

1 Mosaic patterns of selection in genomic regions associated with diverse human traits

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24 Abstract:

25 Natural selection shapes the genetic architecture of many human traits. However, the prevalence of
26 different modes of selection on genomic regions associated with variation in traits remains poorly
27 understood. To address this, we developed an efficient computational framework to calculate enrichment
28 of different evolutionary measures among regions associated with complex traits. We applied the
29 framework to summary statistics from >900 genome-wide association studies (GWASs) and 11
30 evolutionary measures of sequence constraint, population differentiation, and allele age while accounting
31 for linkage disequilibrium, allele frequency, and other potential confounders. We demonstrate that this
32 framework yields consistent results across GWASs with variable sample sizes, numbers of trait-
33 associated SNPs, and analytical approaches. The resulting evolutionary atlas maps diverse signatures of
34 selection on genomic regions associated with complex human traits on an unprecedented scale. We
35 detected positive enrichment for sequence conservation among trait-associated regions for the majority of
36 traits (>77% of 290 high power GWASs), which was most dominant in reproductive traits. Many traits also
37 exhibited substantial enrichment for population differentiation and recent positive selection, especially
38 among hair, skin, and pigmentation traits. In contrast, we detected widespread negative enrichment for
39 balancing selection (51% GWASs) and no evidence of enrichment for selection signals in regions
40 associated with late-onset Alzheimer's disease. These results support a pervasive role for negative
41 selection on regions of the human genome that contribute to variation in complex traits, but also
42 demonstrate where diverse modes of selection have shaped trait-associated loci. This atlas of signatures
43 of different modes of natural selection across the diversity of available GWASs will enable exploration of
44 the relationship between the genetic architecture and selection in the human genome.

45 Introduction

46 Understanding how natural selection has shaped the human genome is fundamental for
47 evolutionary genomics and medicine ¹. As humans expanded out of Africa, they encountered
48 diverse climates, underwent dietary changes, experienced demographic shifts, and mixed with
49 Neanderthals and other hominins. The selective pressures exerted by these events shaped the
50 genetic basis of modern human traits ²⁻⁵. Two well known examples include the strong positive
51 selection on adult milk consumption that shaped frequencies of lactase persistence alleles ⁶⁻⁸
52 and a Denisovan introgressed haplotype that contributed to high-altitude adaptation of Tibetans
53 ^{9,10}. Although the evolutionary histories of these and several other specific loci and traits have
54 been studied ¹¹⁻¹⁴, the extent and types of evolutionary forces that have acted on the genomic
55 regions associated with variation in the human phenome remain poorly understood.

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57 Multiple measures have been developed to infer evolutionary forces from patterns of genetic
58 variation within and between species ¹⁵. For example, comparing human genomes to those of
59 related species using measures like PhyloP and PhastCons enable testing hypotheses about
60 decreases and increases in the substitution rate over evolutionary time that are often indicative
61 of the action of negative and positive selection, respectively ^{16,17}. Identification of clusters of
62 variants at intermediate allele frequencies in human populations by measures such as the Beta
63 Score enables inference of balancing selection ^{18,19}. Similarly, measures such as F_{ST} and XP-
64 EHH rely on single nucleotide polymorphism (SNP) and haplotype structures to detect local
65 adaptation or recent positive selection between human populations ²⁰. It is also possible to
66 estimate the time to the most recent common ancestor of different haplotypes and quantify the
67 age/origin of variants using ancestral recombination graphs ²¹. Driven by increasing amounts of
68 whole genome sequence data and computational power, more recent methods, such as
69 RELATE ²² and CLUES ²³, use locally constructed genealogies and ancestral recombination
70 graphs to infer allele histories and detect recent directional selection. Other methods rely on
71 parametric models of neutral evolution ²⁴ or analyze patterns of singleton variants ²⁵ that
72 incorporate population level genomic data and GWAS summary statistics to estimate the
73 strength of selection and evidence for directional selection ^{13,26}. Together, these evolutionary
74 measures capture evidence for a diverse set of evolutionary forces from signatures in genetic
75 variation.

76
77 Despite advances in these methods, which mainly focus on individual regions, mapping the
78 evolutionary pressures on complex traits remains challenging for several reasons. First,
79 genomic attributes that influence ascertainment and power in association studies, e.g., allele
80 frequency and linkage disequilibrium (LD), also influence the expected distribution of many
81 evolutionary metrics. Thus, the genomic background does not provide an appropriate null when
82 interpreting overlaps between trait associations and signatures of selection. Second, population
83 stratification is common in genome-wide association studies (GWASs). As GWASs became
84 more prevalent and demonstrated that most common traits are polygenic, new trait-focused
85 approaches to detect evidence of recent polygenic selection emerged. Polygenic scores, which
86 can be derived by summing across trait-associated alleles from a GWAS after weighting by the
87 effect size, enable prediction of phenotype from genotype. Several studies computed polygenic
88 scores across populations and interpreted systematic differences and the alleles that drive them

89 as evidence of polygenic adaptation^{27–29}. For example, human height increasing alleles
90 identified from GWAS were found to be at consistently higher frequencies in Northern European
91 populations compared to Southern Europeans²⁹. However, subsequent analyses revealed that
92 residual population stratification in the GWASs and a resulting lack of consistent effects across
93 populations drove the initial signatures of selection^{30–34}. Detecting and correcting for residual
94 stratification is an ongoing challenge in the field. Despite these complications, some clear
95 evolutionary patterns have emerged; regions of the genome that have been associated with
96 complex traits, such as hair color, body mass index, waist-to-hip ratio etc., consistently show
97 evidence of recent and directional selection^{13,22,35,36}.

98
99 In this study, we describe a unified approach to determine enrichment for evolutionary forces
100 acting on regions associated with variation in diverse complex traits. This approach is
101 complementary to previous work on polygenic adaptation that focused on the traits themselves
102^{24,35,37} because we are characterizing the evolutionary history of the genomic regions that
103 contribute to complex trait variation. To protect against biases from stratification, our approach:
104 1) does not directly incorporate effect sizes at trait-associated regions (e.g. as in polygenic
105 scores), 2) builds a null distribution from allele frequency and LD-matched SNPs, and 3)
106 enables flexible enrichment testing at different association thresholds. We generate an atlas of
107 11 evolutionary measures on regions identified from GWASs of over 900 polygenic traits
108 (totalling 210,109 genomic regions). We find widespread enrichment for signatures of negative
109 selection, a dearth of balancing selection, and several groups of GWASs that show distinct
110 enrichments for signals of population differentiation and recent positive selection. By mapping
111 the evolutionary landscape of genomic regions that underlie specific complex traits, these
112 results reveal that human trait-associated regions have been shaped by a mosaic of different
113 modes of selection.

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115

116 **Results**

117 **An efficient permutation-based approach to detect evolutionary forces on GWAS** 118 **loci**

119 To quantify genomic signatures of diverse evolutionary forces acting on genomic regions
120 associated with complex human traits, we developed an empirical framework that infers
121 enrichment for diverse evolutionary signatures from GWAS summary statistics. For a given
122 GWAS, we consider independent trait-associated genomic regions accounting for LD ($r^2 > 0.9$,
123 GWAS p-value $< 5e-8$, Figure 1a).

124

125 To define an appropriate background distribution for each analysis, we randomly select genomic
126 regions matched on minor allele frequency, LD patterns, and gene proximity for each trait-
127 associated region. The matching is repeated until we have 5,000 sets that each contain the
128 same number of genomic regions as the trait-associated regions (Figure 1b). For each
129 evolutionary measure, we build a background distribution from each matched set. We then
130 compare the observed trait-level evolutionary values to the background distribution and
131 calculate an empirical p-value (Figure 1c,d). To summarize each comparison, we define the
132 standardized evolutionary enrichment as the difference between the observed trait-level mean

133 and the matched-background mean divided by the genome-wide standard deviation for the
 134 evolutionary measure (Figure 1e). However, we note that any summary statistic could be used.

