359 PD studies to demonstrate positive and significant genetic overlap with melanoma (Figure 1 and 360 Table 4) bolsters confidence in our results. Importantly, although the PD and melanoma genetic 361 correlation point estimates for the three individual PD studies appear different, their 95% confidence intervals overlap which indicates that the effect size estimates are not significantly 362 different (Figure 1 and Table 4). The genetic overlap between the independent PD datasets 363 364 supported their meta-analysis, and the genetic correlation between the meta-analyzed PD dataset and melanoma provided the most precise estimate (genetic correlation: 0.17, 95% CI 365 0.10 to 0.24; $P = 4.09 \times 10^{-06}$; Figure 1 and Tables 3-4). Further increases in precision may 366 result from incorporating additional independent GWAS summary statistic datasets and thus our 367

368 analyses should be repeated as these become available for both melanoma and PD. Similarly, our FTD genetic correlation results should be interpreted with caution as the current sample size 369 is at least one order of magnitude smaller than the other disease datasets. For example, among 370 the individual PD datasets, we only observe a positive genetic correlation between FTD and 371 372 Nalls2019. Parkinsonism has been observed in about 20% on individuals with FTD [2, 7], and this result may suggest that individuals with FTD with parkinsonism were included among the 373 374 UKB-proxy cases in the Nalls2019 dataset. Alternatively, a positive genetic correlation between FTD and the other PD datasets may be observed from the use of a larger FTD GWAS summary 375 statistic dataset. Thus, our analyses should be repeated as larger GWAS summary statistic 376 datasets become available. 377

378 We infer disease-associated gene expression profiles [35] using melanoma and meta-379 analyzed PD GWAS summary statistics and investigate for their overlap at the level of tissues [57] and genes [38] to provide bioinformatically-driven biological context to our melanoma and 380 381 PD genetic correlation results. We identify significant cross-tissue overlap (disease-inferred gene expression correlation: 0.14, 95% CI 0.06 to 0.22; $P: 7.87 \times 10^{-04}$) and significant individual 382 tissue overlap in suprapubic non-sun-exposed skin (disease-inferred gene expression 383 correlation: 0.37, 95% CI 0.17 to 0.57; *P*: 7.58 × 10⁻⁰⁴). We also observe positive, nominal 384 385 disease-inferred gene expression correlation in peripheral tissues with PD relevance like the 386 heart atrial appendage (disease-inferred gene expression correlation: 0.31, P < 0.05, Table 5) which may reflect the cardiac sympathetic denervation associated with PD [32, 82] - or the 387 minor salivary glands (disease-inferred gene expression correlation: 0.45, P < 0.05, Table 5) -388 389 which have been reported in some, but not all, studies as containing alpha synuclein aggregates 390 in the context of PD [46, 85]. In terms of PD-relevant brain tissues, we observe positive, nominal disease-inferred gene expression correlation in the substantia nigra and basal ganglia caudate 391 nucleus (disease-inferred gene expression correlation: 0.42 and 0.29, respectively; P < 0.05, 392

393 Figure 2 and Table 5). Importantly, the available GTEx v7 inferred gene expression reference model for brain tissues are based on substantially fewer samples than most peripheral tissues, 394 for example the brain substantia nigra reference is derived from 80 donors compared to 335 395 donors for the suprapubic skin reference (Table 5). Consequently, our disease-inferred gene 396 397 expression risk profile overlap analyses should be repeated as larger reference panels become 398 available. Similarly, another limitation of the GTEx dataset is the inclusion of tissues from 399 individuals with extended post-mortem intervals. As this can be expected to result in an 400 underrepresentation of short-lived transcripts in the inferred gene expression reference panels, 401 our analyses should be repeated as reference panels based on tissues from individuals with shorter post-mortem intervals become available. 402

