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Regulation of protein abundance in genetically diverse mouse populations

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Abstract

Proteins constitute much of the structure and functional machinery of cells, forming signaling networks, metabolic pathways, and large multi-component complexes. Protein abundance is regulated at multiple levels spanning transcription, translation, recycling, and degradation to maintain proper balance and optimal function. To better understand how protein abundances are maintained across varying genetic backgrounds, we analyzed liver proteomes of three genetically diverse mouse populations. We observe strong concordance of genetic and sex effects across populations. Differences between the populations arise from the contributions of additive, dominance, and epistatic components of heritable variation. We find that the influence of genetic variation on proteins that form complexes relates to their co-abundance. We identify effects on protein abundance from mutations that arose and became fixed during breeding and can lead to unique regulatory responses and disease states. Genetically diverse mouse populations provide powerful tools for understanding proteome regulation and its relationship to whole-organism phenotypes.

Key words

proteomics, protein complex, systems genetics, pQTL, Collaborative Cross, Diversity Outbred, inbred strains

42 INTRODUCTION

43 Protein abundance is regulated at multiple levels spanning transcription, translation, recycling,
44 and degradation. It is responsive to genetic variation, as observed in yeast (Picotti et al., 2013),
45 mice (Chick et al., 2016; Williams et al., 2016; Wu et al., 2014), and human cell lines, tissues,
46 and populations (Battle et al., 2015; He et al., 2020; Liu et al., 2015; Suhre et al., 2020). Genetic
47 effects on protein abundance can be broadly divided into two classes. Local variants occur in
48 the vicinity of the coding gene and commonly affect protein abundance by altering the rate of
49 transcription or stability of the transcript (Pai et al., 2012). In contrast, distal genetic variants
50 are found at loci far from the coding gene and their effects on protein abundance are often
51 post-translational and conferred via a diffusible intermediate; often another protein. Within
52 these categories, multiple modes of regulation are possible (Chick et al., 2016). The complexity
53 of genetic regulation is compounded for proteins that are members of complexes where
54 stoichiometry imposes varying degrees of constraint (Chick et al., 2016; Huttlin et al., 2020;
55 Romanov et al., 2019; Szklarczyk et al., 2019; Taggart et al., 2020). The number of genetic loci
56 that affect a protein can range from a single locus (monogenic) to many (multi-genic to
57 polygenic). Their individual effects can be additive or dominant/recessive, thus dependent on
58 the extent of heterozygosity in the population. Loci can also interact, resulting in epistatic
59 effects.

60 Resource populations with high levels of genetic diversity can help to identify and characterize
61 the genetic loci that affect protein abundance. The Collaborative Cross (CC) (Churchill et al.,
62 2004; Collaborative Cross Consortium, 2012) and Diversity Outbred (DO) (Churchill et al., 2012)
63 mouse populations are powerful tools for genetic analysis. They are descended from a common
64 set of eight inbred strains (*i.e.*, the founder strains), representing three subspecies of the house
65 mouse, *Mus musculus* (Yang et al., 2007, 2011) and encompass genetic variation from across
66 laboratory and wild mice. DO mice are available in large numbers and each individual possesses
67 a unique, highly recombinant and heterozygous genome, supporting powerful, fine resolution
68 mapping of genetic variants (*e.g.*, Svenson *et al.* 2012; French *et al.* 2015; Keller *et al.* 2018,
69 2019). The CC consists of ~60-70 strains that are largely inbred, with many strains homozygous
70 at most loci (>99%), and residual heterozygous regions known and characterized (Collaborative
71 Cross Consortium, 2012; Shorter et al., 2019; Srivastava et al., 2017). Due to fewer outbreeding
72 generations than in the DO, the CC possess larger linkage disequilibrium (LD) blocks. The
73 reproducible genomes of CC strains enable replicate study designs (Mosedale *et al.* 2017, 2019)
74 and the characterization of genetic effects on strain-specific phenotypes (Philip *et al.* 2011;
75 McMullan *et al.* 2016). CC strains can serve as models for human diseases including colitis
76 (Rogala et al., 2014), susceptibility to Ebola infection (Rasmussen et al., 2014), influenza A virus
77 (Noll et al., 2020), SARS-coronavirus (Gralinski et al., 2015), and peanut allergy (Orgel et al.,
78 2019).

79 In this work, we obtained multiplexed mass spectrometry (mass-spec) quantification of proteins
80 in liver samples of 116 CC mice representing female/male pairs from 58 strains. We previously
81 collected proteomics data from the livers of 192 DO mice and 32 mice representing the eight
82 founder strains (two animals of each sex per founder strain) (Chick et al., 2016) (**Figure 1a**). We
83 map protein quantitative trait loci (pQTL) in both the CC and DO and find consistency of the
84 genetic and sex effects on protein abundance across the populations. We characterize the
85 genetic architecture of protein-complexes in both the CC and DO, and find that complexes with
86 highly co-abundant members are heritable and exhibit non-additive and/or highly polygenic
87 genetic architecture, which is more apparent in the CC. Finally, we identify genetic variants that
88 originated and became fixed during the of the CC and contribute to CC strain-specific
89 phenotypes. Our work demonstrates the complementary strengths of these genetic resources
90 and offers new insights into the genetic regulation of protein abundance.

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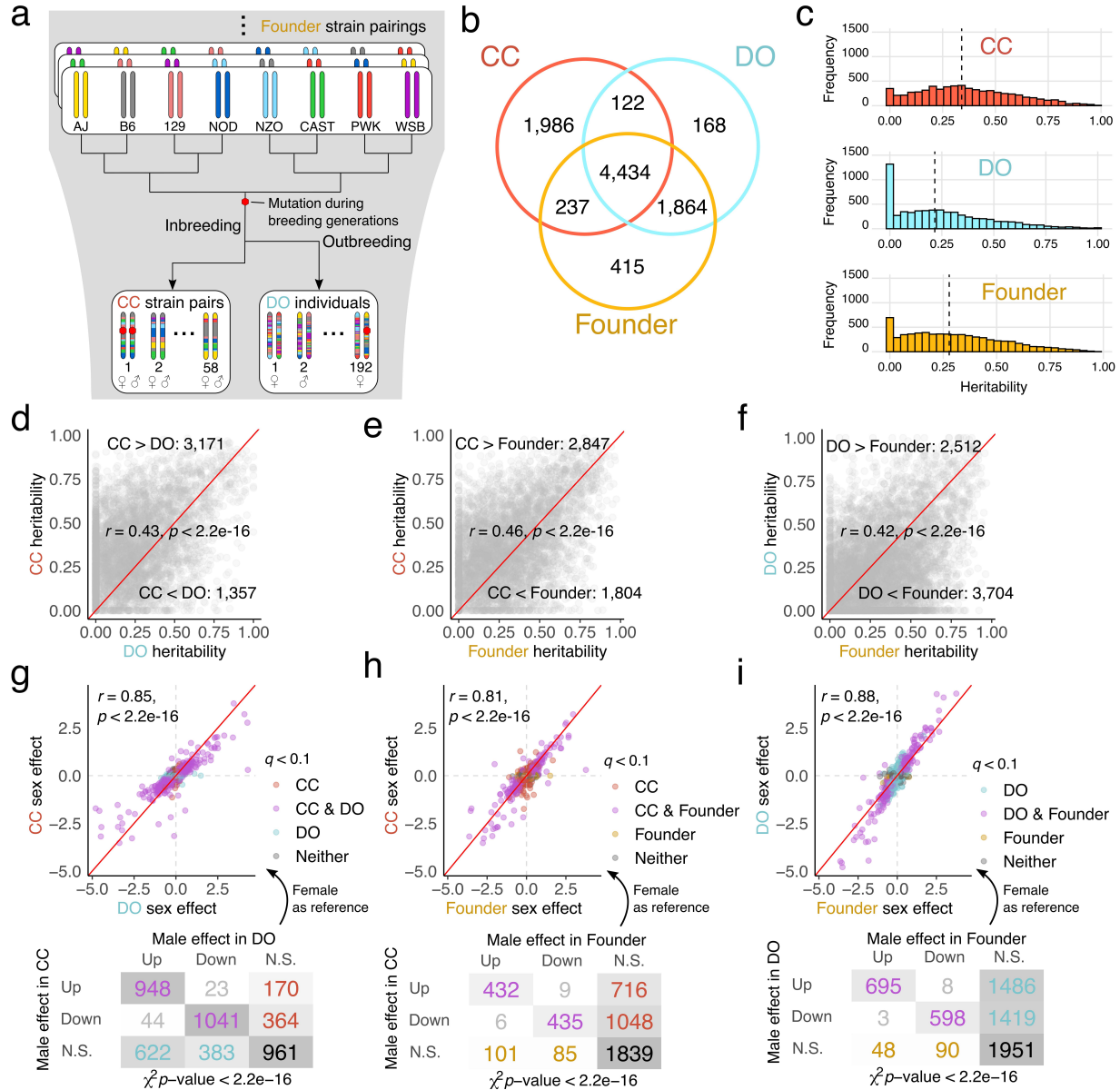
92 **RESULTS**

93 **Genetic effects on proteins are shared across the CC, DO, and founder strains**

94 We quantified the abundance of 6,779 proteins in liver tissue from 58 inbred CC strains, one
95 female and male per strain. We previously reported quantification of proteins from liver tissue
96 of 192 outbred DO mice and 32 mice representing the eight founder strains (two per sex per
97 strain) (Chick et al., 2016). The data for DO and founder strains were re-analyzed for this study
98 to ensure that all data were processed consistently (**Methods**), resulting in the quantification of
99 6,588 and 6,950 proteins, respectively. From the 9,226 total proteins detected, 4,434 were
100 observed in all three populations (**Figure 1b**; **Table S1**).

101 We estimated protein abundance heritability (h^2), which reflects the combined effects of
102 genetic variants, their allele frequencies, and the large-scale genetic background (*e.g.*, inbred or
103 outbred) of each population (**Figure 1c**). Heritability estimates were highest in the CC and
104 founder strains, due to capturing variation from non-additive genetic effects (*i.e.*, recessive and
105 epistatic) in inbred strains. Heritability estimates in the DO are limited to additive genetic
106 factors (narrow sense heritability; Lynch and Walsh 1998) that are captured in the genetic
107 relatedness of our sample. Heritability estimates were significantly correlated across
108 populations ($r > 0.4$, $p < 2.2e-16$), suggesting that much of the controlling genetic variation is
109 conserved across populations.

110 In order to identify the genetic loci that drive variation in protein abundance, we carried out
111 protein quantitative trait locus (pQTL) mapping in the CC and DO (**Table S2**). To determine
112 significant pQTL, we first applied a permutation analysis (Doerge and Churchill, 1996) to control
113 genome-wide error rate for each protein and then applied a false discovery rate adjustment
114 (FDR < 0.1) across proteins (Chesler et al., 2005) to establish a stringent detection threshold for
115 pQTL. Using this criterion, we identified 1,199 local and 289 distal pQTL in the CC and 1,652



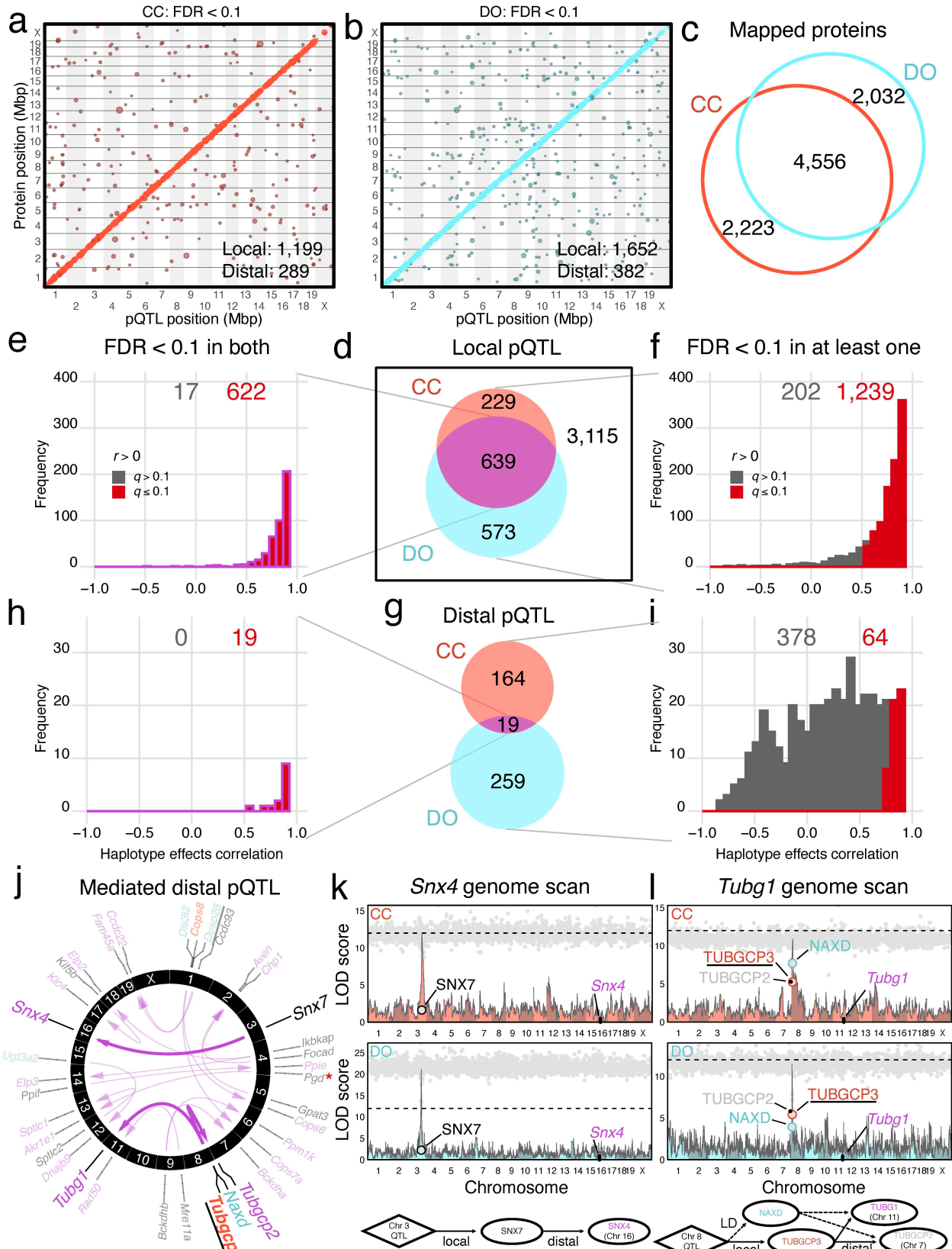
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118 **Figure 1. Comparisons of genetic and sex effects on protein abundance among the CC, DO, and founder strains reveal strong**
119 **concordance.** (a) The CC strains and DO mice are descended from the same eight inbred founder strains. Mutations occur
120 during the breeding generations of the CC and DO and can become fixed in the CC, allowing their effects to be characterized. (b)
121 Venn diagram of the analyzed proteins in the CC, DO, and founder strains. The founder strains and DO samples were from a
122 joint experiment with the same protein annotations, resulting in greater overlap. (c) Overall genetic regulation, measured as
123 heritability, is greater in the inbred CC and founder strains, reflecting contributions from non-additive genetic effects. Dashed
124 vertical lines represent the median heritability in each population. Comparisons of the heritability of individual protein
125 abundance for (d) CC and DO, (e) CC and founder strains, and (f) DO and founder strains. The correlation in heritability is high
126 and significant regardless of which populations are being compared ($r > 0.42$, $p < 2.2e-16$). Heritability estimates in the CC are
127 higher in part due to inclusion of a bridge sample across the batches of mass-spec experiments, which improved quantification
128 accuracy (**Methods**). Red diagonal lines included for reference. Comparisons of the sex effects (with female as the reference) on
129 individual protein abundance for (g) CC and DO, (h) CC and founder strains, and (i) DO and founder strains. N.S. indicates
130 proteins that did not have significant sex effects. Sex effects are even more strongly correlated than heritability ($r > 0.81$, $p <$
131 $2.2e-16$). Red diagonal lines included for reference. A breakdown of the number of proteins with significant sex effects and their
direction is shown for each comparison of populations.