135
 136 We apply this approach for 11 evolutionary measures that detect patterns of genomic variation
 137 consistent with the action of different modes of selection, such as directional selection,
 138 balancing selection, local adaptation, and negative selection. All evolutionary measures had
 139 high coverage (83-99%) across the set of SNPs used in our study (Methods, Table 1).

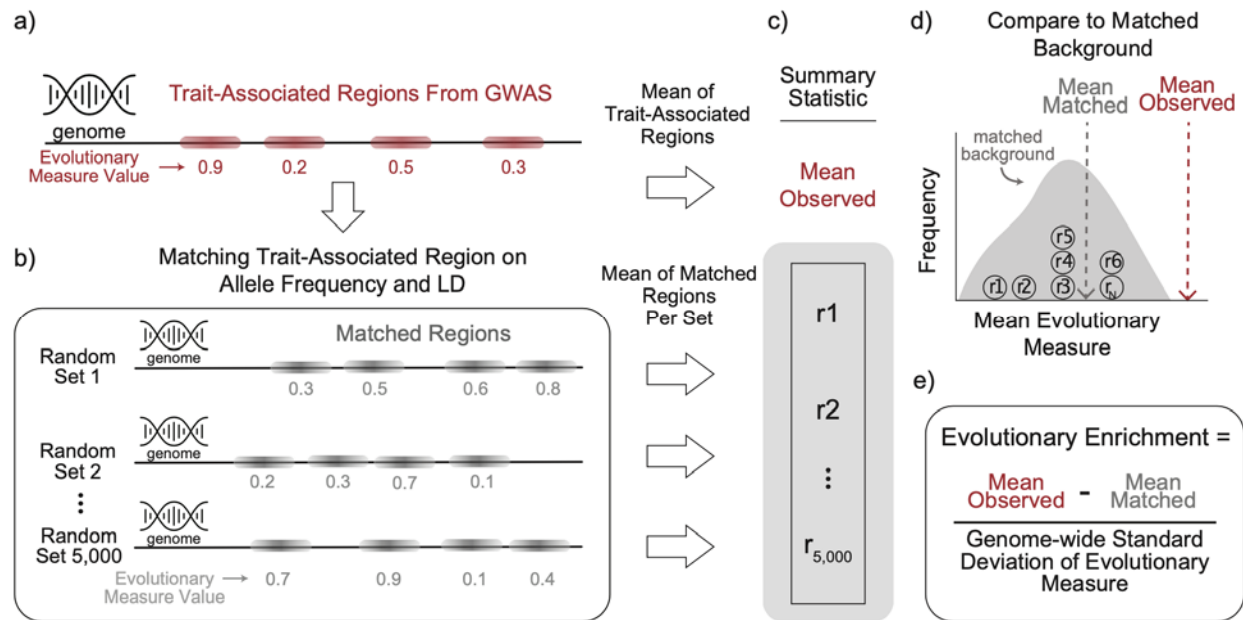
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Evolutionary Measure	Type of Evolutionary Force	Time Scale	%SNPs covered
ARGweaver (TMRCA)	Evolutionary Origin	Human population	99%
Beta Score	Balancing Selection	Human Population	99%
PhyloP	Positive/Negative selection	Across species	98%
PhastCons	Negative Selection	Across species	98%
LINSIGHT	Negative Selection	Across species & Human populations	98%
F_{ST} afr-eas F_{ST} afr-eur F_{ST} eas-eur	Positive Selection	Human populations	99%
XP-EHH afr-eas XP-EHH afr-eur XP-EHH eas-eur	Positive Selection	Human populations	83-86%

144 **Table 1: Evolutionary measures used to quantify different types of evolutionary forces on**
 145 **trait-associated regions.**

146 For each evolutionary measure (rows), the type of evolutionary force inferred and the
 147 corresponding time scale is given. “%SNPs covered” is the proportion of SNPs from 1000
 148 Genomes Phase III after quality control (n=9,535,059) that have an annotation for the given
 149 evolutionary measure. For F_{ST} and XP-EHH, we used the following 1000 genomes
 150 superpopulation comparisons: afr-eas, afr-eur, eas-eur. XP-EHH: cross-population extended
 151 haplotype homozygosity (EHH). TMRCA: time to most recent common ancestor derived from
 152 ARGweaver.

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157 **Figure 1: Computational framework for detecting enrichment for genetic signatures of**
 158 **evolutionary forces from genome-wide association studies (GWASs).**

159 (a) Given the GWAS of a complex trait, we define trait-associated regions by first identifying
 160 variants of genome-wide significance and then clumping based on linkage disequilibrium (LD;
 161 e.g., $r_2 > 0.9$). For each region, we identify the maximum value of an evolutionary measure of
 162 interest. (b) For each trait-associated region, we identify 5,000 randomly selected genomic
 163 regions ("matched regions") that have similar minor allele frequency and linkage disequilibrium
 164 patterns (Methods). (c) Across the trait-associated regions and their matched random genomic
 165 regions, we calculate a summary statistic. To illustrate our approach, we take the mean of the
 166 evolutionary measure to generate an (d) empirical background distribution and (e) calculate
 167 enrichment by comparing the mean observed evolutionary measure to the mean of the matched
 168 background distribution. We divide by the standard deviation of the evolutionary measure
 169 across the genome to standardize the enrichment. However, any summary statistic of interest
 170 could be used.

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173 **Evolutionary signals are consistent across multiple GWASs for height**

174 To evaluate the robustness of our computational framework against potential differences in
 175 GWAS size, population, study design, and analysis strategy, we compared four GWASs
 176 performed in UK Biobank individuals for standing height (Table 2): Berg-2019³⁰, Neale-2017³⁸,
 177 GIANT-2018³⁹, and Loh-2018⁴⁰. The four studies were selected to represent different
 178 methodological approaches. They were conducted in either unrelated white British individuals
 179 (Berg-2019, Neale-2017) or a more broadly defined population of European ancestry (GIANT-
 180 2018, Loh-2018). The Berg-2019 dataset is not corrected for population stratification, since they
 181 were evaluating its effects. The Neale-2017 and GIANT-2018 studies used ten genetic principal
 182 components while the Loh-2018 study used a linear mixed model (BOLT-LMM,⁴¹) shown to be
 183 robust against population stratification. The GIANT-2018 meta-analysis had the largest sample

184 size with 700K individuals whereas the other three had sample sizes of 335-460K individuals.
185 The number of independent regions based on our LD-pruning approach increased with sample
186 size except for the linear mixed model from Loh-2018 ($n=6,903$), which was the highest (Table
187 1). A Benjamini-Hochberg p-value correction (p_{adj}) was performed across 11 evolutionary
188 measures for each GWAS.

189
190 Regions associated with height were enriched for signatures of negative selection (e.g.
191 LINSIGHT, PhyloP, PhastCons) and differentiation between human populations (F_{ST}) in each of
192 the four GWASs ($p_{adj} < 0.05$, Figure 2a). Overall, nine out of the 11 evolutionary measures had
193 statistically significant deviations from the expected values (Figure 2a). These patterns relative
194 to the background distributions were consistent across all GWASs and evolutionary measures.
195 However, some measures (e.g. ARGWeaver, Beta Score, F_{ST}) showed greater variability for the
196 mean observed trait value and background distributions than others (e.g. LINSIGHT, PhyloP,
197 PhastCons). The two evolutionary measures (XP-EHH afr-eas and XP-EHH eur-eas) for which
198 the statistical significance of the deviations from the background is not maintained across all
199 four GWASs both measure population differentiation, and the two GWASs that do not show
200 significant deviations (Neale-2017 and Berg-2019) both only include white British individuals.

201
202 We also randomly sampled trait-associated regions from the Loh-2018 GWAS without
203 replacement to evaluate how evolutionary patterns varied based on the number of trait-
204 associated regions. Across measures, we found that the background distribution and trait-
205 associated value converged rapidly with an increasing number of trait-associated regions
206 (Supplementary Figure 1).