403 We identify seven cross-tissue, eGene-disease associations passing the FDR threshold 404 for both melanoma and PD (Figure 3 and Table 6), most of which are located on the chromosomes which we identified as being enriched for the genetic correlation between these 405 406 two diseases. Importantly, the UTMOST software currently only provides a compatible reference panel based on the GTEx v6 release which is derived from fewer donor samples per tissue 407 compared to GTEx v7 release. In addition, the GTEx v6 reference panel does not include four 408 409 tissues - brain substantia nigra, brain spinal cervical spinal cord, brain amygdala, and minor 410 salivary gland - which we observed to demonstrate positive disease-inferred gene expression 411 overlap for melanoma and PD (Table 5). Additional eGenes may pass the FDR threshold for both PD and melanoma in analyses based on the larger GTEx v7 reference panel. Thus, our 412 analyses should be repeated when this or other larger reference panels become available for 413 UTMOST. Nevertheless, using the smaller GTEx v6 reference panel we identify seven genes 414 415 that may be commonly regulated by melanoma and PD-associated variants under the FDR threshold (Figure 3 and Table 6), including *PIEZO1* (Melanoma P: 2.74×10^{-11} ; METAPD P: 416

417 5.65 × 10⁻⁰⁵); *TRAPPC2L* (Melanoma *P*: 2.36 × 10⁻¹¹; METAPD *P*: 8.47 × 10⁻⁰⁵); and *SOX6* 418 (Melanoma *P*: 1.30 × 10⁻⁰⁴; METAPD *P*: 5.97 × 10⁻⁰⁵).

419 PIEZO1 encodes a recently described mechanosensitive cation channel [15] with several biological functions including human T cell activation [52], direction of lineage choice in 420 421 human neural stem cells [71], and mediating the age-related loss of function of oligodendrocyte progenitor cells [79]. PIEZO1 is expressed in the neurons of the human substantia nigra [20, 422 76] and also is ubiquitously expressed in human enteric neurons [58], both neuronal types 423 impacted by PD [10, 43]. Interestingly, the expression of PIEZO2 - PIEZO1's paralog - is 424 425 regulated by, putatively melanocyte-derived, dopamine signaling in mouse primary sensory neurons [69] but whether this regulation is relevant for *PIEZO1* is currently unknown. Similarly, a 426 427 role for PIEZO1 in melanoma remains largely unexplored though PIEZO1 has been identified to 428 contribute to the migration of invasive melanoma cells [39].

429 TRAPPC2L is a component of transport protein particle (TRAPP) complexes which 430 function in intracellular vesicle-mediated transport and autophagy [60, 61, 78]. This gene is 431 expressed in human substantia nigra neurons [20] and a homozygous missense variant in it causes a neurodevelopmental disorder characterized by progressive encephalopathy and 432 episodic rhabdomyolysis [60]. The intergenic variant rs12921479 - which is an eQTL for 433 TRAPPC2L in the brain [34, 74] – was reported to be associated with PD ($P: 9.31 \times 10^{-07}$) in an 434 autopsy-confirmed cohort of PD [3], but is only nominally associated with PD in our meta-435 analyzed PD dataset (P: 1.01×10^{-02}). A role for TRAPPC2L in melanoma remains to be 436 explored. 437

SOX6 is a transcription factor which was recently identified as a determinant of
 substantia nigra neuron development and maintenance [70]. Its expression was observed to
 localize to pigmented and tyrosine hydroxylase positive neurons but not to pigment-negative

441 neurons within the substantia nigra [70]. In addition, SOX6 expression was diminished in the 442 substantia nigra of individuals with PD and deletion of SOX6 in mice was observed to decrease dopamine levels and innervation in the striatum [70], a brain region that is also impacted in PD 443 444 [9]. In a separate study, a large deletion in SOX6 was identified in a patient with global 445 developmental delay and progressive parkinsonian symptoms including rest tremor [77]. 446 Interestingly, SOX6 has been identified as a determinant of gastric dopaminergic neuron 447 development [59], which may suggest a role for this gene in the enteric nervous system dysfunction and pathology observed in PD. SOX6 may also have a role in melanoma. In a 448 cancer cell line expression study, SOX6 was found to be highly expressed in melanoma cells 449 but was not detectable in eight other cancers [86]. Additionally, SOX6 was identified as a 450 candidate melanoma driver gene [72] in a screen and SOX6 may be a melanoma stem cell 451 452 marker [51].