132 local and 382 distal pQTL in the DO (**Figure 2a & b**), where local is defined as when the pQTL is
133 located within 10 Mbp of the midpoint of the protein-coding gene. We also identified a local
134 pQTL on the mitochondrial genome in the CC for *mt-Nd1* (**Figure S1d**). In order to prevent the
135 exclusion of true pQTL due to overly stringent control of false positive and accurately compare
136 pQTL discovery across populations, we carried out a parallel analysis with more lenient FDR
137 control (FDR < 0.5; **Figure S1a & b**).

138 We next restricted the set of proteins to the 4,556 that were detected and analyzed in both
139 populations in order to compare genetic effects between the CC and DO. (**Figure 2c**). Among
140 1,441 local pQTL stringently detected for the shared proteins, 639 were detected in both
141 populations (**Figure 2d**). To determine if these local pQTL were driven by the same genetic
142 variants, we compared the estimated haplotype effects at each pQTL (**Methods**) and found that
143 622 (97.3%) were significantly positively correlated (**Figure 2e; Table S3**). To assess whether
144 pQTL detected in only one population are population-specific or fell below the detection
145 threshold in the other population but are likely driven by the same genetic variation, we
146 compared the haplotype effects of detected pQTL to effects estimated at the corresponding
147 locus in the other population, finding that 1,239 (86.0%) were significantly positively correlated.
148 This trend for local pQTL holds true for lenient detection (**Figure S1f & g; Table S3**), indicating
149 that local genetic effects on proteins are highly conserved between the CC and DO populations,
150 even when failing to pass the threshold of detection in one of the populations.

151 The founder strains provide additional support for local pQTL in the CC and DO (**Figure S1k-n**),
152 particularly for those that are challenging to detect due to rare alleles in the CC or DO (*e.g.*, a
153 founder allele observed in three or fewer CC strains). We selected all genes with rare local
154 founder alleles that did not have a leniently detected pQTL in the CC, representing 2,410 genes.
155 We correlated the haplotype effects estimated at the locus closest to the gene transcription
156 start sites (TSS) with the protein abundance in the founders (**Methods**) and found significant
157 positive correlation for 196 genes, such as *Cyp2j5*. The conservation of local genetic effects
158 among the founder strains, CC, and DO can be striking (such as in *Gosr2* and *Cyp2d22*; **Figure**
159 **S2**). The three populations together identify a more complete catalog of local pQTL than any
160 one alone. In total we find evidence to support local genetic effects on abundance for 3,004
161 proteins observed across the CC and DO.

162 Of the 183 distal pQTL stringently detected in the CC and 278 in the DO for the shared set of
163 proteins, we found 19 that were stringently detected in both populations. All 19 also had
164 significantly correlated haplotype effects (**Figure 2h; Table S3**). Overall, the distal pQTL are
165 weaker than the local pQTL (*e.g.*, Chick *et al.* 2016; Albert *et al.* 2018) which may contribute to
166 an increased rate of false negative findings. By comparing haplotype effects for pQTL that were
167 only stringently detected in only one population, we identified an additional 45 shared distal
168 pQTL between the CC and DO populations (**Figure 2i; Table S3**).



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Figure 2. Genetic effects of loci are highly consistent between the CC and DO. Stringently detected pQTL (FDR < 0.1) in the (a) CC and (b) DO. The pQTL are plotted by the genomic positions of proteins against pQTL coordinates. Dot size is proportional to

172 strength of association (LOD score). (c) Venn diagram of overlap of analyzed proteins for the CC and DO. (d) Venn diagram of
173 overlap of local pQTL detected in the CC and DO. (e) The haplotype effects of local pQTL detected in both populations are highly
174 consistent, as measured by the correlation coefficient comparing the effects in the CC and DO. (f) More local pQTL have
175 consistent effects between the populations when also considering pQTL detected in only one of them. Red bars represent the
176 number of pQTL that had significantly correlated effects ($FDR < 0.1$). (g) Venn diagram of overlap of distal pQTL detected in the
177 CC and DO. (h) All 19 distal pQTL detected in both populations have consistent haplotype effects. (i) Considering distal pQTL
178 detected in either of the populations resulted in 64 with correlated effects in the other population. (j) Circos plot of the 19
179 distal pQTL detected in both CC and DO. Arrows connect candidate drivers identified through mediation analysis to their targets
180 (proteins with distal pQTL). Gene names in black represent the top candidates identified in both the CC and DO, whereas red
181 and blue gene names were specific to the CC or DO, respectively. For candidate mediators that do not match between the CC
182 and DO, the stronger candidate based on shared membership in protein families is underlined. The red asterisk denotes *Pgd* as
183 a likely false positive protein mediator, observed in both the CC and DO. Examples of mediation agreement (*Snx4*) and
184 disagreement (*Tubg1*) between the CC and DO are highlighted, with overlaid pQTL and mediation scans. Gray dots represent
185 mediation scores for individual proteins. A strong mediator is local to the pQTL and has a steep drop from the peak LOD score.
186 (k) *SNX7* was detected as the driver of the *Snx4* distal pQTL in both the CC and DO. (l) *TUBGCP3* was identified as the best
187 candidate driver for the *Tubg1* distal pQTL in the CC, and *NAXD* as the third-best mediator; however, in the DO, *NAXD* was the
188 stronger mediator. *Tubgcp3*, as well as *Tubg1* and *Tubgcp2*, are members of the tubulin superfamily, providing further support
189 for *TUBGCP3* as the stronger candidate driver of the distal pQTL of *Tubg1* and *Tubgcp2*. Horizontal dashed lines at LOD of 12
190 included for reference. Lenient mapping results ($FDR < 0.5$) further support the consistency of genetic regulation between the
191 CC and DO (**Figure S1**).

192 The genetic variants that drive distal pQTL are generally thought to act through diffusible
193 intermediates, one or more of which are under local genetic control by the same variants. We
194 used mediation analysis (Baron and Kenny 1986; MacKinnon *et al.* 2007), adapted for the DO
195 (Chick *et al.* 2016; Keller *et al.* 2018) and CC (Keele *et al.*, 2020), to identify candidate drivers of
196 distal pQTL (**Methods; Table S4**). Mediation analysis can provide support for one or more
197 candidate mediators for a pQTL. Furthermore, alternative candidates cannot be ruled out if
198 they unobserved, such as proteins that failed to be detected by mass-spec or functional non-
199 coding RNA. We identified candidate mediators for each of the 19 shared distal pQTL (**Figure**
200 **2j**). The same mediator was identified in the CC and DO for 14 of these – we note that *BCKDHB*,
201 *CCDC93*, and *IKBKAP* are each candidate mediators of the distal pQTL for two proteins (*CCDC93*
202 for *Ccdc22* and *Fam45a*, *BCKDHB* for *Bckdha* and *Ppm1k* for, and *IKBKAP* for *Elp2* and *Elp3*). For
203 four of the 19 distal pQTL, the best candidate mediator was different for the CC and DO. For
204 example, *TUBGCP3* is the strongest mediator of distal pQTL of *Tubg1* and *Tubgcp2* in the CC but
205 *NAXD* is the strongest mediator in the DO (**Figure 2l**). Given that *Tubg1*, *Tubgcp2*, and *Tubgcp3*
206 are all members of the tubulin superfamily, *TUBGCP3* is a stronger candidate based on
207 functional overlap. A true mediator should possess a local pQTL that co-localizes with the distal
208 pQTL. Notably, the local pQTL for *Naxd* is much stronger in comparison to the local pQTL of
209 *Tubgcp3* in the DO ($LOD_{Naxd} = 40.2$ compared to $LOD_{Tubgcp3} = 11.0$) whereas in the CC, they are
210 more comparable ($LOD_{Naxd} = 9.7$ compared to $LOD_{Tubgcp3} = 7.9$). The local pQTL of *Naxd* in the
211 DO may be acting as a surrogate for the local genotype, and thus outperforming *TUBGCP3*.
212 Similarly, *COPS8* is the strongest mediator for distal pQTL of *Cops6* and *Cops7a* in the CC,
213 whereas in the DO, *COPS8* was not detected as a mediator and notably did not have a local
214 pQTL (even leniently detected) in the DO, suggesting that *COPS8* may be less accurately
215 measured in our DO sample population. The final case of discordance was for the distal pQTL of

216 *Dnajb9*, for which UGT3A2 was identified as a mediator in the DO whereas no candidate was
217 found in the CC. *Ugt3a2* possesses local pQTL in both the CC and DO, which is notably strong in
218 the DO ($LOD_{Ugt3a2} = 36.9$), suggesting it may be a false positive mediator in the DO and the true
219 mediator was unobserved for both populations. We considered all distal pQTL that were
220 stringently detected ($FDR < 0.1$) in one of the populations, and evaluated the corresponding
221 pQTL status (stringent, lenient, or not detected) and mediation status (*e.g.*, same or different
222 mediator) in the other population (**Figure S1o**), which revealed similar levels of concordance in
223 mediation for pQTL detected in only one of the populations.

224 When we expand our pQTL comparisons to include all leniently detected pQTL, 1,462 proteins
225 have significantly correlated local pQTL (223 more than with stringent detection in one
226 population; **Figure S1g**). In contrast, fewer distal pQTL with significantly correlated effects are
227 detected (22 compared to 64) (**Figure S1j**). Some pQTL that fail to meet the stringent threshold
228 in either population still replicate across the CC and DO, as is the case for 12 of the 41 proteins
229 with distal pQTL and correlated haplotype effects (**Figure S1i**). One interesting example is *Erc3*
230 (**Figure S3**), which has distal pQTL near a region on chromosome 7 that contains *Gtf2h1* and
231 *Erc2*, which all strongly associate with each other based on protein-protein interactions
232 (Bioplex; Huttlin *et al.* 2020) and exhibit consistent but more extreme pQTL effects in the CC
233 compared to the DO.