207
208 These results also demonstrate the importance of matching the background distribution to the
209 regions studied. For example, the observed Beta Scores for the Loh-2018 and GIANT-2018
210 regions are very different in magnitude (Figure 2a). Nonetheless, they are both similarly low
211 compared to their appropriate background distributions. However, if the Beta Score values for
212 GIANT-2018 had been compared to the Loh-2018 background distribution, we would have
213 come to the opposite and incorrect conclusion that they were significantly higher than expected.
214 Overall, these results suggest that our approach is robust across GWASs and not substantially
215 affected by their methodological differences.

216

217 **Some evolutionary signals vary across effect size**

218 Based on evolutionary theory and recent observations¹³, we expect stronger signatures of
219 selection at regions with higher effect sizes. Thus, we stratified the trait-associated regions from
220 the Loh-2018 GWAS into five bins with equal number of regions based on the GWAS effect size
221 at each lead SNP. We observed several trends. Evolutionary measures of negative selection
222 (LINSIGHT, PhastCons, PhyloP) had similar values and enrichment across bins (Figure 2b). In
223 contrast, measures related to local adaptation (F_{ST}), recent positive selection between human
224 populations (XP-EHH), and balancing selection (Beta Score) had the highest values in bins with
225 the smallest effect size (Figure 2b). Evolutionary enrichment was also strongest in bins with the
226 smallest magnitude for F_{ST} but generally similar across bins for XP-EHH (bar color, Figure 2b).
227 When trait-associated regions were stratified by GWAS p-value instead, we generally saw

228 similar trends with higher evolutionary measure values and enrichment for trait-associated
229 regions with the smallest p-values (Supplementary Figure 2).

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231

GWAS Name	Stratification Correction	Population	Sample Size	Independent Genomic Regions
Berg-2019	uncorrected	Unrelated white British	337K	2,505
Neale-2017	10 PCs	Unrelated white British	337K	3,598
GIANT-2018	10 PCs	European ancestry	700K	5,230
Loh-2018	Mixed effects Model	European ancestry	459K	6,903

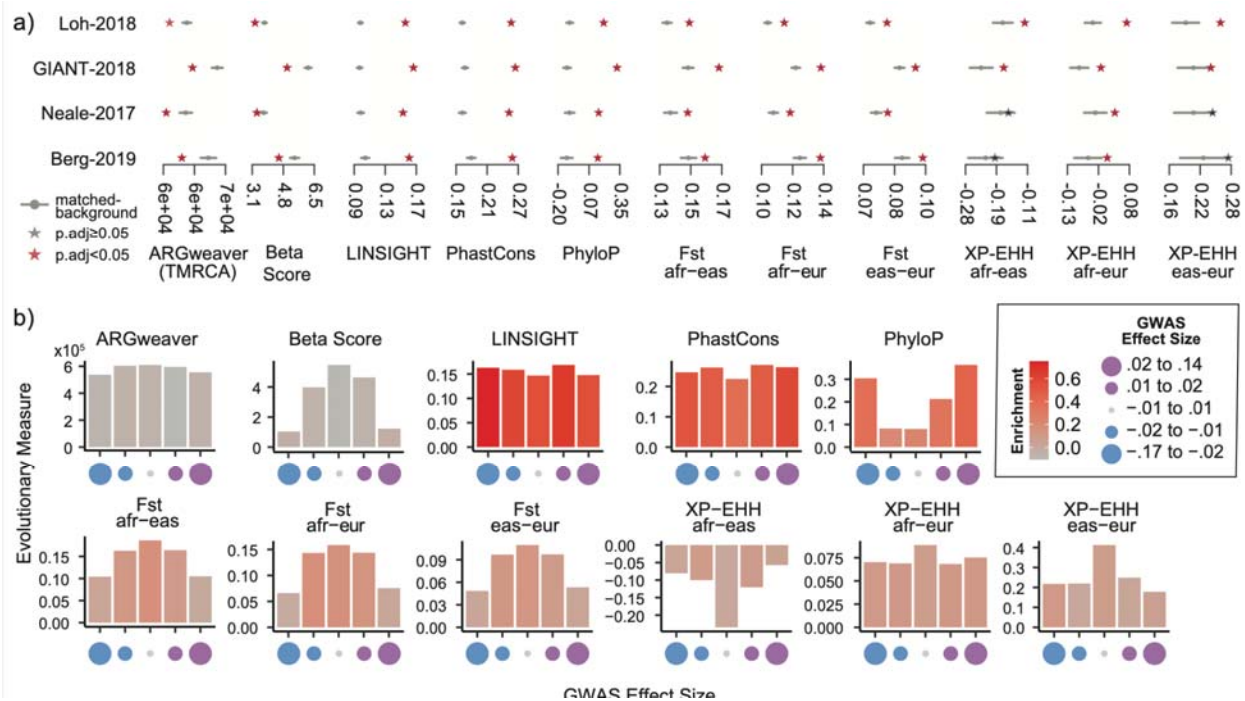
232 **Table 2: GWASs on standing height used to evaluate robustness of our approach.**

233 We used four published GWASs performed in the UK Biobank on standing height to evaluate
234 the robustness of our approach. The year in the name is when the GWAS was published. Any
235 correction for population stratification (“Stratification Correction”) and the specific GWAS
236 population (“Population”) is noted. Using the same criteria for LD-pruning (Methods), we
237 identified independent trait-associated genomic regions (“Independent Genomic Regions”). The
238 Loh-2018⁴⁰ GWAS used a linear mixed model (BOLT-LMM) shown to be robust against
239 population stratification⁴¹.

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Figure 2: The genomic signatures of evolutionary forces are consistent across multiple GWASs of the same trait.

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A mosaic of diverse evolutionary forces on regions associated with complex traits

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To generate an atlas of evolutionary forces on complex-trait-associated regions, we analyzed the GWAS summary statistics of 972 traits (Methods). Summary statistics were downloaded from diverse sources including the Neale lab UK Biobank PheWAS (n=202 traits)³⁸, the GWAS Catalog (n=312)⁴², GWAS Atlas (n=297)⁴³, manual NCBI searches, and large consortia (Psychiatric Genomics Consortium, DIAGRAM, GIANT etc.). We applied our evolutionary enrichment computational framework to each GWAS. The resulting enrichments and trait-level statistics for eleven evolutionary measures can be downloaded from FigShare repository so researchers can explore traits of interest.

269

270 The number of trait-associated regions varied widely (mean: 183, median: 9, maximum: 5,678
271 regions). In our evolutionary atlas, 888 out of 972 traits had at least one trait-associated region
272 meeting genome-wide significance (GWAS p-value < 5E-8). For traits with fewer than 50
273 associated regions, many (n=432) lacked any statistically significant evolutionary enrichments
274 (p-value<0.05 after multiple testing correction for the number of GWAS analyzed, Methods).
275 Therefore, we focus here on describing evolutionary trends for traits (n=290) with well-powered
276 GWASs with 50 or more trait-associated regions.