While we observe evidence for differential expression between neuropathologically-453 confirmed PD and controls for PIEZO1, TRAPPC2L, and SOX6 in at least one substantia nigra 454 microarray dataset, these results should be interpreted with caution. Neurodegenerative 455 456 diseases like PD are characterized by dramatic changes in cell-type proportions [49] which will 457 impact differential expression results. Thus, the PD-associated differential expression of the 458 eGenes highlighted in this study should be confirmed in larger, RNA-sequencing-based 459 datasets - as these become available - in order to allow for the inclusion of important covariates like cell-type proportions, sex, age of death, and RNA quality among others. Nevertheless, the 460 461 fact we observe differential expression of SOX6 in the same direction as previously published [70] is reassuring. 462

Investigating for differential expression of the eGenes highlighted in this study in the
context of melanoma is challenging given our focus on the risk of developing melanoma.
Nevertheless, a recent GTEx v8-based, multi-tissue TWAS resource (phenomexcan.org) [73]

466 provides some evidence for a link between the eGenes we highlight and melanoma-associated 467 pigmentation traits included in the UK Biobank study. For example, *PIEZO1* is associated with 468 red hair (*P*: ~0), ease of skin tanning (*P*: 3.74×10^{-175}), and skin colour (*P*: 3.41×10^{-121}); 469 *TRAPPC2L* is associated with red hair (*P*: 3.28×10^{-181}), ease of skin tanning (*P*: 1.06×10^{-71}), 470 and skin colour (*P*: 6.24×10^{-55}); and *SOX6* is associated with ease of skin tanning (*P*: 1.40×10^{-121})

471 10⁻¹³), skin colour (*P*: 1.55×10^{-11}), and childhood sunburn occasions (*P*: 3.92×10^{-11}).

Together, these results support a biologically plausible role for *PIEZO1*, *TRAPPCL2*, and *SOX6* in the genetic correlation between melanoma and PD, but these findings require confirmation and further investigation with future experimental work.

475 PD and melanoma are clinically heterogenous diseases [16, 18] for which spatiotemporal environmental exposures are relevant [14, 16] and may be necessary, in 476 addition to innate genetic susceptibility, for the development of sporadic disease. Consequently, 477 the moderate genetic correlation we observe should not be interpreted as suggesting that these 478 479 diseases will always be co-morbid. However, our results of replicable and significant genetic 480 correlation, regardless of the magnitude of effect, do suggest that these two very different diseases share common biological pathways. Thus, even if only a minority of individuals with 481 PD ultimately develop melanoma, understanding the genetic correlation between these disease 482 483 at the molecular level – for example, if and how the regulation of *PIEZO1*, *TRAPPC2L*, and SOX6 and their related biological pathways contribute to PD etiopathogenesis – may provide 484 mechanistic insight that is generalizable to all individuals with PD. Our results support such 485 future research efforts. 486

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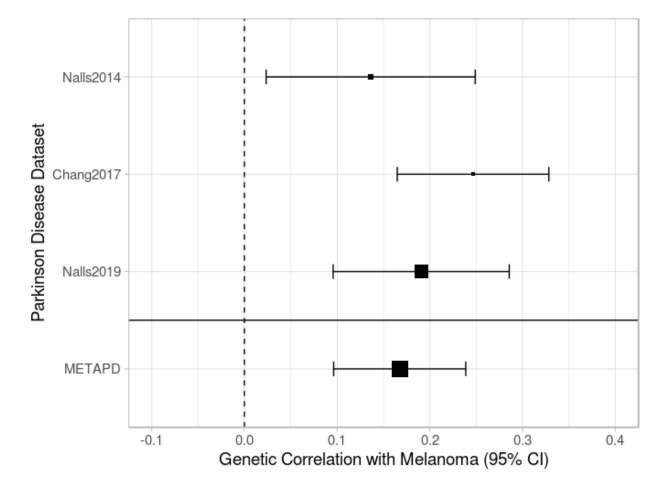
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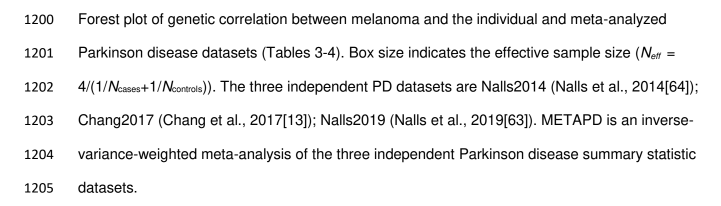
1193

- 1195 Figures
- 1196
- 1197 Figure 1. GNOVA Genetic Correlation Results for Parkinson Disease and Melanoma