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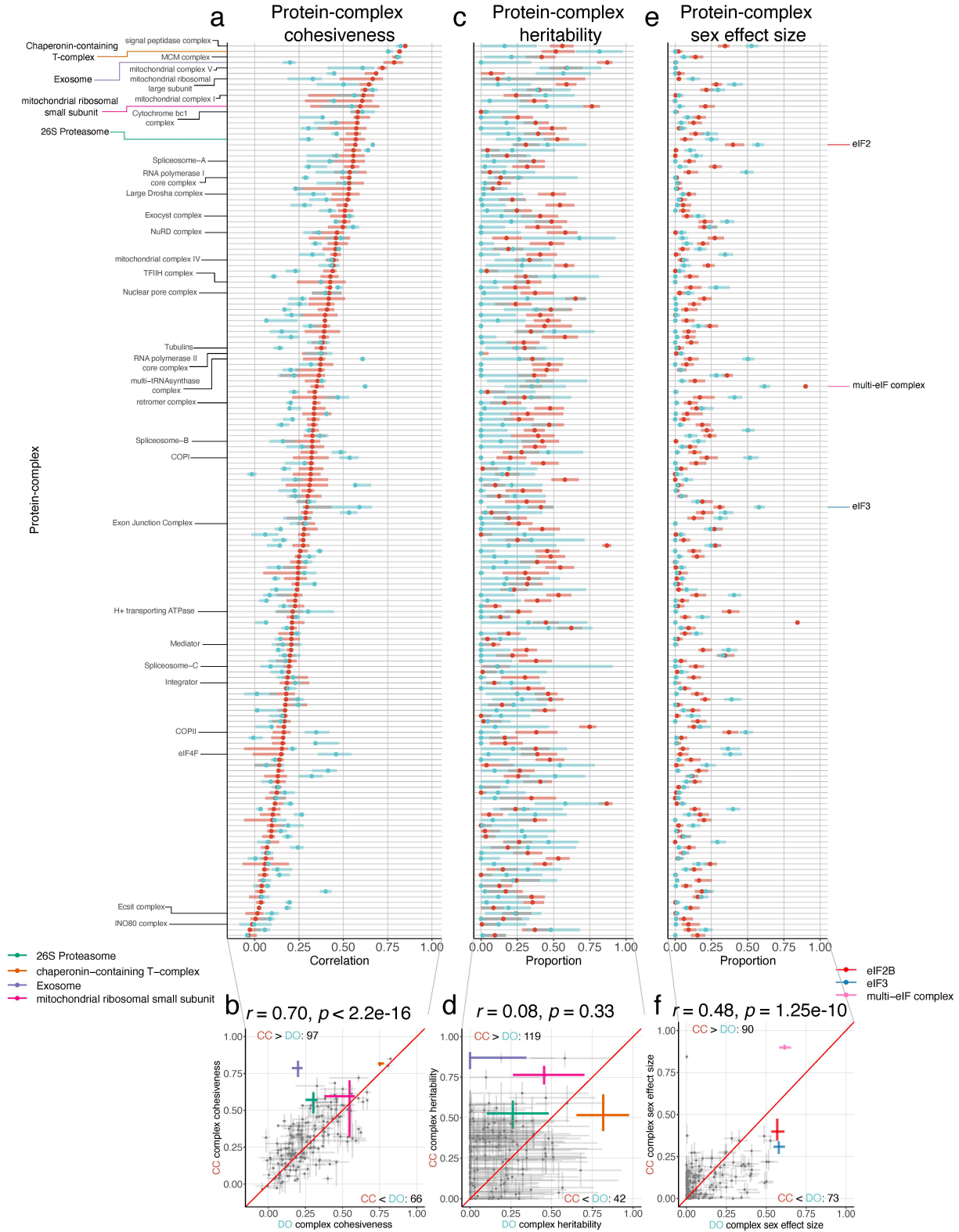
235 **Sex effects on protein abundance are extensively shared across populations**

236 Protein abundances can differ between sexes (Chick *et al.*, 2016), and this pattern can extend
237 collectively to protein complexes (Romanov *et al.*, 2019). We characterized sex effects in the
238 CC, DO, and founder strains (**Methods; Table S1**). Significant sex effects ($FDR < 0.1$) were
239 detected for 3,721 (54.9%) proteins in the CC, 4,376 (66.4%) proteins in the DO, and 1,549
240 (22.3%) proteins in the founder strains. The differences between male and female were
241 overwhelmingly in the same direction for all populations (**Figure 1g-i**). Gene set enrichment
242 analysis revealed that proteins related to ribosomes, translation, and protein transport gene
243 ontology (GO) terms were more abundant in male livers whereas proteins related to catabolic
244 and metabolic processes, including fatty acid metabolism, were more abundant in female livers
245 in all populations.

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247 **Drivers of variation in the abundance of protein-complexes**

248 Members of protein-complexes exhibit varying degrees of co-abundance, with some groups of
249 proteins being tightly co-abundant, *i.e.*, correlated, and others less so (Romanov *et al.*, 2019).
250 Correlation between members of a complex suggests some degree of co-regulation. We found
251 that individual protein members of annotated protein-complexes (Giurgiu *et al.*, 2019; Ori *et al.*,
252 2016; Vinayagam *et al.*, 2013; **Table S5**) are less heritable and fewer of them possess pQTL than



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Figure 3. Genetic and sex effects on protein-complexes. (a-b) The co-abundance of complex members, *i.e.*, complex-cohesiveness, is consistent between the CC (red) and DO (blue) ($r = 0.70, p < 2.2e-16$). Cohesiveness is summarized (Methods) for each of 163 annotated protein complexes (Ori et al., 2016). (c-d) Complex-heritability, based on the first principal

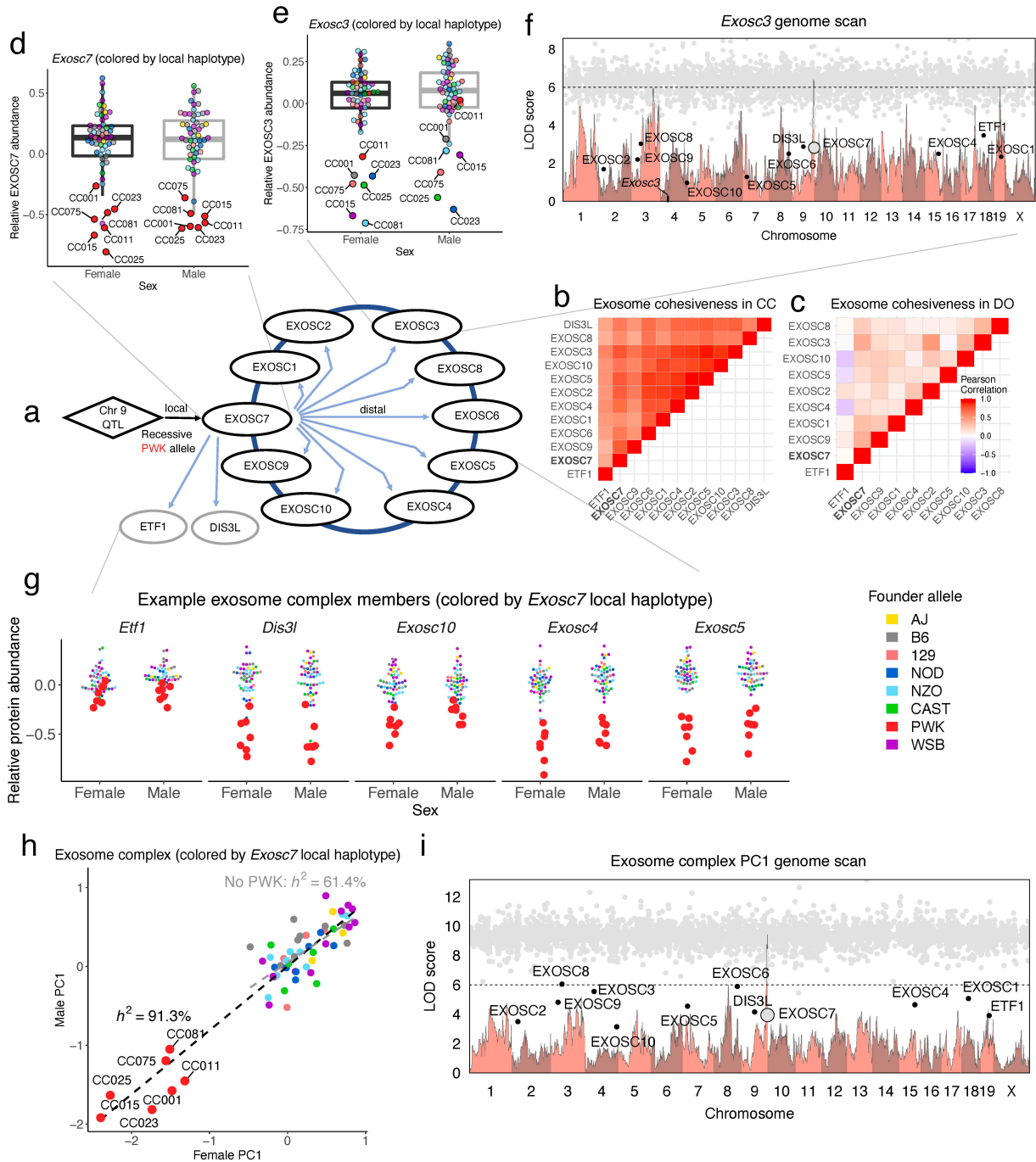
257 component (PC1) from each complex (**Methods**), were inconsistent between the CC and DO ($r = 0.08, p = 0.33$), with heritability
258 notably higher more often in the CC. (e-f) Complex-sex effect size, representing the proportion of variability in the complex PC1
259 explained by sex, was consistent between the CC and DO ($r = 0.48, p = 1.25e-10$). For complex-cohesiveness, the intervals
260 represent the interquartile range, and points represent the median. For complex-heritability and complex-sex effect size,
261 intervals represent 95% subsample intervals. Exosome, chaperonin-containing T-complex, 26S Proteasome, and the
262 mitochondrial ribosomal small subunit are highlighted as examples of highly heritable protein-complexes that are examined in
263 detail (**Figures 4, 5, 6, S5, & S6**). Multi-eIF complex, eIF2B, and eIF3 are highlighted as complexes with large sex effect sizes in
264 both the CC and DO, consistent with a previous study (Romanov et al., 2019). Red diagonal line included for reference.

265 proteins that are not members of complexes (**Figure S4a-d; Table S6**). However, protein-
266 complexes can be influenced by genetic variation, as we previously reported for the Chaperonin
267 containing T (CCT)-complex that is stoichiometrically regulated in response to the low
268 abundance of a member protein, CCT6A (Chick et al., 2016). We evaluated the extent to which
269 members of annotated protein-complexes were internally correlated, as well as how genetic
270 factors and sex contribute to variation in their joint abundance. We summarized co-abundance,
271 which we refer to as complex cohesiveness, as the median pairwise correlation between
272 complex members (Romanov et al., 2019). To assess the contributions from genetic factors and
273 sex, we summarized each protein-complex using the PC1 from member proteins. We first
274 filtered proteins with local pQTL (FDR < 0.5) or strong distal pQTL (FDR < 0.1) to minimize the
275 influence of individual proteins with independent genetic effects in order to focus on the
276 shared genetic effects on a protein-complex. We estimated heritability and the proportion of
277 variation explained by sex for each complex-specific PC1 (**Table S7**).

278 Complex-cohesiveness was correlated between the CC and DO ($r = 0.70, p < 2.2e-16$) (**Figure 3a**
279 **& b**), and within each population, correlated with complex-heritability: $r = 0.33, p = 4.37e-5$ in
280 the CC and $r = 0.17, p = 0.03$ in the DO (**Figure S4e**), suggesting that cohesiveness does reflect
281 genetic factors that control protein abundance at some level. Notably, complex-heritability is
282 consistently higher in the CC than DO (119 out of 163 complexes; 73.0%) and uncorrelated with
283 complex-heritability in the DO ($r = 0.08, p = 0.33$), in contrast to the heritability of individual
284 proteins ($r = 0.43, p < 2.2e-16$) (**Figure 3c & d**). The greater consistency between cohesiveness
285 and heritability of protein-complexes in the CC is consistent with variation in overall protein-
286 complex levels being influenced by non-additive genetic effects. The complex-heritability in CC
287 captures the greater similarity of strain replicates with each other compared to other CC
288 strains. The proportion of variation explained by sex for protein-complex abundance was
289 correlated between the CC and DO ($r = 0.48, p < 1.25e-10$; **Figure 3e-f**). Protein-complexes
290 previously shown to be driven by sex, such as eIF2B (Romanov et al., 2019), were confirmed in
291 both the CC and DO.

292 293 **Genetic and stoichiometric regulation of the exosome**

294 We identified the exosome complex as a novel and striking example of a protein-complex with
295 stoichiometric genetic regulation in the CC (**Figure 4**). The exosome complex had the highest
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 299 **Figure 4. PWK allele at *Exosc7* drives low abundance of the exosome complex and related proteins.** (a) Genetic variation at
 300 *Exosc7* has strong effects on the other members of the exosome complex and *Dis3l* and *Etf1*, which were not annotated
 301 members (Ori et al. 2016) though functionally related. The complex proteins were more tightly correlated in the (b) CC than the
 302 (c) DO, potentially due to the observance of the homozygous PWK genotype and strain replicates in the CC (**Figure S5**). (d) A
 303 local PWK allele resulted in lower EXOSC7 abundance, as observed in seven CC strains (CC001, CC011, CC015, CC023, CC25,
 304 CC075, and CC081). Points are colored by the haplotype allele at *Exosc7*. (e) Local genetic variation at *Exosc3* does not explain
 305 protein abundance patterns, which instead match the genetic variation at *Exosc7*, representing a distal pQTL. (f) The genome
 306 scan for *Exosc3* reveals the distal pQTL near *Exosc7*, and mediation analysis identifies EXOSC7 as a strong mediator of the distal
 307 pQTL (large gray dot) as well as the other exosome-related proteins as mediators (black dots), reflecting shared genetic
 308 regulation. The remaining smaller gray dots represent mediation scores for all other quantified proteins. Horizontal dashed line
 309 at LOD of 6 included for reference. (g) The PWK allele at *Exosc7* distally control other members of the exosome complex, with
Etf1, *Dis3l*, *Exosc10*, *Exosc4*, and *Exosc5* shown here. Points are colored by the founder haplotype at *Exosc7*. (h) The first

310 principal component (PC1) from the exosome complex, plotted males against females for the CC strains. Points are colored by
311 the founder haplotype at *Exosc7*. CC strains separate based on whether they possess the PWK allele of *Exosc7*, reflected in a
312 complex-heritability of 91.3%. After removal of the seven strains that possess the PWK allele, the complex-heritability is 61.4%,
313 which suggest there are remaining loci that affect the abundance of the exosome. The black dashed line is the best fit line
314 between males and females for the complex PC1, based on all 58 CC strains. The gray dashed line excludes the seven CC strains
315 with the PWK allele at *Exosc7*. (i) The genome scan of the complex PC1 reveals a similar scan to *Exosc3*, though the pQTL near
316 *Exosc7* and its mediation are stronger, due to the consistent effects across complex members.

317 complex-heritability in the CC (87.0%, [80.0-90.5%]) while being essentially non-heritable in the
318 DO (0.0%, [0.0-34.6%]). Low EXOSC7 abundance is driven by a local PWK allele in the CC, which
319 is homozygous in seven CC strains, whereas in our DO cohort, there were no mice homozygous
320 for the PWK allele (**Figure S5b**). Heterozygous carriers of the PWK allele in the DO mice
321 exhibited no discernable effect, suggesting that the exosome complex is regulated by a
322 recessive PWK allele at *Exosc7* (**Figure S5c**), which is consistent with founder PWK mice having
323 low EXOSC7 abundance (**Figure S5d**). Mediation analysis indicates that genetic variation at the
324 *Exosc7* locus distally regulates the complex as well as the functionally related genes, *Dis3L* and
325 *Etf1*. These two genes were not included in the original complex annotations, and their co-
326 regulation suggests new biological interactions discovered via our approach. The complex-
327 heritability (with *Dis3L* and *Etf1* included as well as complex members previously filtered out
328 due to possessing pQTL) was 91.3%, and after removing the seven strains with the PWK allele at
329 *Exosc7*, was reduced but still high at 61.4%, indicating the presence of additional genetic factors
330 that affect the abundance of the exosome.