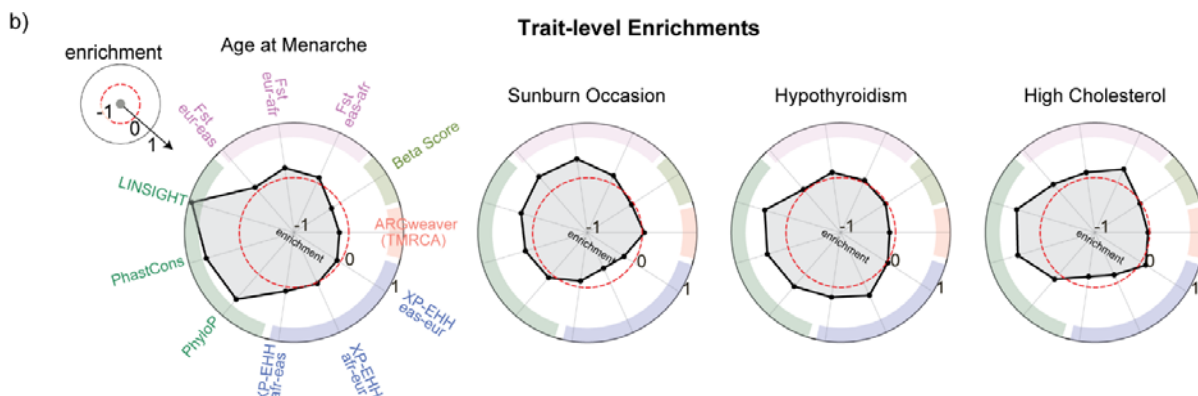
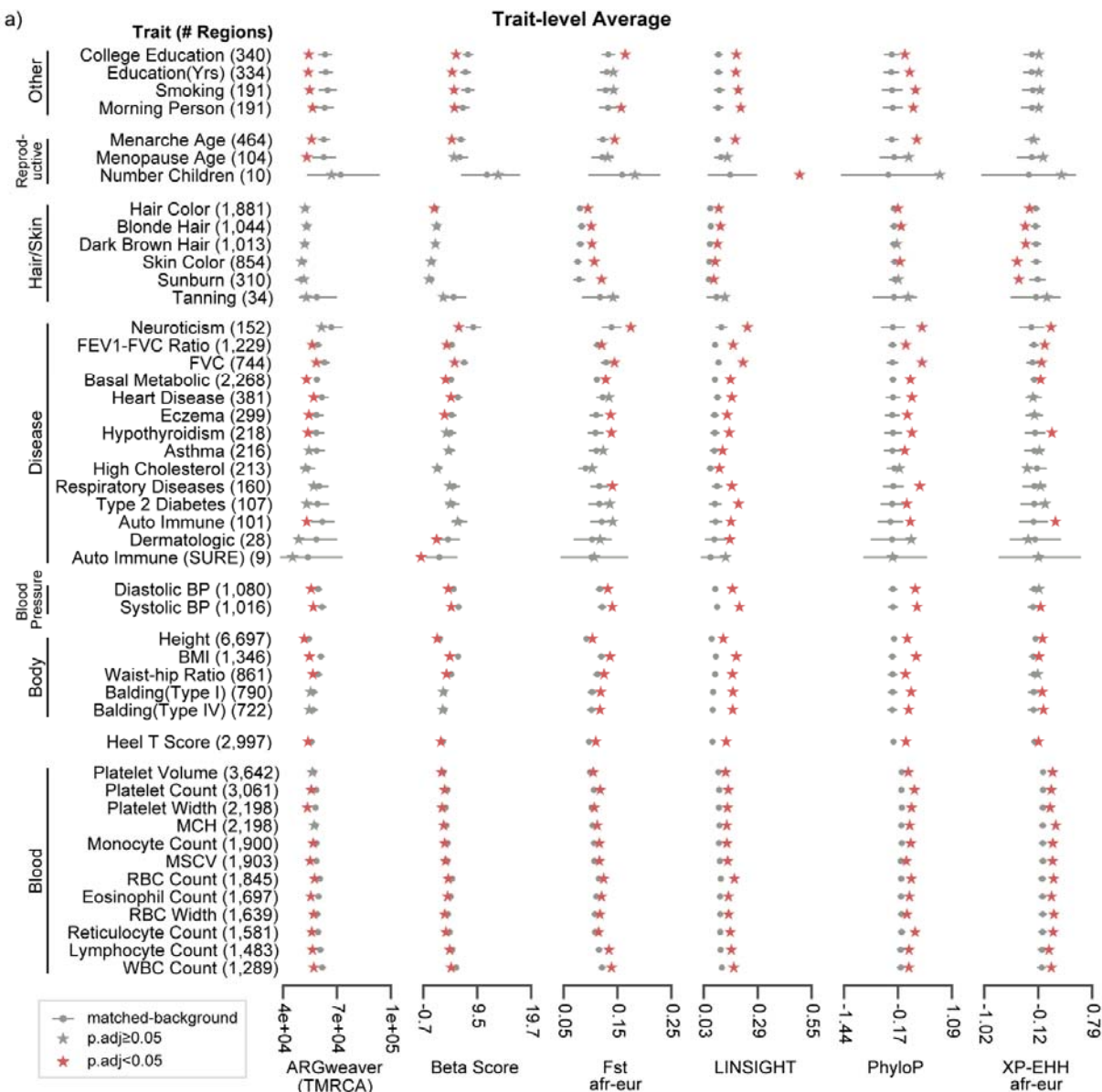
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278 For each evolutionary measure, we counted the number of GWASs with a significant deviation
279 from the background (p-value < 0.05 after multiple testing correction for the total number of
280 GWAS; Methods, Supplementary Table 1). Genomic signatures of negative selection were the
281 most prevalent: 95% of GWASs had statistically significant enrichment for PhastCons (281/290),
282 PhyloP (222/290), and LINSIGHT (278/290). We also commonly detected signals for the other
283 modes of selection. More than half of the GWASs had significant enrichment for local adaptation
284 (F_{st} , n=152 to 194 traits), negative enrichment for balancing selection (Beta Score, n=147
285 traits), and younger than expected allele ages (ARGweaver, n=166 traits). Significant genomic
286 signatures for cross-population positive selection (XP-EHH) were most prevalent for the African-
287 European comparison (n=138 traits) and less prevalent between Africans-East Asians (n=37
288 traits), and Europeans-East Asians (n=87 traits) comparisons. Though these differences may be
289 driven in part by the bias towards European-ancestry individuals in genomic studies.

290

291 To illustrate the evolutionary patterns we observed across diverse traits, we plot the results for a
292 subset of 47 GWASs carried out using the same BOLT-LMM mixed-effects model in the UK
293 Biobank (Figure 3a)⁴⁰. We refer to this analysis as the “BOLT-LMM set” (Methods). The BOLT-
294 LMM set demonstrated the same general trends across evolutionary measures as we observed
295 in the larger evolutionary atlas (Figure 3a, Supplementary Table 1). As examples of distinct
296 evolutionary profiles, we highlight four traits: Age at Menarche, Sunburn Occasion (Sunburn),
297 Hypothyroidism, and High Cholesterol (Figure 3b). Out of the four, age at menarche had the
298 strongest enrichments for negative selection measures and negative enrichment for balancing
299 selection and younger than expected allele ages. Sunburn’s evolutionary profile was
300 predominantly enriched for within human population genomic signals of positive selection (F_{st} ,
301 XP-EHH). Hypothyroidism had signatures of both negative selection and within human-
302 population positive selection (XP-EHH). Similar to age at menarche, high cholesterol had strong
303 signals of negative selection in addition to positive selection (F_{ST} , XP-EHH). Altogether, each
304 trait is characterized by distinct evolutionary profiles.

305



307 **Figure 3: Mosaic evolutionary architecture across 47 well-powered GWASs of human**
308 **complex traits.**

309 From our evolutionary atlas of 972 GWASs, we plot a subset of 47 GWASs (BOLT-LMM set)
310 performed using the same approach and from the same cohort (Methods).

311 (a) For each evolutionary measure (columns) and a given trait (row), we calculated the trait-
312 averaged value (x-axis, stars) and compared it with the matched genomic background
313 distribution (gray dots: mean values, gray bars: 5th, 95th percentiles). Traits are manually
314 grouped based on type and similarity. The number of trait-associated regions is provided in
315 parentheses. Red stars ($p_{\text{adj}} < 0.05$) represent statistically significant deviation after multiple
316 testing correction (Methods). Results are shown for six evolutionary measures; see
317 Supplementary Figure 3 for all 11 evolutionary measures.

318 (b) We calculated enrichment as described in Figure 1d and highlight four traits with distinct
319 evolutionary profiles. Spokes represent different evolutionary measures (colored by type of
320 force) and concentric rings represent levels of evolutionary enrichment. Red dashed circles
321 represent the expected values (i.e., no enrichment).