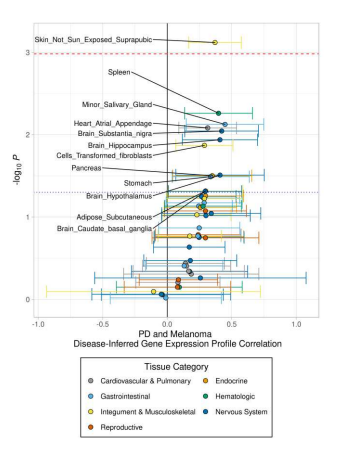
1198 GWAS Summary Statistic Datasets





1206 Figure 2. Parkinson Disease (PD) and Melanoma Tissue-specific, Disease-inferred Gene



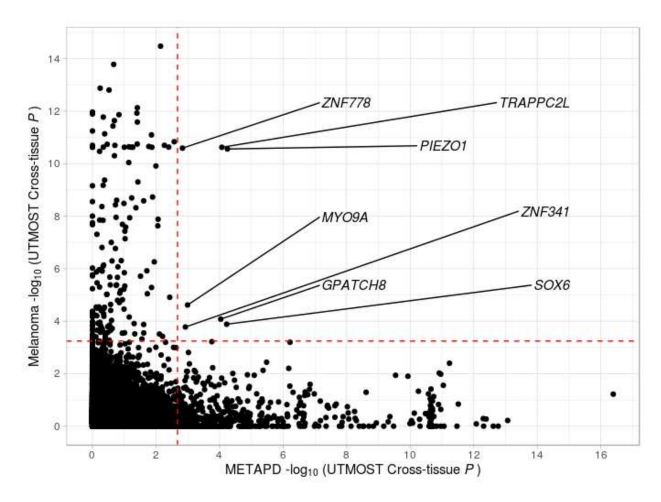


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PD and Melanoma disease-inferred gene expression profile correlation at the level of 48 specific 1209 tissues included in the GTEx v7 reference panel (Table 5). Disease-inferred gene expression 1210 profiles were generated from the processed melanoma and METAPD summary statistics using 1211 1212 FUSION/TWAS software and correlation between these profiles was estimated using RHOGE software. METAPD is an inverse-variance-weighted meta-analysis of the three independent 1213 Parkinson disease summary statistic datasets. The red dashed line demarks the multiple test 1214 corrected *P* threshold of 1.04×10^{-03} (0.05 / 48) while the blue dotted line demarks the nominal 1215 threshold, P = 0.05. 1216

1218 Figure 3. Cross-tissue eGenes Associated with Both Parkinson Disease (PD) and





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Conjunction plot of the cross-tissue PD and melanoma eGene -log₁₀ *P* values. We generated cross-tissue eGene-disease results (Supplementary Tables 3-4, online resource) from the processed melanoma and METAPD summary statistics using UTMOST software. METAPD is an inverse-variance-weighted meta-analysis of the three independent Parkinson disease summary statistic datasets. The red dashed lines demark the false discovery rate (FDR) threshold of 0.05. Labels and lines indicate eGenes associated with both PD and melanoma under the FDR threshold.

1229 Table 1. Number of Overlapping Variants in Processed GWAS Summary Statistic

1230	Datasets
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alasels							
Dataset	Melanoma Law 2015	PD Nalls 2014	PD Chang 2017	PD Nalls 2019	METAPD	AD Kunkle 2019	FTD Ferrar 2014
Law 2015	1,038,973	-	-	-	-	-	-
Nalls 2014	997,418	1,015,955	-	-	-	-	-
Chang 2017	1,038,516	1,015,498	1,075,906	-	-	-	-
Nalls 2019	1,007,785	983,012	1,033,569	1,034,607	-	-	-
METAPD	1,007,521	983,023	1,032,819	1,033,287	1,033,303	-	-
Kunkle 2019	1,038,796	1,015,849	1,075,582	1,034,409	1,033,126	1,077,308	-
Ferrari 2014	979,084	973,381	993,831	961,697	961,512	994,078	994,33