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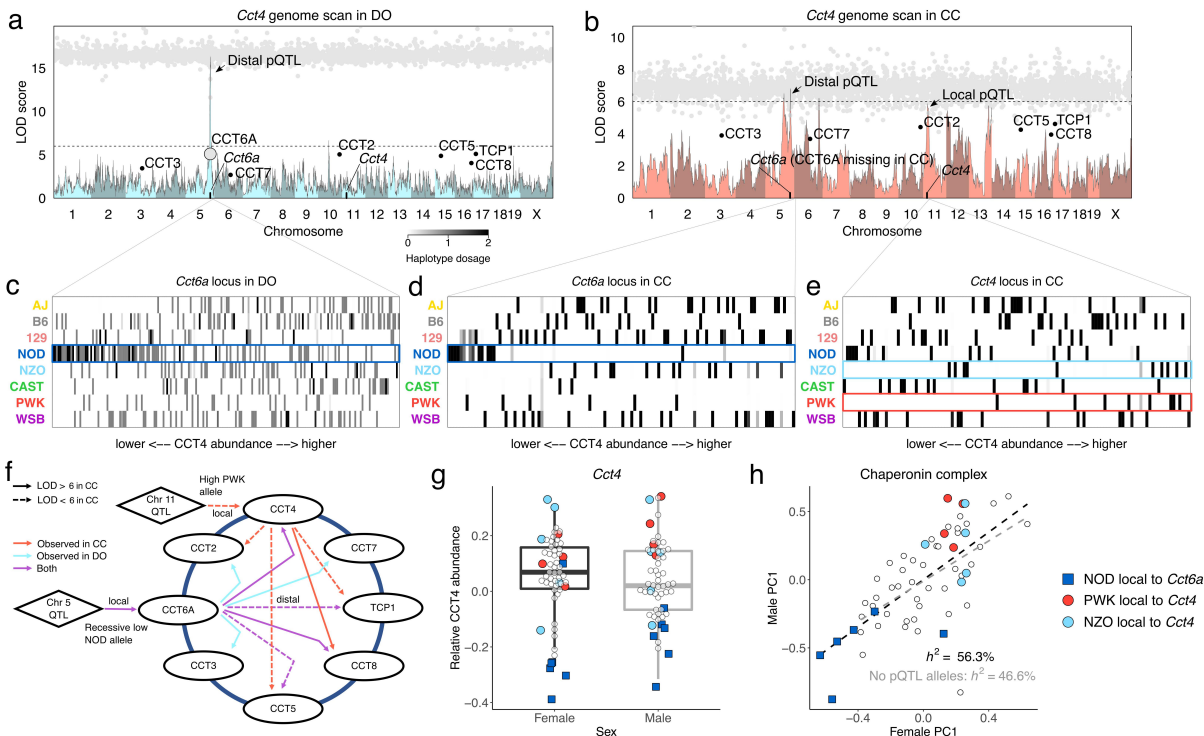
332 **Secondary genetic effect on the chaperonin complex**

333 Previously we found that the CCT complex was stoichiometrically regulated and driven by low
334 abundance of CCT6A when the NOD haplotype is present (Chick et al., 2016). The CCT complex
335 (**Figure 5**) has higher heritability in the DO (81.6%, [65.3-97.5%]) than in the CC (51.6%, [41.8-
336 64.4%]) (**Figure 3c & d**). The DO sample is well-powered to detect the low NOD allele that drives
337 the complex-wide regulation, due to 19 (9.9%) of the mice being homozygous NOD. The CC
338 strains replicate the distal pQTL at the locus of *Cct6a* through a low NOD effect for complex
339 members (*Cct4*, *Cct5*, *Cct8*, and *Tcp1*) – but CCT6a itself was not quantified in the CC samples.
340 The effect of the pQTL at *Cct6a* drives less of the overall variation in the CC sample due to fewer
341 occurrences of NOD homozygotes (12 CC mice in comparison to 19 DO mice). The CC reveals a
342 secondary genetic effect mediated through CCT4, with high NZO and PWK alleles. After
343 including complex members that were previously filtered out for possessing pQTL, the complex-
344 heritability was 56.3%, which, after excluding CC strains that are NOD at *Cct6a* and NZO or PWK
345 at *Cct4*, was still significant at 46.6% (**Figure 5h**), indicating that, as with the exosome complex,
346 additional genetic effects contribute to CCT complex abundance.

347

348 **Independent genetic effects on the components and subcomplexes of the 26S proteasome**

349 The 26S proteasome is composed of a 20S proteasome catalytic core (PSMA and PSMB
350 proteins), which in the constitutive form has the constitutive subunits, PSMB5, PSMB6, and



351
 352
 353 **Figure 5. Genetic effects on the chaperonin-containing T-complex.** Genome scans overlaid with mediation analysis for *Cct4*, a
 354 member of chaperonin-containing T (CCT)-complex, in the (a) DO and (b) CC, reveal a shared distal pQTL on chromosome 5. The
 355 CC also have a suggestive local pQTL for *Cct4* on chromosome 11. CCT4 was previously revealed through mediation analysis to
 356 be distally controlled through genetic variation at *Cct6a* (large gray dot) in the DO (Chick et al., 2016). The other CCT member
 357 proteins were detected as mediators (black dots) as well, stemming from shared genetic effects. The remaining smaller gray
 358 dots represent mediation scores of all other quantified proteins. The mediation scan of the distal pQTL on chromosome 5 in the
 359 CC similarly detects the other members of the CCT complex, though the likely causal intermediate CCT6A was not quantified in
 360 the CC. Horizontal dashed line at LOD of 6 included for reference. The founder haplotype inheritance, represented as heatmaps
 361 with founder allele dosage (expected allele counts) as rows and individual mice as columns, ordered by CCT4 abundance, at the
 362 chromosome 5 locus near *Cct6a* in the (c) DO and (d) CC and the (e) chromosome 11 locus near *Cct4* in the CC. A low NOD
 363 effect, potentially recessive and highlighted with dark blue boxes, drives the distal pQTL (likely through CCT6A) on CCT4 in both
 364 the DO and CC. The local effects on *Cct4* were not as strong, though NZO and PWK had subtle high effects, highlighted as light
 365 blue and red boxes. (f) A diagram of the population-specific and shared effects observed through pQTL and mediation analyses
 366 in the DO and CC. Many of the pQTL in the CC are weak, with all falling below the FDR < 0.1 threshold, and some with LOD < 6
 367 (FDR > 0.5), but they are supported by overlap with the DO (chromosome 5) or being local to a complex member (chromosome
 368 11). (g) In the CC, CCT4 abundance is affected by genetic variation at *Cct6a* as well as its local haplotype. The dark blue squares
 369 represent the six strains that possess the NOD allele at *Cct6a*, which had low CCT4. The light blue and red circles are the strains
 370 with high CCT4 that possess NZO and PWK alleles at *Cct4*. (e) The first principal component (PC1) for the complex, plotted males
 371 against females, reveals the same pattern of low strains (NOD allele of *Cct6a*; dark blue squares) and the high strains (NZO and
 372 PWK alleles of *Cct4*; light blue and red circles, respectively). The complex-heritability was high at 56.3% and after removal of the
 373 strains possessing notable alleles from pQTL at *Cct6a* and *Cct4*, dropped to 46.6%. The black dashed line represents the best fit
 374 line between males and females for complex PC1 based on all strains, and the gray dashed line is the best fit line with strains

375 PSMB7, and is capped by two 19S regulators (composed of the PSMC and PSMD proteins). The
 376 constitutive form can be modified into the immunoproteasome by replacing the respective
 377 constitutive subunits with the three immunoproteasome inducible subunits, PSMB8, PSMB9,
 378 and PSMB10, and the 19S regulators with the 11S regulators (made up of PSME proteins)
 379 (Marshall and Vierstra, 2019; **Figure 6a**). The immunoproteasome is a highly efficient form of
 380 the proteasome that is predominantly expressed in immune cells (Kimura et al., 2015).

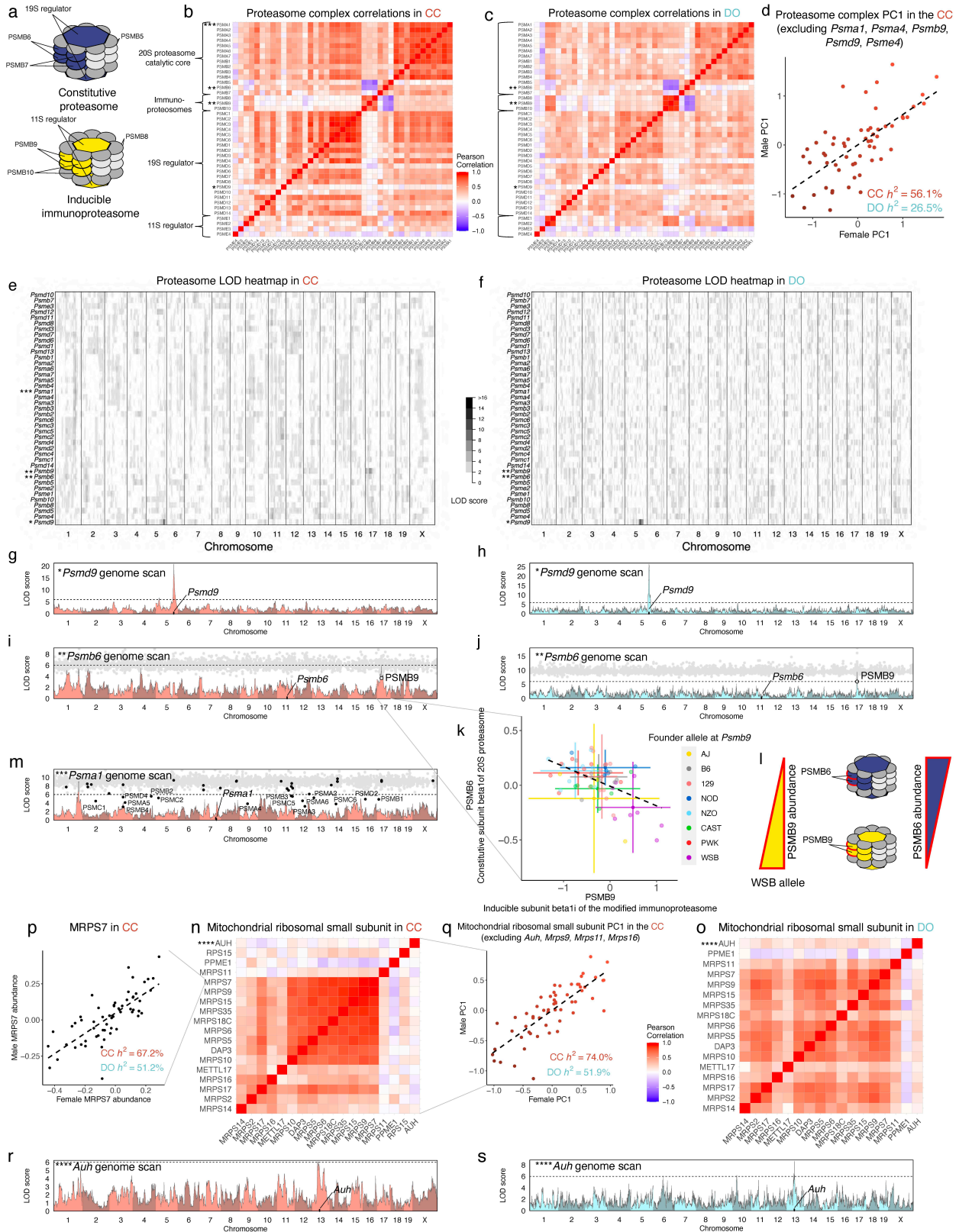
381 This alternation between two different forms of the proteasome is apparent in the anti-
382 correlation between the inducible and immune components in both the CC and DO (**Figure 6b**
383 **& c**) and founder strains (**Figure S6a**). The inverse relationship between the constitutive and
384 inducible forms suggests that although individual samples vary in the relative abundance of the
385 constitutive proteasome and immunoproteasome, they predominantly express one of the
386 forms. Across the founder strains, this relationship appears to be genetically regulated, with the
387 WSB, AJ, and NZO strains expressing more immunoproteasome, and the others expressing
388 more of the constitutive form (**Figure S6b-g**). In the recombinant CC and DO, we identified a
389 genetic variant that controls the balance between PSMB6 (constitutive) and PSMB9
390 (immunoproteasome). In both the CC and DO, genetic variation near *Psmb9* affected PSMB9
391 abundance, as well as PSMB6 abundance, confirmed through mediation analysis (**Figure 6i & j**).
392 Consistent with the anti-correlation between inducible and constitutive subunits as well as the
393 balance in the founder strains, the relationship between PSMB9 and PSMB6 is negative, *e.g.*,
394 mice that inherited the WSB haplotype at the *Psmb9* locus have high PSMB9 abundance and
395 low PSMB6 abundance (**Figure 6k & l**). The pQTL only explains the balance of PSMB6/PSMB9,
396 and while consistent with the overall balance of the constitutive and inducible forms of the
397 proteasome observed in the founder strains, it does not directly affect the other
398 interchangeable members of the proteasome, which do not possess their own strong pQTL. The
399 loss in the CC and DO of the co-regulation of all constitutive and inducible members suggests
400 epistatic factors may interact with pQTL at *Psmb9* and those relationships are broken apart
401 in the recombinant populations, thus decoupling PSMB9 from the other immunoproteasome
402 subunits.