322

323

324 **Skin and hair traits show signatures of local adaptation**

325 Our analyses revealed strong signals of local adaptation for GWASs of hair and skin traits
326 (Figure 3a). In the BOLT-LMM set, the GWASs for hair color traits were highly polygenic with
327 over 1,000 trait-associated genomic regions. GWASs for skin-related traits (sunburn, tanning,
328 skin color) had variable degrees of polygenicity (34 to 854 trait-associated regions), while the
329 GWASs for the two balding traits had around 700 trait-associated regions. Except for the GWAS
330 for the tanning trait, all others demonstrated strong signatures of negative selection (LINSIGHT,
331 PhastCons, Figure 3a). They also exhibited strong genomic signatures of local adaptation (F_{ST})
332 across the three 1000 genomes superpopulations. Hair/skin color and tanning trait-associated
333 regions had signatures of recent positive selection in the European superpopulation (negative
334 XP-EHH afr- eur) compared to the African superpopulation. Meanwhile, the balding trait-
335 associated regions had evidence of recent positive selection in the African superpopulation
336 compared to the European. Similarly, evidence of recent selection between African and East
337 Asian superpopulations was observed for GWASs of dark hair and skin color. Recent selection
338 between East Asian and European super populations was observed for GWASs of hair color,
339 skin color, tanning and sunburn.

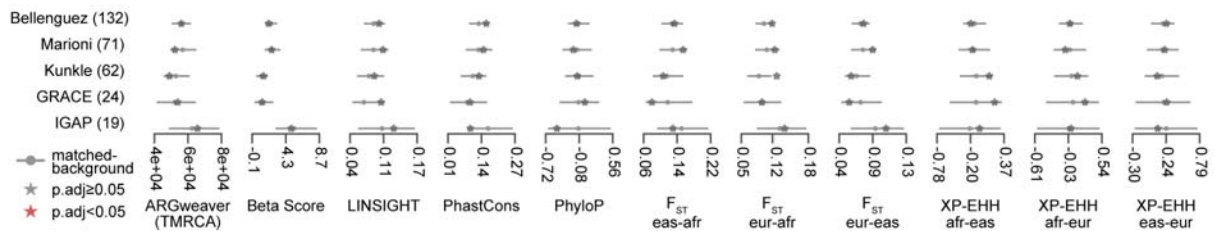
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342 **Alzheimer's disease associated genomic regions lack enrichment for selective**
343 **signatures**

344 The GWASs of nearly all traits in the BOLT-LMM set had diverse genomic signatures of
345 selection. In contrast, we observed that genomic regions associated with late-onset Alzheimer's
346 disease exhibited no significant enrichment for any evolutionary measure (Figure 4). This result
347 held across five published GWASs of late-onset Alzheimer's disease. The GWASs had between
348 19 to 132 trait-associated regions identified in European-ancestry populations: Bellenguez⁴⁴,
349 Marioni⁴⁵, Kunkle⁴⁶, GRACE⁴⁷, IGAP⁴⁸. Across the 11 evolutionary measures we tested, all
350 GWASs had trait-associated evolutionary values that overlap the expected range from their

351 matched backgrounds ($p > 0.05$, Figure 4). Consequently, we did not detect any genomic
352 signatures of enrichment across the evolutionary measures we tested. Thus, we hypothesize
353 that genomic regions associated with late-onset traits may be less likely to have strong
354 signatures of selection.
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358 **Figure 4: Loci for late-onset Alzheimer's disease lack enrichment for evolutionary forces.**

359 Across five GWASs conducted on Alzheimer's Disease (y-axis), we plot the trait-averaged value
360 (red or black stars) across evolutionary measures (x-axis) compared to their matched genomic
361 background values (gray bars, 5th, 95th percentiles). We did not find any significant enrichment
362 for any evolutionary measures ($p_{\text{adj}} < 0.05$ with multiple testing correction, Methods). This
363 pattern held across all five GWASs considered. This suggests that genomic regions contributing
364 to the development of Alzheimer's Disease are not enriched for specific evolutionary forces.
365
366

367 **Discussion**

368 Natural selection has influenced patterns of variation in genomic regions associated with many
369 human complex traits. However, the role of different modes of selection and the extent of their
370 influence on genomic regions associated with complex human traits remain challenging to
371 study. Here, we couple the availability of summary statistics from 972 GWASs with 11
372 evolutionary statistics to identify enrichment for different evolutionary forces on genomic regions
373 that contribute to variation in the human phenome. Our empirical approach quantifies
374 enrichment compared to background genomic regions matched to those identified for each trait.
375 The analysis pipeline can flexibly incorporate any evolutionary measure with genome-wide SNP
376 level annotation and quantify a trait-level summary and enrichment. We make our evolutionary
377 atlas and efficient open-source software available for the research community
378 [PLACEHOLDER_REF].
379

380 We observe several consistent trends across regions associated with diverse complex traits.
381 Signatures of negative selection, both within and between species, are enriched among variants
382 associated with nearly all complex traits. This indicates that, as expected, trait-associated
383 variation is enriched in functional regions with significant evolutionary constraint. We also
384 consistently observe significantly younger ages for trait-associated alleles, which suggests that
385 recent variants make a substantial contribution to the common-variant mediated variation in
386 most complex traits. We also observe enrichment for signatures of differentiation/positive
387 selection between populations for a substantial fraction of traits, most notably those involved in
388 hair, skin, blood measurements, and the immune system. This is consistent with recent

389 population-specific adaptation driven by these traits with particular relevance to survival in new
390 environments¹¹. Overall, regions associated with most traits show strong enrichment for
391 multiple evolutionary patterns, suggesting that a mosaic of selective pressures commonly
392 shaped variants associated with complex traits.

393
394 Our approach generalizes the common strategy of analyzing evolutionary forces on individual
395 loci of interest to comprehensively characterize all regions associated with a trait. This region-
396 focused approach has several advantages. Previous empirical work (Labella 2020,^{14,35} has
397 shown the promise of quantification of region-level pressures to understand evolutionary forces
398 on a handful of traits and interpret associated loci. Calculating a standardized enrichment for
399 each trait and measure from an appropriate background enables us to compare across different
400 evolutionary measures and, consequently, generate evolutionary profiles across GWASs of
401 different traits. Our findings are consistent with several recent genome-wide analyses that use
402 different approaches and identify widespread global differentiation³⁵, negative selection¹³, and
403 polygenic adaptation⁴⁹ on complex traits.

404
405 Differences in the average polygenic risk score between populations and the correlation
406 between polygenic risk scores and geographic clines^{27–29} or time⁴⁹ have been used to argue for
407 polygenic adaptation on traits such as height. However, such approaches can yield false
408 signatures of adaptation due to inflated differences arising from population stratification in the
409 GWASs^{30,31,50}. Our approach is distinct from and complementary to recent methods for
410 detecting polygenic selection from GWAS in several key aspects. First, it separates the
411 identification of genomic signatures of different evolutionary forces from the trait(s) that drove
412 the selection. While both are challenging problems, identifying the specific traits driving
413 selection is not necessary to infer that selection occurred in genomic regions associated with
414 these traits. Rigorous detection of polygenic adaptation would require detailed phenotypic and
415 environmental measurements over time and/or across different populations. The difficulties
416 accounting for stratification in previous studies of height illustrate these challenges. Such an
417 approach is not currently possible at scale since both modern and ancient phenotype data are
418 very sparse for most traits and many of these pressures happened deep in our evolutionary
419 history. Thus, our atlas provides a complementary high level overview of the currently
420 detectable evolutionary forces on genomic regions that underlie complex traits. We anticipate
421 that this can help generate hypotheses about which traits may have experienced different
422 selective pressures.

423
424 A second major difference is that we do not directly consider effect size or direction inferred
425 from GWAS therefore reducing the potential effect of inflated or unstable estimates between
426 populations. However, we note that effect size is indirectly taken into account in the selection of
427 genomic regions that are associated with a trait. Nonetheless, our framework enables us to
428 evaluate the relationship between effect size and evolutionary signatures of selection (Figure 2).
429 We observe for height that the most extreme scores and strongest enrichment for evolutionary
430 measures focused on differences between human populations (FST, XP-EHH) occur at lower
431 effect sizes.