All GWAS summary statistic datasets were standardized and filtered using the same pipeline. 1231 We annotated all variants with dbSNP v151 rs-identifiers and gnomAD non-Finnish European 1232 1233 (NFE) allele frequencies. We filtered variants as to only include bi-allelic variants with rsidentifiers and further removed variants with an effect allele frequency less than 0.05, variants 1234 with strand ambiguous alleles, variants with limited support, i.e. those supported by a low 1235 sample or study number, and variants that were not reported in the HapMap3 study. Presented 1236 1237 are the numbers of variants overlapping between each dataset. METAPD is an inverse-1238 variance-weighted meta-analysis of the three independent Parkinson disease summary statistic datasets. PD: Parkinson disease; AD: Alzheimer disease; FTD: Frontotemporal dementia. 1239

1241 Table 2. GNOVA Genetic Correlation Results for independent Parkinson Disease

1242 Datasets

Parkinson Disease Dataset	Nalls2014	Chang2017	Nalls2019	METAPD
Nalls2014 n _{Case} = 9,581 n _{Control} = 33,245	-	-	-	-
Chang2017 n _{Case} = 6,476 n _{Control} = 302,042	0.95 [0.77, 1.12] (4.16 × 10⁻²⁶)	-	-	-
Nalls2019 n _{Case} = 33,674 n _{Control} = 449,056	1.07 [0.90, 1.25] (7.91 × 10⁻³⁴)	0.94 [0.80, 1.09] (1.43 × 10⁻³⁶)	-	-
$\begin{array}{c} \textbf{METAPD} \\ n_{Case} = 49,731 \\ n_{Control} = 784,343 \end{array}$	1.00 [0.83, 1.18] (1.04 × 10⁻²⁸)	0.71 [0.56, 0.86] (8.09 × 10⁻²¹)	1.06 [0.91, 1.21] (6.10 × 10⁻⁴²)	-

We estimated the genetic correlation between the independent Parkinson disease datasets using GNOVA software. All correlation estimates, 95% confidence intervals – presented in square brackets - and p-values - presented in parentheses - are corrected for any potential sample overlap. GNOVA genetic correlation estimates are unbounded and thus may be greater than 1. METAPD is an inverse-variance-weighted meta-analysis of the three independent Parkinson disease summary statistic datasets.

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1252 Table 3. GNOVA Genetic Correlation Results for Meta-analyzed Parkinson Disease,

1253 Melanoma, and Comparator Neurodegenerative Diseases GWAS Summary Statistic 1254 Datasets

Summary Statistic Dataset	Melanoma Law2015	PD METAPD	AD Kunkle2019	FTD Ferrari2014
Melanoma Law2015 n _{Case} = 12,814 n _{Control} = 23,203	-	-	-	-
PD METAPD n _{Case} = 49,731 n _{Control} = 784,343	0.17 [0.10, 0.24] (4.09 × 10⁻⁰⁶)	-	-	-
AD Kunkle2019 n _{Case} = 21,982 n _{Control} = 41,944	-0.02 [-0.11, 0.07] (0.73)	0.01 [-0.06, 0.09] (0.71)	-	-
FTD Ferrari2014 $n_{Case} = 2,154$ $n_{Control} = 4,308$	-0.13 [-0.37, 0.12] (0.32)	0.27 [0.07, 0.47] (8.43 × 10⁻⁰³)	0.22 [-0.05, 0.49] (0.11)	-

We estimated the genetic correlation between diseases using processed disease-specific GWAS summary statistic datasets and GNOVA software. All correlation estimates, 95% confidence intervals – presented in square brackets - and p-values - presented in parentheses - are corrected for any potential sample overlap. METAPD is an inverse-variance-weighted meta-analysis of the three independent Parkinson disease summary statistic datasets. PD: Parkinson disease; AD: Alzheimer disease; FTD: Frontotemporal dementia.

1262 Table 4. GNOVA Genetic Correlation Results for Independent Parkinson Disease,

Melanoma, and Comparator Neurodegenerative Diseases GWAS Summary Statistic 1263 Datasets