403 Genetic factors also influence other components of the 26S proteasome. We identified a strong
404 local pQTL that is consistent in both the CC and DO for *Psmid9* that does not affect other
405 members of 19S regulator (**Figure 6g & h**), thus explaining the lack of cohesiveness of PSMD9
406 within the proteasome, which was previously observed in the DO (Romanov et al., 2019). A
407 distal pQTL hotspot comprising members of the 20S catalytic core and 19S regulator mapped to
408 a short interval on chromosome 1 in the CC, most strongly observed in *Psma1* (**Figure 6m**). No
409 single, strong mediator was detected for the hotspot, though PEX19 was the best candidate for
410 a number of the proteins, suggesting there may be multiple drivers of the hotspot or that the
411 true driver was not observed at the protein-level.

412

413 **Polygenic regulation of the mitochondrial ribosomal small subunit**

414 The mitochondrial ribosomal small subunit was highly cohesive in both populations (**Figures 3a,**
415 **b, 6n, & o**). The complex-heritability is also high, and after including all annotated members,
416 was 74.0% in the CC and 51.9% in the DO. Despite high complex-heritability, we detected few
417 pQTL for individual members of the complex. The one notable exception is *Auh*, for which we
418 detected a strong local pQTL in the CC and DO (**Figure 6r & s**).



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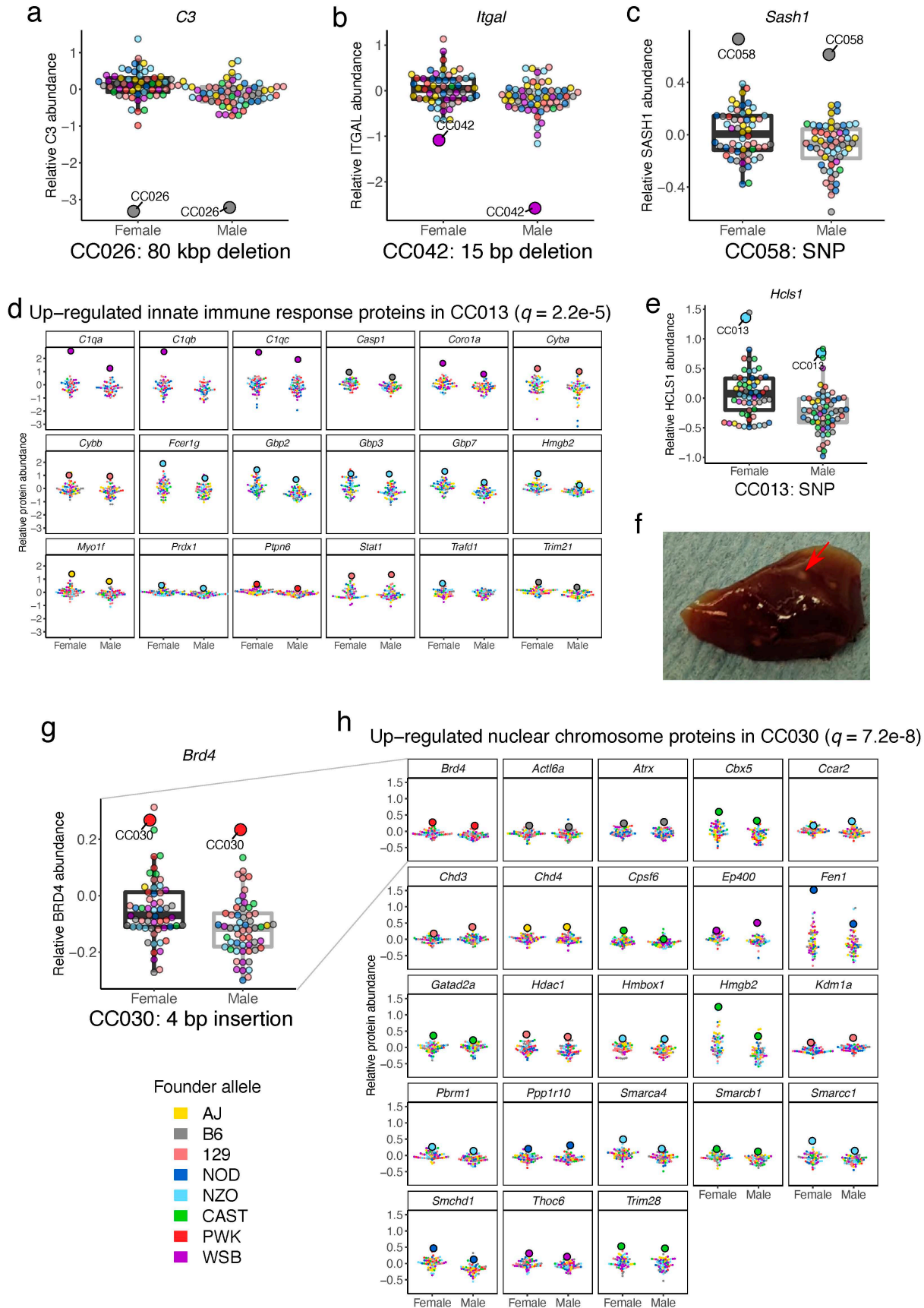
Figure 6. Polygenic regulation of the 26S proteasome and the mitochondrial ribosomal small subunit. (a) The 26S proteasome is composed of multiple subcomplexes: the 20S proteasome catalytic core (*PsmA* and *PsmB* genes) and 19S regulator (*PsmC* and *PsmD* genes) for the constitutive form, and the inducible immunoproteasomes (*PsmB8*, *PsmB9*, and *PsmB10*) with their 11S

423 regulator (*Psme* genes). The correlation patterns between member proteins reflect these subcomplexes, which are highly
424 consistent between the (b) CC and (c) DO (and founder strains; **Figure S6a**). The overall correlation pattern reflects an inverse
425 relationship between the constitutive proteasome and the inducible immunoproteasome. (d) CC strain identity explains a large
426 portion of the variability in the first principal component (PC1) of the proteasome (after excluding genes with strong pQTL) and
427 the complex-heritability is notably higher in the CC (56.1%) than DO (26.5%). The dashed line represents the best fit line
428 between males and females in the complex PC1. A portion of the complex correlation structure can be explained by pQTL,
429 shown here as a heatmap of genome scans (proteins as rows and genomic coordinate as columns), for the (e) CC and (f) DO.
430 *Psm9* (*) has a strong local pQTL in both the (g) CC and (h) DO, which is not shared with other proteasome proteins, explaining
431 why *Psm9* is poorly correlated with other members of the 19S proteasome. Horizontal dashed line at LOD of 6 included for
432 reference. *Psm6* (**) has a distal pQTL in both the (i) CC and (j) DO, which is mediated by PSMB9. Gray dots represent
433 mediation scores for all quantified proteins, with PSMB9 highlighted as a large gray dot. (k) The relationship is negative –
434 individuals with low abundance of PSMB9 (most notably CC strains with the WSB allele at *Psm9*) have high abundance of
435 PSMB6. Intervals represent mean \pm 2 standard deviation bars. The dashed line represents the best fit line between PSMB6 and
436 PSMB9 for the CC mice. (l) PSMB6 is a constitutive subunit of the 20S proteasome and PSMB9 is the corresponding inducible
437 subunit in the modified immunoproteasome, suggesting that the inverse relationship between constitutive proteasome and
438 immunoproteasome is genetically controlled for at least PSMB6 and PSMB9 in the CC and DO (though more broadly controlled
439 in the founder strains; **Figure S6b-g**). A distal pQTL hotspot was observed in the CC on chromosome 1 (~170 Mbp) which was
440 not present in the DO, for which (m) *Psm1* (***) has the strongest signal. Though a strong mediator was not detected from the
441 proteins near the pQTL, many members of the 20S and 19S proteasomes present as strong mediators (black dots) due to the
442 strong correlation among the proteins that map to the hotspot. The mitochondrial ribosomal small subunit was more heritable
443 and cohesive than the 26S proteasome (**Figure 3**), which is evident in the correlation patterns for both the (n) CC and (o) DO.
444 The individual proteins, such as (p) MRPS7, and the (q) complex PC1 (after excluding genes with strong pQTL) are highly
445 consistent within CC strains despite few strong pQTL related to the complex. The dashed line represents the best fit line
446 between males and females in MRPS7 and complex PC1. Similar to *Psm9* within the 26S proteasome, a less cohesive member
447 of the complex, *Auh* (****), can be explained by a unique local pQTL detected in both the (r) CC and (s) DO.

448 AUH was incohesive with the core of the complex, composed of mostly MRPS proteins. The
449 complex PC1 and individual proteins are highly consistent within CC strain ($r = 0.81$ for the
450 complex PC1; $r = 0.76$ for MRPS7) while displaying continuous variation across the CC
451 population (**Figures 6p & q**). This distribution contrasts with the bimodal pattern of abundance
452 for the exosome complex, which is driven by a single strong pQTL (**Figure 4h**) and suggests that
453 many loci with small effects influence the overall abundance of the mitochondrial ribosomal
454 small subunit.

455 456 **Strain-private variants affect protein abundance**

457 Inbred mouse strains can accumulate mutations, and these variants can lead to phenotypic
458 abnormalities across classical inbred strains, *e.g.*, new mutations that become fixed in sub-
459 strains of B6 (Kumar et al., 2013) and in the parental B6 and DBA stocks (Anderson et al., 2002)
460 used across different epochs of the BxD panel (Ashbrook et al., 2019; Mulligan et al., 2012).
461 New mutations originate and became fixed in the CC strains (Shorter et al., 2019; Srivastava et
462 al., 2017) (**Figure 1a**). As a proof of concept, we first confirmed the loss of expression of
463 proteins with known deletions, such as the 80 kbp deletion in CC026 that includes *C3* and a 15
464 bp deletion for CC042 in *Itgal* (**Figure 7a-b**) that increases susceptibility to tuberculosis (Smith
465 et al., 2019) and salmonella (Zhang et al., 2019). Next, we estimated CC strain-specific effects
466 for each protein in the CC and identified CC strains with extreme abundance of a given protein,
467 referred to as strain-protein outliers (**Methods**). In total, we identified 6,046 strain-protein



469 **Figure 7. Strain-private genetic variants affect protein abundance and potentially influence larger protein networks.** The
470 effects of known strain-private deletions on protein abundance were confirmed for (a) *C3* in CC026 and (b) *Itgal* in CC042. The
471 color or the dots indicates the founder allele at the gene. (c) A novel SNP allele in *Sash1* specific to CC058 is associated with an
472 increase in SASH1 abundance. (d) Proteins related to innate immune response and other related pathways were high abundant
473 in CC013. Large dots represent the specified CC strain with extreme protein abundance. (e) CC013 also possesses a strain-
474 specific SNP allele in *Hcls1*, a gene involved in leukocyte differentiation, which may contribute to its unique protein abundance
475 patterns in immune pathways. (f) CC013 has a unique liver phenotype, characterized by white granules, highlighted with red
476 arrow, which may be related to the increased abundance of immune proteins. CC030 possesses a (g) private insertion in *Brd4*,
477 which may contribute to (h) increased abundance in genes related to nuclear chromosome. Additional functionally related
478 strain-specific protein dynamics are shown in **Figure S7**.

479 outliers, representing 4,323 proteins across all 58 CC strains. The strain-protein outliers
480 coincided with 69 known strain-private genetic variants (Srivastava et al., 2017), meaning the
481 strain-protein outlier could represent a local effect of the private variant associated with the
482 protein's coding gene, and this level of enrichment was significant per permutation ($p = 3.7e-4$).
483 Interestingly, not all of the observed outliers associated with genetic variants were low
484 extremes, as would be expected with a mutation that results in loss of the protein; we also
485 observed increases in protein abundance in strains with private variants, such as *Sash1*, which
486 harbors a novel SNP allele in CC058 (**Figure 7c**). CC004 has a unique SNP allele associated with
487 low abundance of *Plek*, a gene which also has a weak local pQTL based on genetic variation
488 from the founder strains (**Figure S7e & f**).

489 The presence of biologically related proteins with extreme abundances specific to CC strains
490 further supports the impact of strain-specific genetic variants and regulatory patterns. We
491 identified strain-specific dynamics by testing the set of outlying proteins specific to each CC
492 strain for enrichment in GO and KEGG pathway terms (**Tables S8 and S9**). In CC013, we
493 observed increased abundance in proteins with GO annotations for the innate immune system
494 (**Figure 7d**), leukocytes, and other immune system-related GO terms. CC013 possesses a unique
495 SNP allele in *Hcls1* that was associated with increased HCLS1 abundance (**Figure 7e**), a gene
496 involved in myeloid leukocyte differentiation that may contribute to the high abundance of
497 immune-related proteins in CC013. CC013 also expressed a unique liver phenotype,
498 characterized by white granules across the tissue (**Figure 7f**), which may relate to the excess of
499 immune-related proteins. CC030 has a 4 bp insertion in *Brd4* that was associated with high
500 BRD4 abundance and may contribute to increased abundance in proteins related to nuclear
501 chromosome and chromatin (**Figure 7g-h**). The strain-specific protein outlier sets were enriched
502 in a wide range of GO biological functions (**Figure S7g-j**), including low and high abundance of
503 proteins from the mitochondrial respiratory complex I in CC007 and high abundance of
504 cytosolic ribosome proteins in CC009.

505

506 **DISCUSSION**

507 Multiparent populations are key resources for understanding genetic architecture; developing
508 new phenotypic models of disease; and producing robust results that translate from model

509 organisms to genetically diverse outbred populations such as humans. Here we show that the
510 CC, DO, and their founder strains broadly share protein heritability, pQTL, their mediators, and
511 sex effects on proteins. In comparison to individual proteins, protein-complexes were less
512 consistent across these populations, highlighting the role of non-additive genetic effects in
513 controlling protein-complex interactions. We observed a wide range of genetic effects on
514 protein complexes, ranging from stoichiometric regulation in response to a large-effect locus
515 (exosome and CCT complexes) to multi-locus and highly polygenic regulation (*e.g.*, the 26S
516 proteasome and mitochondrial ribosomal small subunit, respectively). Within the CC, we
517 observed strong effects on protein abundance from new mutations in specific CC strains. Lastly,
518 we highlight individual CC strains with both aberrant protein regulation based on strain-specific
519 mutations that broadly disrupt protein regulatory networks.