432

433 Third, by summarizing the distribution of evolutionary measures at the local region and then
434 genome-wide level, we obtain a richer characterization rather than considering a single tag
435 SNP, which may be subject to substantial variation and not truly causal. Moreover, this allows
436 us to build an appropriate background distribution. This is especially important, since the
437 strength of selection is not uniform but often varies based on functional annotations across the
438 genome^{13 51}. We are also able to corroborate observations by incorporating multiple
439 evolutionary measures capturing similar evolutionary forces (e.g., PhyloP, PhastCons,
440 LINSIGHT). Finally, our framework flexibly considers many different evolutionary forces, not just
441 adaptation. We are also able to compare the enrichment for signatures of selection across traits
442 and effect sizes.

443

444 Our approach also has some limitations. First, as noted above, if the goal is to find traits under
445 selection, then identification of selection acting on genomic regions associated with a trait does
446 not necessarily imply that selection acted on the trait itself. Linking genomic signatures of
447 selection to traits is complicated by pleiotropy, especially antagonistic pleiotropy, e.g., regions
448 associated with heart disease and lifetime reproductive success exhibit antagonistic effects⁵².
449 Furthermore, the omnigenic model suggests that pleiotropy is extremely pervasive across
450 human traits⁵³; thus, attributing the contributions of selection on different genomic regions to
451 individual traits is likely to be a considerable challenge. Second, rare variants contribute to
452 variation in many complex traits⁵⁴, and our use of GWAS data limits our analyses to relatively
453 common variants. Nonetheless, our approach can be used to analyze known rare variants, and
454 increasing GWAS sample sizes are enabling the detection of effects for increasingly rare
455 variants. Finally, given the limited availability of GWAS data from non-European populations⁵⁵,
456 we have focused on trait-associated regions identified in Europeans.

457

458 The flexibility of our approach enables several future directions. As new evolutionary measures
459 are developed, they can easily be integrated into our framework. Evolutionary enrichment at the
460 trait level can be used to better understand pleiotropy and whether the enrichment varies across
461 functional regions of the human genome for a given trait. As more diverse GWASs conducted in
462 non-Europeans become available, our framework can be used to compare genomic signatures
463 of selection across human populations. This will enable additional tests for evidence of
464 polygenic adaptation, such as heterogeneity among loci and non-parallelism between replicated
465 populations⁵⁶. Additionally, our framework is not limited to the human species; the same
466 approach can be applied to GWAS conducted in any species such as mice⁵⁷, non-human
467 primates⁵⁸, or fungi⁵⁹. In summary, our quantification of genomic signatures of selection on
468 trait-associated regions advances our understanding of the genetic architecture of complex
469 traits and illuminates the diverse forces that have shaped functional regions of the human
470 genome.

471

472

473 **Methods**

474

475 **Detecting genomic signatures of evolutionary forces from summary statistics**

476 Our empirical framework to detect evolutionary signatures relies on building a matched
477 background to compare trait-associated regions. For a given trait, we identify independent trait-
478 associated regions by pruning using LD ($r^2 \geq 0.9$), genomic distance ≤ 500 kbases, and GWAS
479 p-value $< 5E-8$ (Figure 1a). This is obtained by running the --clump flag in PLINKv2 with the
480 following parameters: --clump-kb 500, --clump-r2 0.9, --clump-p1 5E-8, --clump-p2 5E-8. We
481 refer to the independent regions identified by LD clumping as trait-associated regions and
482 variants with GWAS p-value $< 5E-8$ within the clumped regions as trait-associated variants. All
483 genomic coordinates are GRCh37.

484
485 For each trait-associated region, we match using an approach motivated by SNPSNAP⁶⁰ and
486 described previously¹⁴. Briefly, for each lead variant (variant with lowest p-value) in a trait-
487 associated region, we randomly select 5,000 control variants matched on the following features:
488 allele frequency ($\pm 5\%$), LD ($r^2 > 0.9$, $\pm 10\%$ LD buddies, gene density ($\pm 500\%$) and distance
489 to nearest gene ($\pm 500\%$) (Figure 1B). We implemented the matching as a python script.
490 Matched variants were drawn from 1000 Genomes subset of the European superpopulation. To
491 match on LD patterns for each trait-associated region, we first identified the number of trait-
492 associated variants in LD ($r^2 > 0.9$) with the lead SNP. To match on LD patterns, we randomly
493 selected a variant for each trait-associated variant in LD ($r^2 > 0.9$) with the lead SNP for all trait-
494 associated regions.

495
496 Next, for every evolutionary measure, we calculated a trait-level average using two steps. First,
497 we calculate for each region (matched or trait-associated) a 'region-average' defined as the
498 greatest absolute value across all trait-associated variants. For the second step, we calculate
499 the trait-level average across all the region-averages for the trait-associated regions and each of
500 the 5,000 matched sets, where each set includes a matched region for each trait-associated
501 region (Figure 1c). The 5,000 averaged evolutionary values make up the background
502 distribution that we use to compare the trait-average evolutionary measure value (Figure 1d).
503 We derive unadjusted p-values by quantifying the number of averaged matched evolutionary
504 values as or more extreme than the trait-average out of the 5,000. We adjust this p-value for
505 multiple testing in each analysis. Additionally, using this background distribution, we define
506 evolutionary enrichment as the difference between the trait-level mean and the mean of the
507 background distribution divided by the genome-wide standard deviation of the evolutionary
508 measure (Figure 1e). This standardization allows us to compare the relative enrichment across
509 different evolutionary measures. In summary, this approach starts with GWAS summary
510 statistics and quantifies a trait-level average and enrichment for a given evolutionary measure.

511 512 **Source of evolutionary measures**

513 In this study, we downloaded or calculated eleven evolutionary measures (Table 1) for all trait-
514 associated and matched control variants as described in our previous study¹⁴. Briefly,
515 VCFTools (v0.1.14)⁶¹ was used to calculate pairwise F_{ST} , the R package rehh(v2.0) was used
516 to calculate XP-EHH using phase 3 1KG data. BetaScan software¹⁹ was used to calculate Beta
517 Score. PhyloP⁶², PhastCons 100-way⁶³, LINSIGHT⁶⁴, and Allele Age^{21,65} were downloaded
518 from their publications or the UCSC Table Browser⁶⁶.

519

520 **Evaluating robustness of evolutionary signatures using height GWAS summary** 521 **statistics**

522 GWAS summary statistics for standing height were downloaded from four different studies
523 (Table 2). The Berg-2019 analysis performed a linear regression with age, sex, and sequencing
524 array as covariates on unrelated British ancestry individuals in the UK Biobank³⁰. We
525 downloaded the summary statistics labeled “UKBB_noPCs” from
526 datadryad.org/stash/dataset/doi:10.5061/dryad.mg1rr36. The Neale-2017 analysis also
527 performed a linear regression with the first genetic 10 principal components and sex as
528 covariates on unrelated white british individuals [cite: <http://www.nealelab.is/uk-biobank/>].
529 Summary statistics were obtained by downloading the file
530 50_raw.gwas.imputed_v3.both_sexes.tsv from the “GWAS round 2” repository hosted at
531 nealelab.is/uk-biobank. The GIANT-2018 summary statistics were obtained from a meta-
532 analysis of previous height GWAS on European ancestry combined with the UK Biobank cohort
533 that included age, sex, recruitment center, genotyping batches and 10 genetic principal
534 components⁶⁷. The summary statistics were downloaded from:
535 [https://portals.broadinstitute.org/collaboration/giant/images/6/63/Meta-](https://portals.broadinstitute.org/collaboration/giant/images/6/63/Meta-analysis_Wood_et_al%2BUKBiobank_2018.txt.gz)
536 [analysis_Wood_et_al%2BUKBiobank_2018.txt.gz](https://portals.broadinstitute.org/collaboration/giant/images/6/63/Meta-analysis_Wood_et_al%2BUKBiobank_2018.txt.gz). The Loh-2018 analysis used a linear mixed
537 model on individuals of European ancestry from the UKBiobank⁴⁰. The height summary
538 statistics were downloaded from <https://alkesgroup.broadinstitute.org/UKBB/> (file name:
539 [body_HEIGHTz.sumstats.gz](https://alkesgroup.broadinstitute.org/UKBB/)).