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Summary Statistic Dataset	Melanoma Law2015 n _{Case} = 12,814 n _{Control} = 23,203	AD Kunkle2019 n _{Case} = 21,982 n _{Control} = 41,944	FTD Ferrari2014 n _{Case} = 2,154 n _{Control} = 4,308
PD Nalls2014 $n_{Case} = 9,581$ $n_{Control} = 33,245$	0.14 [0.02, 0.25] (1.79 × 10⁻⁰²)	-0.11 [-0.22, 0.00] (4.94 × 10 ⁻⁰²)	0.27 [-0.06, 0.60] (0.10)
PD Chang2017 n _{Case} = 6,476 n _{Control} = 302,042	0.25 [0.16, 0.33] (3.31 × 10⁻⁰⁹)	-0.01 [-0.11, 0.09] (0.87)	-0.16 [-0.45, 0.12] (0.26)
PD Nalls2019 n _{Case} = 33,674 n _{Control} = 449,056	0.19 [0.10, 0.29] (8.28 × 10⁻⁰⁵)	0.05 [-0.04, 0.14] (0.27)	0.40 [0.14, 0.66] (2.78 × 10⁻⁰³)

We estimated the genetic correlation between diseases using processed disease-specific GWAS 1265 summary statistic datasets and GNOVA software. All correlation estimates, 95% confidence 1266 intervals - presented in square brackets - and p-values - presented in parentheses - are corrected 1267 1268 for any potential sample overlap. PD: Parkinson disease; AD: Alzheimer disease; FTD: 1269 Frontotemporal dementia.

GTEx v7 Tissue	Number of Samples in	Melanoma vs. METAPD	
GTEX V/ TISSUE	Tissue Reference Panel	ρ ge	p-value
Adipose Subcutaneous	385	0.30 [0.01, 0.59]	4.82 × 10-
Adipose Visceral Omentum	313	0.23 [-0.03, 0.49]	9.39 × 10-
Adrenal Gland	175	0.25 [-0.10, 0.59]	1.73 × 10-
Artery Aorta	267	0.14 [-0.16, 0.44]	3.64 × 10-
Artery Coronary	152	0.19 [-0.34, 0.71]	4.93 × 10-
Artery Tibial	388	0.15 [-0.19, 0.49]	3.93 × 10-
Brain Amygdala	88	0.25 [-0.10, 0.60]	1.77 × 10 ⁻
Brain Anterior cingulate cortex BA24	109	0.17 [-0.28, 0.62]	4.58 × 10 ⁻
Brain Caudate basal ganglia	144	0.29 [0.01, 0.58]	4.89 × 10-
Brain Cerebellar Hemisphere	125	0.18 [-0.18, 0.54]	3.38 × 10 ⁻
Brain Cerebellum	154	0.17 [-0.11, 0.45]	2.32 × 10 ⁻
Brain Cortex	136	-0.04 [-0.51, 0.43]	8.75 × 10 ⁻
Brain Frontal Cortex BA9	118	-0.05 [-0.58, 0.49]	8.67 × 10 ⁻
Brain Hippocampus	111	0.41 [0.12, 0.70]	1.15 × 10 ⁻
Brain Hypothalamus	108	0.41 [0.07, 0.75]	3.09 × 10 ⁻
Brain Nucleus accumbens basal ganglia	130	0.34 [-0.04, 0.73]	9.04 × 10 ⁻
Brain Putamen basal ganglia	111	0.30 [-0.04, 0.64]	9.60 × 10 ⁻
Brain Spinal cord cervical c-1	83	0.26 [-0.56, 1.08]	5.49 × 10 ⁻
Brain Substantia nigra	80	0.42 [0.14, 0.71]	9.02 × 10 ⁻
Breast Mammary Tissue	251	0.24 [-0.09, 0.57]	1.64 × 10 ⁻
Cells EBV-transformed lymphocytes	117	0.09 [-0.39, 0.58]	7.11 × 10 ⁻
Cells Transformed fibroblasts	300	0.29 [0.07, 0.51]	1.35 × 10 ⁻
Colon Sigmoid	203	-0.01 [-0.44, 0.42]	9.60 × 10 ⁻
Colon Transverse	246	0.24 [-0.10, 0.57]	1.70 × 10 ⁻
Esophagus Gastroesophageal Junction	213	0.28 [-0.00, 0.56]	5.88 × 10 ⁻
Esophagus Mucosa	358	0.13 [-0.17, 0.43]	3.92 × 10 ⁻
Esophagus Muscularis	335	0.24 [-0.02, 0.51]	7.36 × 10 ⁻
Heart Atrial Appendage	264	0.31 [0.09, 0.54]	8.27 × 10 ⁻
Heart Left Ventricle	272	0.08 [-0.24, 0.41]	6.22 × 10 ⁻
Liver	153	0.25 [-0.07, 0.56]	1.36 × 10 ⁻
Lung	383	0.17 [-0.27, 0.60]	4.54 × 10 ⁻
Minor Salivary Gland	85	0.45 [0.15, 0.75]	7.49 × 10 ⁻
Muscle Skeletal	491	0.17 [-0.07, 0.42]	1.70 × 10 ⁻
Nerve Tibial	361	0.27 [-0.00, 0.53]	5.61 × 10 ⁻