520

521 **Conservation of pQTL between CC and DO**

522 Consistent with expectations, the effects of local genetic variation are highly conserved
523 between the CC and DO (and often with the founder strains). In cases where discordance
524 occurred between populations, we can often explain based on differing founder allele
525 frequencies between the populations, such as for the local pQTL of *Ercc2* that is not detected in
526 the DO because only one NOD homozygote was observed in the DO sample. Differences
527 between populations stemming from different allele frequencies has been observed in human
528 populations (Mogil et al., 2018). These discordant cases highlight how dominance and large-
529 scale homozygosity contribute to the genetic effects on proteins, exposed by comparing the CC
530 and DO. Distal (*i.e.*, *trans*) pQTL are often harder to detect due to weaker effects and are thus
531 more likely to be discordant based on differences in allele frequencies and the presence of non-
532 additive effects. Here we showed that the 19 distal pQTL that were detected in both
533 populations at FDR < 0.1 are highly consistent, both in terms of haplotype effects and their
534 mediation candidates. Discordance in effects between the CC and DO was much greater for
535 distal pQTL detected in a single population, which is in striking contrast to the strong
536 concordance of local genetic effects detected in only one population. The extent of discordance
537 in distal pQTL effects compared to local effects suggests that additional factors are contributing
538 beyond differing allele frequencies and may represent distal genetic effects that are not
539 conserved across population.

540 Mediation analysis results mirror the concordance of distal pQTL effects, with largely the same
541 mediators detected for distal pQTL detected with high confidence in both populations. There
542 are some key caveats with mediation analysis, such as its accuracy being dependent on the true
543 mediator being present in the data. If a candidate mediator is correlated with the true but
544 unobserved mediator, possibly due to LD, it will likely be identified as a false positive mediator.
545 This issue is problematic for proteomics studies if the mediator is not captured in the protein

546 data, such as non-coding RNAs or lowly abundant proteins that go undetected in the mass-spec
547 analysis. For example, PGD was detected as a candidate mediator of a distal pQTL of *Akr1e1* in
548 both the CC and DO; however, previous studies identified zinc finger proteins (*Rex2* and *Zfp985*)
549 as the most likely mediator of the pQTL (Hamilton-Williams et al., 2010; Keele et al., 2020). Zinc
550 finger proteins are lowly expressed and not prevalent in proteomics data, and in their absence,
551 the nearby protein PGD stands out as the best mediator in both populations. Variable
552 measurement error across candidate mediators can also cause preference for a false mediator
553 due to noise reducing the correlation between the distally controlled protein and its true
554 mediator. This is likely occurring in the DO where NAXD is the stronger mediator for the *Tubg1*
555 distal pQTL than TUBGCP3, the strong biological candidate. Furthermore, the presence of
556 variable measurement error in proteomics is likely, stemming from the overall magnitude of
557 each protein's expression, the size of the protein and the number of peptides used to
558 summarize it, and how specifically those peptides map to the protein. Comparisons of
559 mediation analysis of two independent genetic experiments can provide replication of findings,
560 as well as assess a more complete set of candidate mediators and correct misidentifications.

561

562 **Non-additive genetic effects on proteins and protein-complexes**

563 Protein-complexes can be viewed as emergent phenotypes that can be driven by independently
564 regulated members or sub-complexes, as well as higher order dynamics like protein-protein
565 stoichiometry that control complex assembly. We observed a spectrum of cohesiveness
566 (Romanov et al., 2019) across the protein-complexes and our three populations. These
567 differences reflected components and sub-complexes that are semi-independent of the greater
568 complex, as well as the underlying genetic architectures of the populations. Because high
569 cohesiveness does not necessarily imply that a protein-complex is genetically regulated (*e.g.*,
570 stoichiometry could produce tight correlation among complex members due to non-genetic
571 factors), we employed genetic analyses (*e.g.*, QTL and heritability) to characterize genetic
572 control of the protein-complexes. We found a number of strong and distinct examples of
573 protein-complexes that are genetically regulated. Despite the challenges imposed by separate
574 experiments and the relative nature of mass-spec proteomics (O'Brien et al., 2018), comparing
575 the CC to the DO reveals the impact of an inbred genetic background on a number of protein-
576 complexes, due to recessive effects, or conversely, the lack of dominance genome-wide.
577 Furthermore, CC strain replicates can capture multi-locus interactions, *i.e.*, epistasis, by fixing
578 alleles at multiple loci within a strain. These results support the CC as a distinctly powerful tool
579 for genetic analyses of emergent phenotypes like protein-complexes and as a companion
580 resource to the DO for disentangling complicated genetic mechanisms.

581 The examples of genetically regulated protein-complexes that we identified represented a
582 continuum of polygenicity underlying their genetic architecture. Examples ranged from the

583 exosome with a single large effect pQTL local to a single member driving most variation, to the
584 highly polygenic mitochondrial ribosomal small subunit, reflected in its continuous distribution
585 across the CC strains and lack of pQTL. Intermediate to these extremes are examples like the
586 CCT complex with its combination of a large effect pQTL with subtle secondary genetic effects
587 revealed in the CC, and the 26S proteasome with its well-defined sub-complexes representing
588 two distinct forms for altered biological function, the balance of which is genetically controlled
589 to various degrees in these three populations. A common feature shared by these example
590 protein-complexes examples is some degree of polygenicity. Even the fairly monogenic
591 exosome and CCT complexes have some residual heritability after removing the CC strains that
592 possess the contrasting alleles of the pQTL, suggesting other loci have secondary effects. The
593 polygenic effects that drives the residual heritability may also be non-additive. In the DO, the
594 exosome and CCT complex – after accounting for the strong pQTL of *Cct6a* – appear non-
595 heritable, either due to large-scale dominance or the inability to capture epistatic effects in an
596 outbred genetic background. These examples highlight the challenge of dissecting polygenic
597 and non-additive effects down to individual loci and their specific mechanisms, emphasizing the
598 value of related genetic resource populations with differing genetic architecture, like the CC
599 and DO.

600

601 **New models of aberrant protein functional networks and disease**

602 The unique biology of individual mouse strains has led to a variety of discoveries of genetically
603 based disease phenotypes. We leveraged the CC population to identify strains exhibiting
604 aberrant protein dynamics that could not be tied to the founder strains, such as being
605 downstream of mutations unique to specific CC strains. They may also result from unique
606 combinations of alleles of upstream drivers of a shared functional network. Examples include
607 increased abundance of immune-related proteins and unique liver phenotype in CC013, which
608 we are following up, and altered mitochondrial respiratory complex I function in CC007. Due to
609 the replicability of the CC population, these strain-specific protein networks can be followed up,
610 confirmed, and the underlying mechanisms dissected to reveal new biology and develop new
611 models of disease.

612

613 **Integrative genetic resource populations**

614 Proteins are a more functionally relevant measure of physiology and disease than the more
615 commonly measured gene transcripts. In our examination of protein regulation across three
616 related populations, we found that local genetic and sex effects on protein abundance were
617 consistent, suggesting that molecular phenotype data and their findings can largely be
618 integrated across these populations. It is easy to conceive of investigators querying specific
619 genes of interest to assess whether their proteins possess sex effects, pQTL and their haplotype

620 effects, or mediators of distal pQTL. To enable these queries, we provide interactive QTL
621 analysis tools for both the CC (<https://churchilllab.jax.org/qtlviewer/CC/Ferris>) and DO
622 (<https://churchilllab.jax.org/qtlviewer/DO/Svenson>). In contrast, higher order molecular
623 phenotypes or characteristics, such as protein-complexes and regulatory networks, showed
624 greater discordance across these populations. Non-additive genetic effects, either at a single
625 locus or across loci, were apparent in the regulation of protein-complexes, driving the
626 discordance between populations based on the genetic architecture of each. Lastly, we
627 identified CC strain-specific aberrant protein abundances, their phenotypic or system relevance,
628 and their putative consistency with previously described mutations present across these
629 strains.

630 In this work, we used these diverse mouse populations to finely dissect genetic effects on
631 protein-complexes in liver tissue, revealing the presence of non-additive effects and a polygenic
632 spectrum of genetic regulatory patterns. In the future, we envision highly expandable joint
633 population resources, covering a range of molecular phenotypes (*e.g.*, RNA-seq, ATAC-seq,
634 mass-spec proteomics) and tissues relevant to human disease.

635

636 **METHODS**

637 **Founder and CC strains**

638 The CC and DO are descended from eight inbred founder strains: A/J (AJ), C57BL/6J (B6),
639 129S1/SvImJ (129), NOD/ShiLtJ (NOD), NZO/H1LtJ (NZO), CAST/EiJ (CAST), PWK/PhJ (PWK), and
640 WSB/EiJ (WSB). We previously collected, processed, and quantified liver proteins from two
641 females and two males from each founder strain as well as 192 DO mice (Chick et al., 2016).

642 We received pairs of young mice from 58 CC strains from the UNC Systems Genetics Core
643 Facility between the summer of 2018 and early 2019. Mice were singly housed upon receipt
644 until eight weeks of age. The 58 CC strains used in this study include: CC001/Unc (CC001),
645 CC002/Unc (CC002), CC003/Unc (CC003), CC004/TauUnc (CC004), CC005/TauUnc (CC005),
646 CC006/TauUnc (CC006), CC007/Unc (CC007), CC008/GeniUnc (CC008), CC009/UncJ (CC009),
647 CC010/GeniUnc (CC010), CC011/Unc (CC011), CC012/GeniUnc (CC012), CC013/GeniUnc
648 (CC013), CC015/Unc (CC015), CC016/GeniUnc (CC016), CC017/Unc (CC017), CC019/TauUnc
649 (CC019), CC021/Unc (CC021), CC023/GeniUnc (CC023), CC024/GeniUnc (CC024),
650 CC025/GeniUnc (CC025), CC026/GeniUnc (CC026), CC027/GeniUnc (CC027), CC029/Unc
651 (CC029), CC030/GeniUnc (CC030), CC031/GeniUnc (CC031), CC032/GeniUnc (CC032),
652 CC033/GeniUnc (CC033), CC035/Unc (CC035), CC036/Unc (CC036), CC037/TauUnc (CC037),
653 CC038/GeniUnc (CC038), CC039/Unc (CC039), CC040/TauUnc (CC040), CC041/TauUnc (CC041),
654 CC042/GeniUnc (CC042), CC043/GeniUnc (CC043), CC044/Unc (CC044), CC045/GeniUnc
655 (CC045), CC046/Unc (CC046), CC049/TauUnc (CC049), CC051/TauUnc (CC051), CC053/Unc
656 (CC053), CC055/TauUnc (CC055), CC057/Unc (CC057), CC058/Unc (CC058), CC059/TauUnc
657 (CC059), CC060/Unc (CC060), CC061/GeniUnc (CC061), CC062/Unc (CC062), CC071/TauUnc
658 (CC071), CC072/TauUnc (CC072), CC075/UncJ (CC075), CC078/TauUnc (CC078), CC079/TauUnc

659 (CC079), CC080/TauUnc (CC080), CC081/Unc (CC081), and CC082/Unc (CC082). More
660 information regarding the CC strains can be found at
661 <http://csbio.unc.edu/CCstatus/index.py?run=AvailableLines.information>.

662

663 **Mouse genotyping, founder haplotype reconstruction, and gene annotations**

664 The 116 CC mice were genotyped on the Mini Mouse Universal Genotyping Array (MiniMUGA),
665 which includes 11,125 markers (Sigmon et al., 2020). Founder haplotypes were reconstructed
666 using a Hidden Markov Model (HMM), implemented in the qtl2 R package (Broman et al.,
667 2019), using the “risib8” option for an eight founder recombinant inbred panel. Notably,
668 heterozygous markers are omitted, and haplotype reconstructions are limited to homozygous
669 states, smoothing over potential residual heterozygous sites that remain in the CC mice. The
670 genotyping and haplotype reconstruction for the DO mice were previously described (Chick et
671 al., 2016); briefly, genotyping was performed on the larger MegaMUGA (57,973 markers)
672 (Morgan and Welsh, 2015), from which founder haplotypes were reconstructed using the
673 DOQTL R package (Gatti et al., 2014).

674 Ensembl version 91 gene and protein annotations were used in the CC, whereas version 75 was
675 previously used in the older DO and founder strains data. If the gene symbol or gene ID differed
676 for a protein ID between versions 75 and 91, we updated them to version 91 in the DO and
677 founder strains. When comparing results (*e.g.*, pQTL, heritability, sex effects) between the CC
678 and the DO or founder strains, we merged based on protein ID. For the more complicated
679 results from mediation analysis, we allowed matches based on mediator gene symbol if the
680 target protein IDs matched.

681

682 **Sample preparation for proteomics analysis**

683 The sample preparation and mass-spec experimentation and analysis for the DO and founder
684 strains were previously described (Chick et al., 2016). Singly housed CC mice had their food
685 removed six hours prior to euthanasia and tissue harvest. Tissues were dissected out, weighed,
686 and snap frozen in liquid nitrogen. Pulverized CC liver tissue were syringe-lysed in 8 M urea and
687 200 mM EPPS pH 8.5 with protease inhibitor and phosphatase inhibitor. BCA assay was
688 performed to determine protein concentration of each sample. Samples were reduced in 5 mM
689 TCEP, alkylated with 10 mM iodoacetamide, and quenched with 15 mM DTT. 200 µg protein
690 was chloroform-methanol precipitated and re-suspended in 200 µL 200 mM EPPS pH 8.5. The
691 proteins were digested by Lys-C at a 1:100 protease-to-peptide ratio overnight at room
692 temperature with gentle shaking. Trypsin was used for further digestion for 6 hours at 37°C at
693 the same ratio with Lys-C. After digestion, 50 µL of each sample were combined in a separate
694 tube and used as the 11th sample in all 12 tandem mass tag (TMT) 11plex. 100 µL of each
695 sample were aliquoted, and 30 µL acetonitrile (ACN) was added into each sample to 30% final
696 volume. 200 µg TMT reagent (126, 127N, 127C, 128N, 128C, 129N, 129C, 130N, 130C, 130N,
697 and 131C) in 10 µL ACN was added to each sample. After 1 hour of labeling, 2 µL of each sample
698 was combined, desalted, and analyzed using mass-spec. Total intensities were determined in
699 each channel to calculate normalization factors. After quenching using 0.3% hydroxylamine, 11
700 samples were combined in 1:1 ratio of peptides based on normalization factors. The mixture
701 was desalted by solid-phase extraction and fractionated with basic pH reversed phase (BPRP)

702 high performance liquid chromatography (HPLC), collected onto a 96 well plate and combined
703 for 24 fractions in total. Twelve fractions were desalted and analyzed by liquid
704 chromatography-tandem mass spectrometry (LC-MS/MS).