540

541 On all four summary statistics, we applied our approach to detect genomic signatures of
542 evolutionary forces. We calculated a trait-associated region average and the distribution of the
543 background set and the evolutionary enrichment as described earlier (Figure 1). For each
544 summary statistic, we corrected for multiple testing across the 11 evolutionary measures using
545 the Benjamini-Hochberg FDR control approach.

546

547 To test for effects of trait-associated p-value obtained from the summary statistics, we created
548 quintiles with an equal number of trait-associated regions based on the GWAS summary
549 statistics p-value at the lead SNP. We then applied our evolutionary analysis on each quintile.
550 We repeated the same steps to test for the effect size from the GWAS summary statistics but
551 instead created quintiles based on the beta coefficient. To test how the number of trait-
552 associated regions affected our evolutionary analyses, we randomly sampled with replacement
553 the number of trait-associated regions to create under-sampled sets. Then for each set, we ran
554 our evolutionary pipeline to calculate a trait-level average (Supplementary Figure 1).

555

556 **GWAS datasets to generate evolutionary atlas**

557 We used multiple sources to identify GWASs that were conducted in individuals of European
558 Ancestry and had complete publically available summary statistics for all analyzed regions
559 reported in human genome version hg19. GWASs that reported only the top hits were excluded.
560 None of the sources required substantial authorization or approval and could be downloaded
561 either from a web browser or via file transfer. Sources for GWASs include, but are not limited to
562 the Neale Lab analysis of the UK-Biobank data, the GWAS Catalog, the GWAS Atlas, NCBI

563 searches, and major GWAS consortia such as the Psychiatric Genomics Consortium and
564 DIAGRAM. All of the analyzed GWASs are reported in Supplemental File 1. This includes, when
565 available, the associated PMID, and link to download the raw GWAS summary statistics.

566
567 For each summary statistic, we applied our approach to detect genomic signatures of
568 evolutionary forces as described earlier. GWASs without any significant independent regions
569 (based on p-value and LD as described above) were not further analyzed. For all GWAS with at
570 least one associated region we retained the summary statistics for every individual trait-
571 associated genomic region and the trait-level enrichment across the entire GWAS. To correct
572 for multiple testing, empirical p-values across all traits for a given evolutionary measure were
573 adjusted using the Benjamini-Hochberg FDR control approach.

574 This data is available on FigShare reports empirical p-value only and should be adjusted
575 accordingly for future analyses (link to be provided upon publication).

576

577 **BOLT-LMM GWASs subset analysis**

578 We further analyzed a subset of 47 traits, which we refer to as the “BOLT-LMM set”, whose
579 summary statistics were generated using a mixed modeling approach⁴⁰. All summary statistics
580 were downloaded from <https://alkesgroup.broadinstitute.org/UKBB/>. We ran our evolutionary
581 analyses to calculate trait-level averages and the background distribution (Figure 3a). Empirical
582 p-values were corrected for multiple testing across traits and evolutionary measures using the
583 Benjamini-Hochberg FDR control method. Next we calculated the evolutionary enrichment for
584 each trait and evolutionary measure.

585

586 **Late-onset Alzheimer’s disease analyses**

587 We performed our evolutionary analysis on five GWAS of the late onset Alzheimer’s trait. The
588 GWAS analyzed were collected from the following sources: Bellenguez et al.
589 (<https://doi.org/10.1038/s41588-022-01024-z>), Marioni et al. (doi:10.1038/s41398-018-0150-6),
590 Kunkle et al. (doi:10.1038/s41588-019-0495-7), GRACE
591 (<https://doi.org/10.1016/j.jalz.2019.06.4950>), IGAP (doi: 10.1038/ng.2802). The most recent
592 GWAS (Bellenguez et al.) was reported in hg38 and converted to hg19 using the Biomart
593 function in R using the archived Ensembl 75: Feb 2014 (GRCh37.p13). Empirical p-values were
594 corrected for multiple testing across all five GWAS and 11 evolutionary measures using the
595 Benjamini-Hochberg FDR control approach.

596

597

598

599 **Data Availability**

600 We have made both the formatted input files and the final output files (both trait and region level
601 results) available for download on our FigShare (link to be provided upon publication). The
602 FigShare repository contains one compressed folder per PubMed ID which contains all the
603 associated input and output files.

604

605

606 **Code Availability**

607 Evolutionary calculations were performed using the GSEL python package (link to be provided
608 upon publication). Scripts with necessary data to replicate manuscripts will be provided upon
609 publication.

610

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620

621 **Contributions**

622 A.A., A.L.L., J.A.C., A.R. conceived and designed the study. A.A. and A.L.L. performed all data
623 curation, analyses, and visualizations. A.A. and A.L.L wrote the original draft and revisions for
624 the manuscript under guidance from J.A.C. and A.R. All authors reviewed and approved the
625 final manuscript.

626

627 **Competing interests**

628 The authors have no competing interests to declare.

629 Supplemental Tables

Evolutionary Atlas		
Annotation	# Traits	Proportion of All Traits (%)
ARGweaver	166	57.2
Beta Score	147	50.7
LINSIGHT	278	95.9
PhastCons	281	96.9
PhyloP	222	76.6
F _{ST} afr-eas	152	52.4
F _{ST} afr-eur	194	66.9
F _{ST} eas-eur	175	60.3
XP-EHH afr-eas	37	12.8
XP-EHH afr-eur	138	47.6
XP-EHH eas-eur	87	30

BOLT-LMM Set				
Annotation	# Traits	Proportion of All Traits (%)	Enrichment (# Traits)	Depletion (# Traits)
ARGweaver	29	61.7	0	30
Beta Score	32	68.1	0	32
LINSIGHT	44	93.6	45	0
PhastCons	47	100	47	0
PhyloP	39	83	40	0
F _{ST} afr-eas	31	66	32	0
F _{ST} afr-eur	35	74.5	37	0
F _{ST} eas-eur	35	74.5	35	0
XP-EHH afr-eas	15	31.9	5	2
XP-EHH afr-eur	29	61.7	24	5

XP-EHH eas-				
eur	19	40.4	13	4

630 **Supplementary Table 1: Count of traits with signals of evolutionary forces.**

631 Number of traits (“# Traits”) in the full Evolutionary Atlas (top) and BOLT-LMM subset (bottom)
632 with statistically significant enrichment for evolutionary measures (rows). Note, only traits with
633 50 or more associated regions are analyzed within the Evolutionary Atlas. The proportion out of
634 all traits analyzed (“Proportion of All Traits (%)”) are shown for the Evolutionary Atlas (n=290
635 traits) and BOLT-LMM set (n=47 traits). Depletion refers to negative enrichment.