1271Table 5. Disease-Inferred Gene Expression Profile Overlap between Melanoma and PD in1272GTEx v7 Reference Panel Tissues

Ovary	122	0.30 [-0.12, 0.71]	1.79 × 10 ⁻⁰¹
Pancreas	220	0.35 [0.04, 0.66]	3.15 × 10 ⁻⁰²
Pituitary	157	0.30 [0.00, 0.59]	5.54 × 10 ⁻⁰²
Prostate	132	0.08 [-0.33, 0.49]	7.10 × 10 ⁻⁰¹
Skin Not Sun Exposed Suprapubic	335	0.37 [0.17, 0.57]	7.58 × 10 ⁻⁰⁴
Skin Sun Exposed Lower leg	414	0.29 [-0.01, 0.58]	5.96 × 10 ⁻⁰²
Small Intestine Terminal Ileum	122	0.29 [-0.01, 0.58]	6.71 × 10 ⁻⁰²
Spleen	146	0.40 [0.13, 0.66]	5.49 × 10 ⁻⁰³
Stomach	237	0.34 [0.04, 0.64]	3.23 × 10 ⁻⁰²
Testis	225	0.09 [-0.22, 0.39]	5.78 × 10 ⁻⁰¹
Thyroid	399	0.26 [-0.02, 0.54]	7.66 × 10 ⁻⁰²
Uterus	101	0.30 [-0.02, 0.61]	8.43 × 10 ⁻⁰²
Vagina	106	-0.11 [-0.93, 0.72]	8.05 × 10 ⁻⁰¹
Whole Blood	369	0.28 [-0.02, 0.57]	7.38 × 10 ⁻⁰²

1273 We generated disease-inferred gene expression profiles based on standardized and processed 1274 GWAS summary statistics using FUSION/TWAS software and the Genotype-Tissue Expression 1275 Project (GTEx) v7 reference panel. We further compared the overlap of these disease-inferred 1276 gene expression profiles using RHOGE software. METAPD is an inverse-variance-weighted 1277 meta-analysis of the three independent Parkinson disease summary statistic datasets. PD: 1278 Parkinson disease; ρ_{GE} : correlation coefficient for inferred transcriptomic overlap; BA: Brodmann 1279 Area.

Gene	Melanoma UTMOST Cross-tissue		PD UTMOST Cross-tissue	
	Test Metric	P	Test Metric	Р
GPATCH8	9.27	8.33 × 10 ⁻⁰⁵	9.18	9.17 × 10 ⁻⁰⁵
MYO9A	10.10	2.41 × 10 ⁻⁰⁵	6.47	1.01 × 10 ⁻⁰³
PIEZO1	176.52	2.74 × 10 ⁻¹¹	9.29	5.65 × 10 ⁻⁰⁵
SOX6	9.02	1.30 × 10 ⁻⁰⁴	9.77	5.97 × 10 ⁻⁰⁵
TRAPPC2L	690.56	2.36 × 10 ⁻¹¹	9.27	8.47 × 10 ⁻⁰⁵
ZNF341	8.42	1.67 × 10 ⁻⁰⁴	6.57	1.19 × 10 ⁻⁰³
ZNF778	219.82	2.55 × 10 ⁻¹¹	6.07	1.47 × 10 ⁻⁰³

1281 Table 6. Cross-Tissue eGene-Disease Associations for Melanoma and PD

1282 We inferred cross-tissue, eGene-disease associations based on standardized and processed 1283 melanoma and METAPD GWAS summary statistics using UTMOST software and the

1284 Genotype-Tissue Expression Project (GTEx) v6 reference panel. METAPD is an inverse-

variance-weighted meta-analysis of the three independent Parkinson disease (PD) summary

1286 statistic datasets.