705

706 **Off-line basic pH reversed-phase (BPRP) fractionation**

707 We fractionated the pooled TMT-labeled peptide sample using BPRP HPLC (Wang et al., 2011).
708 We used an Agilent 1200 pump equipped with a degasser and a photodiode array (PDA)
709 detector. Peptides were subjected to a 50-min linear gradient from 5% to 35% acetonitrile in 10
710 mM ammonium bicarbonate pH 8 at a flow rate of 0.6 mL/min over an Agilent 300Extend C18
711 column (3.5 μ m particles, 4.6 mm ID, and 220 mm in length). The peptide mixture was
712 fractionated into a total of 96 fractions, which were consolidated into 24, from which 12 non-
713 adjacent samples were analyzed (Paulo et al., 2016a). Samples were subsequently acidified with
714 1% formic acid and vacuum centrifuged to near dryness. Each consolidated fraction was
715 desalted via StageTip, dried again via vacuum centrifugation, and reconstituted in 5%
716 acetonitrile, 5% formic acid for LC-MS/MS processing.

717

718 **Liquid chromatography and tandem mass spectrometry**

719 Mass spectrometric data were collected on an Orbitrap Fusion Lumos mass spectrometer
720 coupled to a Proxeon NanoLC-1200 UHPLC. The 100 μ m capillary column was packed with 35
721 cm of Accucore 50 resin (2.6 μ m, 150 \AA ; ThermoFisher Scientific). The scan sequence began with
722 an MS1 spectrum (Orbitrap analysis, resolution 120,000, 350–1400 Th, automatic gain control
723 (AGC) target 5E5, maximum injection time 50 ms). SPS-MS3 analysis was used to reduce ion
724 interference (Gygi et al., 2019; Paulo et al., 2016b). The top 10 precursors were then selected
725 for MS2/MS3 analysis. MS2 analysis consisted of collision-induced dissociation (CID),
726 quadrupole ion trap analysis, automatic gain control (AGC) 1E4, NCE (normalized collision
727 energy) 35, q -value < 0.25, maximum injection time 60 ms), and isolation window at 0.5.
728 Following acquisition of each MS2 spectrum, we collected an MS3 spectrum in which multiple
729 MS2 fragment ions are captured in the MS3 precursor population using isolation waveforms
730 with multiple frequency notches. MS3 precursors were fragmented by HCD and analyzed using
731 the Orbitrap (NCE 65, AGC 3E5, maximum injection time 150 ms, resolution was 50,000 at 400
732 Th).

733

734 **Mass spectra data analysis**

735 Mass spectra were processed using a Sequest-based pipeline (Huttlin et al., 2010). Spectra were
736 converted to mzXML using a modified version of ReAdW.exe. Database search included all
737 entries from an indexed Ensembl database version 90 (downloaded:10/09/2017). This database
738 was concatenated with one composed of all protein sequences in the reversed order. Searches
739 were performed using a 50 ppm precursor ion tolerance for total protein level analysis. The
740 product ion tolerance was set to 0.9 Da. TMT tags on lysine residues, peptide N termini
741 (+229.163 Da), and carbamidomethylation of cysteine residues (+57.021 Da) were set as static
742 modifications, while oxidation of methionine residues (+15.995 Da) was set as a variable
743 modification.

744 Peptide-spectrum matches (PSMs) were adjusted to FDR < 0.01 (Elias and Gygi, 2007, 2010).
745 PSM filtering was performed using a linear discriminant analysis (LDA), as described previously
746 (Huttlin et al., 2010), while considering the following parameters: XCorr, ΔCn, missed cleavages,
747 peptide length, charge state, and precursor mass accuracy. For TMT-based reporter ion
748 quantitation, we extracted the summed signal-to-noise (S:N) ratio for each TMT channel and
749 found the closest matching centroid to the expected mass of the TMT reporter ion. For protein-
750 level comparisons, PSMs were identified, quantified, and collapsed to a peptide FDR < 0.01 and
751 then collapsed further to a final protein-level FDR < 0.01, which resulted in a final peptide level
752 FDR < 0.001. Moreover, protein assembly was guided by principles of parsimony to produce the
753 smallest set of proteins necessary to account for all observed peptides. PSMs with poor quality,
754 MS3 spectra with TMT reporter summed signal-to-noise of less than 100, or having no MS3
755 spectra were excluded from quantification (McAlister et al., 2012).

756

757 **Filtering out peptides that contain polymorphisms**

758 Peptides that contain polymorphisms are problematic for protein quantification in genetically
759 diverse samples because as the reference genome, only the B6 allele is quantified.
760 Polymorphisms (with respect to the B6 genome) function as flags of the presence or absence of
761 the B6 allele rather than reflecting the relative abundance of the peptide. During protein
762 abundance estimation from peptides, a polymorphic peptide can either obscure the signal of a
763 true abundance pQTL or induce a false local abundance pQTL. To avoid these biases, we filtered
764 out peptides that contained polymorphisms based on the genome sequence of the founder
765 strains and that were further confirmed in the data by having local founder haplotype effects
766 that matched the expected distribution pattern of the B6 allele among the founder strains.

767 To determine whether peptides with polymorphisms matched their expected B6 allele
768 distribution patterns, the peptide data was made more comparable by standardization within
769 batches and removal of batch effects. Each peptide was scaled by a sample-specific within-

770 batch scaling factor: $\tilde{y}_i^{\text{pep } k} = \frac{y_i^{\text{pep } k}}{\theta_i}$, where $y_i^{\text{pep } k}$ is the mass-spec intensity of peptide k for

771 mouse i , $\theta_i = \frac{\sum_K y_i^{\text{pep } k}}{\max_{l \in B[i]} (\sum_K y_l^{\text{pep } k})}$, K is the set of all peptides measured for mouse i , and $B[i]$ is the

772 set of samples included in the mass-spec batch of mouse i . For the CC samples, the additional
773 pooled sample (*i.e.*, bridge sample) was also included in each batch and provided an additional

774 standardization across batches: $\tilde{\tilde{y}}_i^{\text{pep } k} = \log_2 \left(\frac{\tilde{y}_i^{\text{pep } k} + 1}{\tilde{y}_{b[i]}^{\text{pep } k} + 1} \right)$, where $b[i]$ represents the bridge

775 sample from the batch of mouse i . For the DO and founder strain samples that did not include

776 bridge samples, $\tilde{\tilde{y}}_i^{\text{pep } k} = \log_2 (\tilde{y}_i^{\text{pep } k} + 1)$. A log transformation is used because peptide

777 intensities are commonly log-linear.

778 Batch effects were removed from the processed peptide data using a linear mixed effect model
779 (LMM) fit with the lme4 R package (Bates et al., 2015). Peptides unobserved for all samples
780 within a batch were recorded as missing (coded as NA). If greater than 80% of samples were
781 missing for a polymorphic peptide, it was removed from the batch correction step and the

782 subsequent evaluation of the B6 allele distribution pattern. The following model was fit for the
 783 CC data:
 784 Equation 1

$$785 \quad \tilde{y}_i^{\text{pep } k} = \mu + \mathbf{x}_{i, \text{covar}}^T \boldsymbol{\beta}_{\text{covar}} + u_{\text{strain}[i]} + u_{b[i]} + \varepsilon_i$$

786
 787 where μ is the intercept, $\boldsymbol{\beta}_{\text{covar}}$ is the effect vector of covariates estimated as fixed effects,
 788 $\mathbf{x}_{i, \text{covar}}^T$ is the i^{th} row of the covariate design matrix, $u_{\text{strain}[i]}$ is the effect of the strain of sample
 789 i , $u_{b[i]}$ is the effect the batch of sample i , and ε_i is the error for sample i with $\varepsilon_i \sim N(0, \sigma^2)$. The
 790 strain and batch effects were estimated as random effects: $\mathbf{u}_{\text{strain}} \sim N(\mathbf{0}, \mathbf{I}\tau_{\text{strain}}^2)$ and
 791 $\mathbf{u}_b \sim N(\mathbf{0}, \mathbf{I}\tau_b^2)$. For the CC and founder strains, sex was included as a covariate. A similar model
 792 was fit for the DO but with no strain effect and diet also included as a covariate with sex. The
 793 batch effects, estimated with best linear unbiased predictors (BLUPs) using restricted maximum
 794 likelihood estimates (REML; Patterson and Thompson 1971), were subtracted from each
 795 peptide measurement: $\tilde{\tilde{y}}_i^{\text{pep } k} = \tilde{y}_i^{\text{pep } k} - \hat{u}_{b[i]}$.

796 For peptides expected to contain a polymorphism, we then fit local genetic effects: haplotype
 797 effects at the marker closest to the TSS of the gene to which the peptide maps in the CC and DO
 798 and strain effects in the founder strains.
 799 Equation 2

$$800 \quad \tilde{\tilde{y}}_i^{\text{pep } k} = \mu + \text{local}[i] + \mathbf{x}_{i, \text{covar}}^T \boldsymbol{\beta}_{\text{covar}} + u_i^{\text{kinship}} + \varepsilon_i$$

801
 802 where local_i is the effect of the local haplotype pair to peptide k for sample i , u_i^{kinship}
 803 represents a random effect that accounts for the correlation structure between individuals that
 804 is consistent with overall genetic relatedness, often referred to as the kinship effect, and all
 805 other terms as previously defined. For the CC, $\text{local}[i] = \mathbf{p}_i^T \boldsymbol{\beta}_{\text{local}}$ where \mathbf{p}_i^T is the founder
 806 haplotype probability vector at the marker closest to the gene TSS (e.g., ordering the founder
 807 strains as AJ, B6, 129, NOD, NZO, CAST, PWK, and WSB, $\mathbf{p}_i^T = [0 \ 1 \ 0 \ 0 \ 0 \ 0 \ 0 \ 0]$ for a CC mouse i
 808 that is B6/B6 at the locus). For the DO, $\text{local}[i] = \mathbf{d}_i^T \boldsymbol{\beta}_{\text{local}}$ where \mathbf{d}_i^T is the founder haplotype
 809 dosage vector, scaled to sum to zero, at the marker closest to the gene TSS (e.g., $\mathbf{d}_i^T =$
 810 $[0.5 \ 0.5 \ 0 \ 0 \ 0 \ 0 \ 0 \ 0]$ for a DO mouse i that is AJ/B6 at the locus). For the founder strains,
 811 $\text{local}_i = \mathbf{x}_{i, \text{strain}}^T \boldsymbol{\beta}_{\text{local}}$ where $\mathbf{x}_{i, \text{strain}}^T$ is the founder strain incidence vector for mouse i (e.g.,
 812 $\mathbf{x}_{i, \text{strain}}^T = [0 \ 1 \ 0 \ 0 \ 0 \ 0 \ 0 \ 0]$ for a B6 mouse). $\boldsymbol{\beta}_{\text{local}}$ is an eight-element vector of founder
 813 haplotype effects, fit as a random effect: $\boldsymbol{\beta}_{\text{local}} \sim N(\mathbf{0}, \mathbf{I}\tau_{\text{local}}^2)$ where \mathbf{I} is an 8×8 identity matrix
 814 and τ_{local}^2 is the variance component underlying the local effects. The kinship effect is included
 815 for the CC and DO and modeled as $\mathbf{u}^{\text{kinship}} \sim N(\mathbf{0}, \mathbf{G}\tau_G^2)$ where \mathbf{G} is a realized genomic
 816 relationship matrix and τ_G^2 is the variance component underlying the kinship effect, accounting
 817 for population structure (Kang et al., 2008, 2010; Lippert et al., 2011; Zhou and Stephens,
 818 2012). Here we used a leave-one-chromosome-out or “loco” \mathbf{G} , in which markers from the
 819 chromosome the peptide is predicted to be located on are excluded from \mathbf{G} estimation in order
 820 to avoid the kinship term absorbing some of $\text{local}[i]$ (Wei and Xu, 2016). We then calculated
 821 $r_{\text{poly}} = \text{cor}(\hat{\boldsymbol{\beta}}_{\text{local}}, \mathbf{q})$, the Pearson correlation coefficient between $\hat{\boldsymbol{\beta}}_{\text{local}}$, the BLUP of $\boldsymbol{\beta}_{\text{local}}$ and

822 \mathbf{q} , the incidence vector of the B6 allele among the founder strains (*e.g.*, $\mathbf{q} = [0\ 1\ 0\ 0\ 0\ 0\ 0]$)
 823 for a peptide with a B6 allele that is missing in the other founder strains). Sets of peptides with
 824 polymorphisms were defined based on having $r_{\text{poly}} > 0.5$ for each of the CC, DO, and founder
 825 strains, to be excluded from further analysis because they would bias protein abundance
 826 estimation.