636

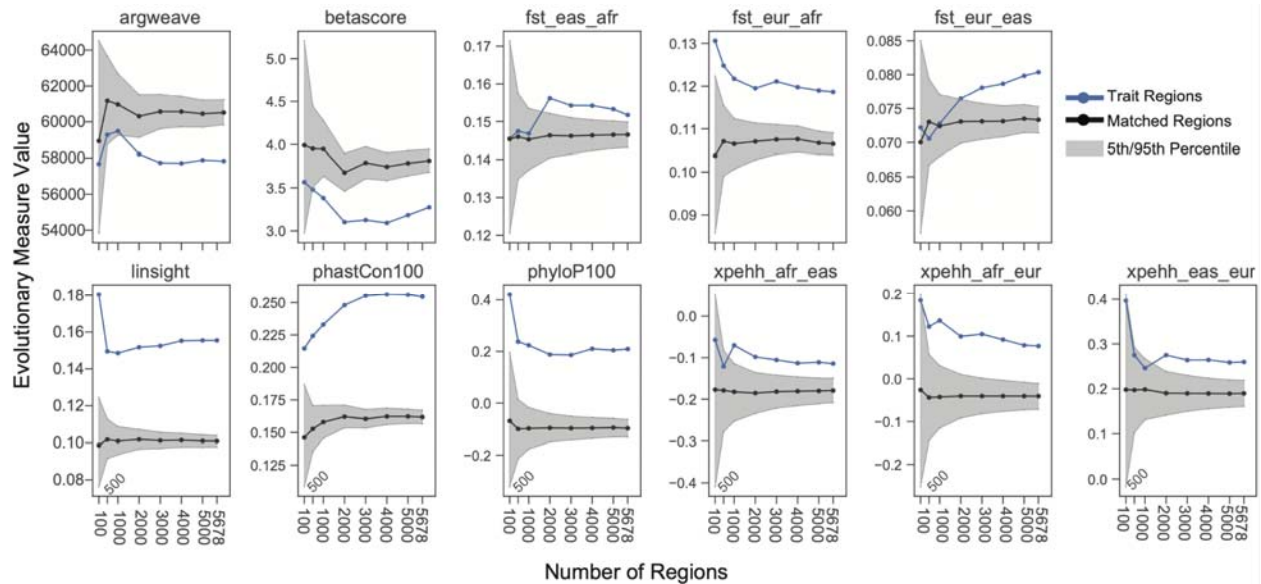
637

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639

640 **Supplemental Figures**

641



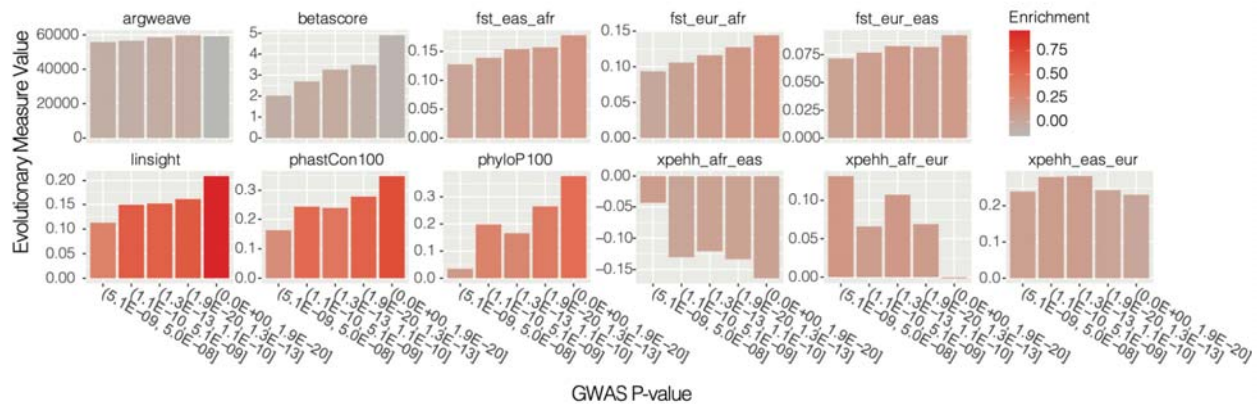
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644 **Supplemental Figure 1: Evolutionary signatures converge rapidly with increasing number**
645 **of trait-associated genomic regions.**

646 Using the Loh et. al. GWAS, we randomly undersampled the number of trait-associated regions
647 without replacement (x-axis) and measured the mean evolutionary measure at trait-associated
648 regions (blue line) and the matched background (mean: black line, gray shading between 5th
649 and 95th percentiles). The observed evolutionary measures for trait-associated regions and
650 their relative values compared to the matched background regions are consistent across
651 different numbers of associated loci considered.

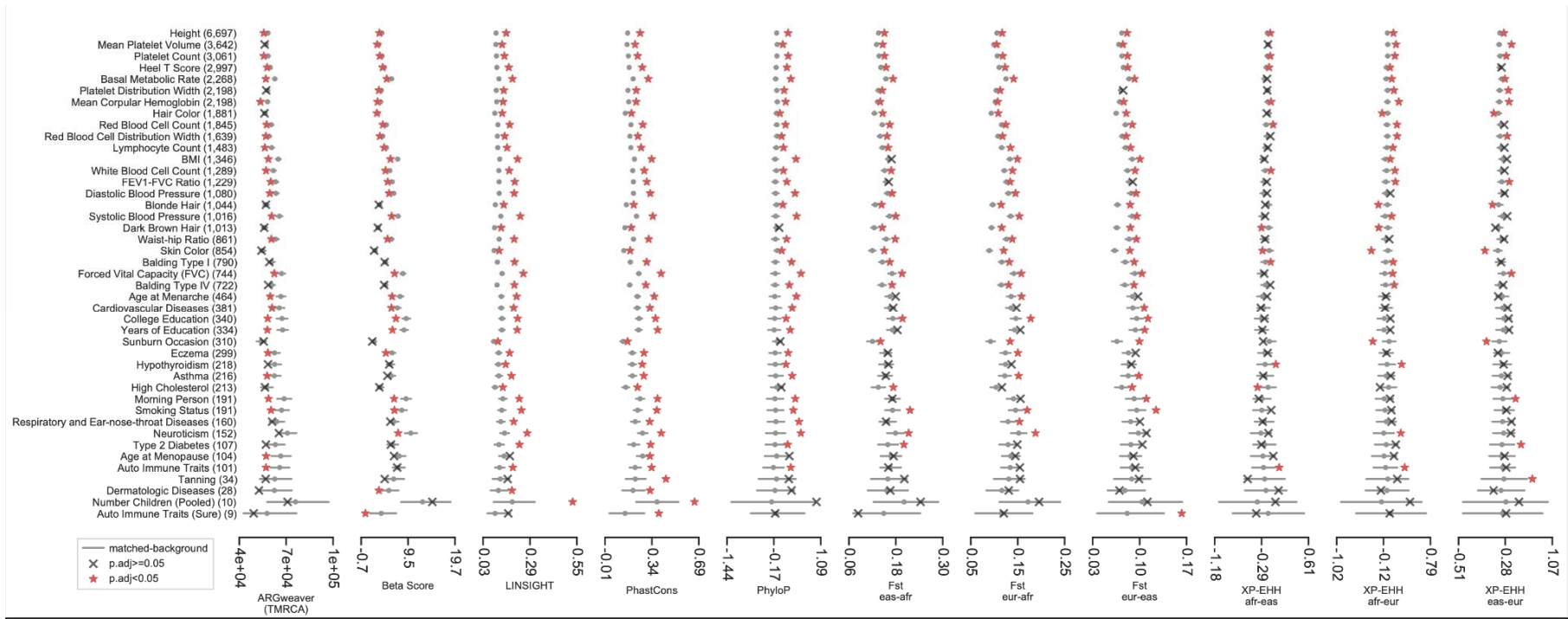
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656 **Supplemental Figure 2: Strongest genomic evolutionary signatures occur in most**
657 **significant trait-associated regions.**

658 Using the Loh-2018 (Figure2) GWAS, we partitioned trait-associated regions into five bins with
659 equal number of regions based on GWAS p-value of the lead SNP in each region. Each plot
660 represents the mean trait value (y-axis) for an evolutionary measure and each bar is colored by
661 the evolutionary enrichment which is calculated as described in Figure 1d.
662



Supplemental Figure 3: Mosaic evolutionary architecture across 47-well-powered GWASs across 11 evolutionary measures.

On a subset of 47 GWASs (y-axis, BOLT-LMM set), the trait-level average (red star or gray 'x') for 11 evolutionary measures (x-axis) compared to its matched background distribution (gray dots: mean values, gray bars: 5th, 95th percentiles) are displayed. The number of trait-associated regions is provided in parentheses. Red stars (p.adj<0.05) represent statistically significant deviation after multiple testing correction (Methods). This figure extends Figure 3a by including all 11 evolutionary measures considered in this study.

Supplemental Files

Supplemental File 1: GWAS_SOURCE_TABLE.xlsx - This excel file contains PMID or web link and the source for each GWAS summary statistics analyzed in this study.

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