827

828 Protein abundance estimation from peptides

829 Protein abundances were estimated from their component peptides after filtering out peptides
 830 that possessed polymorphisms based on founder strain sequences that were confirmed in the
 831 peptide data. The abundance for protein j is calculated as $y_i^{\text{prot } j} = \frac{\sum_M y_i^{\text{pep } m} 1_{i,m}}{\theta_i}$ where M is the
 832 set of peptides that map to protein j , $1_{i,m}$ is the indicator function that peptide m was
 833 observed in mouse i , and θ_i is the scaling factor previously defined (Huttlin et al., 2010). Similar
 834 to the previously described peptide normalization in the CC, proteins were scaled relative to the
 835 bridge sample (pooled sample of all CC mice included in all batches) and log-transformed:

836 $\tilde{y}_i^{\text{prot } j} = \log_2 \left(\frac{y_i^{\text{prot } j} + 1}{y_{b[i]}^{\text{prot } j} + 1} \right)$. For the DO and founder strain samples, there was no bridge sample,

837 and proteins were instead normalized as: $\tilde{y}_i^{\text{prot } j} = \log_2(y_i^{\text{prot } j} + 1)$.

838 Batch effects were removed from the protein data using the same LMM approach described for
 839 the peptide data (Equation 1). If more than 50% of samples were missing a protein, it was
 840 removed from further analysis. Batch effects, estimated as BLUPs, were then removed:

841 $\tilde{\tilde{y}}_i^{\text{prot } j} = \tilde{y}_i^{\text{prot } j} - \hat{u}_{b[i]}$.

842

843 Heritability estimation

844 We estimated heritability for all proteins in the CC, DO, and founder strains. The heritability
 845 model is similar to Equation 2, but for proteins instead of peptides and without the local[i]
 846 term:

847 Equation 3

$$848 \tilde{\tilde{y}}_i^{\text{prot } j} = \mu + \mathbf{x}_{i, \text{covar}}^T \boldsymbol{\beta}_{\text{covar}} + u_i^{\text{kinship}} + \varepsilon_i$$

849

850 where terms are as previously defined. The genomic relationship matrix \mathbf{G} – corresponding to
 851 the kinship term $\mathbf{u}^{\text{kinship}} \sim N(\mathbf{0}, \mathbf{G}\tau_G^2)$ for the CC and DO – is estimated from all markers –“non-
 852 loco” \mathbf{G} – because there are no other genetic terms in the model. In the founder strains, $\mathbf{G} =$
 853 $\mathbf{X}_{\text{strain}} \mathbf{X}_{\text{strain}}^T$ where $\mathbf{X}_{\text{strain}}$ is the founder strain incidence matrix. Sex was modeled as a
 854 covariate for all three populations, and diet as well in the DO. Heritability is then calculated as

$$855 h^2 = \frac{\tau_G^2}{\tau_G^2 + \sigma^2}.$$

856

857 QTL analysis

858 In the CC and DO, we performed a genome-wide pQTL scan for each protein, testing a QTL
 859 effect at positions across the genome, using a model similar to Equation 2:

860 Equation 4

$$861 \quad \tilde{z}_i^{\text{prot } j} = \mu + \text{QTL}_m[i] + \mathbf{x}_{i, \text{covar}}^T \boldsymbol{\beta}_{\text{covar}} + u_i^{\text{kinship}} + \varepsilon_i$$

862

863 where $\tilde{z}_i^{\text{prot } j}$ is the standard normal quantile returned by the inverse cumulative distribution
864 function of the normal distribution on the uniform percentiles defined by the ranks of $\tilde{\mathbf{y}}^{\text{prot } j}$,
865 *i.e.*, the rank-based inverse normal transformation (RINT) (Beasley et al., 2009) of protein j for
866 individual i , $\text{QTL}_m[i]$ is the effect of the putative QTL at marker m on protein j for individual i ,
867 equivalent to the $\text{local}[i]$ term in Equation 2 for the CC and DO, and all other terms as
868 previously defined. The kinship effect was fit based on the “loco” \mathbf{G} specific to the chromosome
869 of marker m . We used RINT for the QTL analysis to reduce the influence of extreme
870 observations that can produce false positives, particularly when they coincide with a rare
871 founder haplotype allele, which are of particular concern in the context of a CC sample
872 population with 58 unique genomes. To test the QTL term, the model in Equation 4 is compared
873 to a null model excluding QTL_m , summarized as the \log_{10} likelihood ratio (LOD) score.

874 The QTL model in Equation 4 was also used for variant association mapping at specific pQTL
875 identified through the haplotype-based analysis by adjusting the $\text{QTL}_m[i]$ term: $\text{QTL}_v[i] =$
876 $\mathbf{p}_{i,v}^T \boldsymbol{\beta}_{\text{QTL}}$, where $\mathbf{p}_{i,v}^T$ is the marginal variant allele probability vector for variant v , which is
877 calculated by collapsing and simplifying the underlying founder haplotype probabilities based
878 on the known variant genotype in the founder strains (SQLite variant database:
879 <https://doi.org/10.6084/m9.figshare.5280229.v3>).

880 For the CC, we mapped pQTL based on strain averages where $\tilde{z}_i^{\text{prot } j}$ is the average
881 of $\tilde{\mathbf{y}}_{\text{male, strain } i}^{\text{prot } j}$ and $\tilde{\mathbf{y}}_{\text{female, strain } i}^{\text{prot } j}$ followed by RINT across the population of strains. Founder
882 haplotype probabilities were reconstructed at the level of individual mice and were also
883 averaged at all markers for strain-level mapping. A strain-level \mathbf{G} was then estimated from the
884 strain-level founder haplotype probabilities. No fixed effect covariates were included when
885 mapping on strain averages. We tried mapping pQTL in the CC on individual-level data, which
886 returned largely consistent results, but notably fewer and weaker pQTL. In the CC, we also
887 mapped pQTL to the mitochondrial genome and Y chromosome by testing whether the founder
888 origin of the mitochondria or Y chromosome was associated with protein abundance. We fit
889 Equation 4, treating the mitochondrial genome or Y chromosome as a single locus $\text{QTL}_Y[i]$ and
890 $\text{QTL}_{\text{MT}}[i]$, respectively, using the “non-loco” \mathbf{G} for the kinship effect. The founder strain of
891 origin for the Y chromosome was determined for all CC strains. For the mitochondrial genome,
892 six strains (CC031, CC032, CC041, CC051, CC059, CC072) possessed ambiguity between AJ and
893 NOD, which we encoded as equal probabilities ($\mathbf{p}_{i,\text{MT}}^T = [0.5 \ 0 \ 0 \ 0.5 \ 0 \ 0 \ 0]$).

894

895 QTL significance thresholds

896 We estimated significance thresholds for pQTL using permutations (Doerge and Churchill,
897 1996). We accounted for varying levels of missing data by performing genome scans on 10,000
898 permutations of the normal quantiles for each level of observed missingness in the CC and DO
899 (ranging from 0 to 50%). Genome scans of the permuted data used the model in Equation 4,
900 while excluding covariates and the kinship term, allowing permutations to be more applicable

901 across proteins with the same level of missingness. To control the genome-wide error rate per
 902 protein and the FDR across proteins (Chesler et al., 2005), the maximum LOD scores from the
 903 permutation scans were used to fit generalized extreme value distributions (GEV) (Dudbridge
 904 and Koeleman, 2004; Valdar et al., 2009) specific to the level of missingness, which were used
 905 to calculate genome-wide permutation p -values for the maximum LOD observed per protein:

906 *Equation 5*

$$p_{\text{perm}}^{\text{prot } j} = 1 - F_{\text{GEV}, n_{\text{NA}}[\text{prot } j]}(\text{max LOD}[\text{prot } j])$$

907
 908 where $F_{\text{GEV}, n_{\text{NA}}[\text{prot } j]}$ is the cumulative density function for the GEV fit from the permutations
 909 of quantiles with n_{NA} number missing values, corresponding to the number missing for protein
 910 j , and $\text{max LOD}[\text{prot } j]$ is the maximum LOD score from the genome scan of protein j . We then
 911 used the Benjamini-Hochberg (BH) procedure (Benjamini and Hochberg, 1995) to calculate FDR
 912 q -values from all observed permutation p -values, which were used to interpolate a permutation
 913 p -value that corresponds to $\text{FDR} < \alpha$: $p_{\text{perm}, \alpha}^{\text{interp}}$ where $\alpha \in [0.1, 0.5]$. Significance thresholds on
 914 the LOD scale, specific to $\text{FDR} < \alpha$ and n_{NA} missing data points, were calculated:

915 $\lambda_{\text{FDR} < \alpha}^{n_{\text{NA}}} = F_{\text{GEV}, n_{\text{NA}}}^{-1}(1 - p_{\text{perm}, \alpha}^{\text{interp}})$ where $F_{\text{GEV}, n_{\text{NA}}}^{-1}$ is the inverse cumulative density function
 916 for the GEV with n_{NA} missing data points. As a final step to reduce random variation between
 917 sets of permutations, we regressed the estimated thresholds for a population and FDR level on
 918 the number of missing data points n_{NA} , and created a table of fitted thresholds: $\hat{\lambda}_{\text{FDR} < \alpha}^{n_{\text{NA}}}$ for $\alpha \in$
 919 $[0.1, 0.5]$ for both the CC and DO. Whether a pQTL met $\text{FDR} < \alpha$ significance, the threshold
 920 corresponding to α with the n_{NA} for protein j was used. For reference, $\hat{\lambda}_{\text{FDR} < 0.1}^0 = 7.89$ and
 921 $\hat{\lambda}_{\text{FDR} < 0.5}^0 = 6.30$ in the CC, and $\hat{\lambda}_{\text{FDR} < 0.1}^0 = 7.87$ and $\hat{\lambda}_{\text{FDR} < 0.5}^0 = 6.44$ in the DO.

923

924 **Consistency of QTL between the CC and DO**

925 We evaluated the consistency of pQTL between the CC and DO by comparing their haplotype
 926 effects. Haplotype effects were estimated at the pQTL marker using the model in Equation 4. To
 927 stabilize the effects, they were modeled as a random effect: $\beta_{\text{QTL}} \sim N(\mathbf{0}, \mathbf{I}\tau_{\text{QTL}}^2)$, where τ_{QTL}^2 is a
 928 variance component underlying the haplotype effects of the pQTL. We then estimated the
 929 haplotype effects as BLUPs ($\tilde{\beta}_{\text{QTL}}$). To declare pQTL consistent between the CC and DO, we
 930 evaluated whether their haplotype effects were significantly positively correlated: $p_{\text{QTL}}^r =$
 931 $\Pr(r_{\text{QTL}} > 0)$ where $r_{\text{QTL}} = \text{cor}(\tilde{\beta}_{\text{QTL}}^{\text{CC}}, \tilde{\beta}_{\text{QTL}}^{\text{DO}})$ and $r_{\text{QTL}}\sqrt{6}(1 - r_{\text{QTL}}^2)^{-1} \sim t_{(6)}$. To account for
 932 multiple testing, we used the BH procedure on the p -values for correlated effects and declared
 933 pQTL with $q_{\text{QTL}}^r < 0.1$ as consistent between the CC and DO.

934 Selecting which marker to fit the haplotype effects at is complicated by the fact that the CC and
 935 DO have different sets of markers and the genomic coordinate of the peak LOD score is also
 936 subject to variation. When comparing pQTL detected in both populations, we fit Equation 4 at
 937 the markers with the highest LOD score specific to each population, meaning they may not be
 938 the markers closest to each other. When comparing pQTL that were detected in only one
 939 population, we selected the marker in the population that failed to map the pQTL that was
 940 closest to the marker in the population that detected it.

941

942 Consistency of local QTL in the CC with the founder strains

943 If the genetic effects on a protein are primarily local, the relative abundances for a protein in
944 the founder strains should match the local pQTL effects observed in the CC and DO, which can
945 be used to confirm findings and better support suggestive pQTL in the CC or DO. We evaluated
946 the consistency of local pQTL in the CC with the founder strains, using an approach similar to
947 how we compared pQTL effects between the CC and DO. For the founder strains, rather than
948 fitting pQTL effects ($\tilde{\beta}_{\text{QTL}}$), we fit the founder effects as random terms (as described for the
949 local term in Equation 2 for the founder strains) summarized as BLUPs ($\tilde{\beta}_{\text{strain}}^{\text{Founders}}$). We then
950 calculated the Pearson correlation between local pQTL effects in the CC and founder effects in
951 the founder strains: $r_{\text{local}} = \text{cor}(\tilde{\beta}_{\text{QTL}}^{\text{CC}}, \tilde{\beta}_{\text{strain}}^{\text{Founders}})$. As when comparing QTL effects between the
952 CC and DO, we then tested $r_{\text{local}} > 0$, and corrected for multiple testing through the BH
953 procedure.

954

955 Mediation analysis

956 We performed mediation analysis on each distal pQTL with LOD > 6 in the CC or DO, which
957 involved a scan analogous to the QTL genome scans. The underlying model is

958 *Equation 5*

$$959 \quad \tilde{z}_i^{\text{prot } t} = \mu + \text{QTL}[i] + \mathbf{x}_{i, \text{covar}}^T \boldsymbol{\beta}_{\text{covar}} + \text{mediator}_q[i] + \varepsilon_i$$

960

961 where $\text{QTL}[i]$ is as defined for $\text{QTL}_m[i]$ in Equation 4, but fixed at the marker m of the
962 detected distal pQTL for target protein t , $\text{mediator}_q[i]$ is the effect of candidate mediator
963 protein q on protein t for individual i , and all other terms as previously defined. The effect of
964 the mediator is modeled as: $\text{mediator}_q[i] = \beta_{\text{prot } q} \tilde{z}_i^{\text{prot } q}$, where $\beta_{\text{prot } q}$ is the regression
965 coefficient for the mediator protein q and $\tilde{z}_i^{\text{prot } q}$ is the RINT quantity of protein q for individual
966 i . The likelihood of Equation 5 is compared to a null QTL model that excludes the QTL_i term,
967 producing a mediation LOD score. The mediation model is fit for all proteins as mediators,
968 excluding protein t , resulting in a mediation scan.

969 We assume that the vast majority of candidates are not true mediators of a specific pQTL and
970 thus the distribution of mediation LOD scores approximates a null distribution. Assuming that
971 the null distribution is approximately normal, we calculate the z-scores of the mediation LOD
972 scores, and define mediators of the QTL at marker m for protein t as proteins with $z_q^{\text{med}} < -4$,
973 where z_q^{med} is the z-score of the mediation LOD score for candidate mediator protein q . The
974 rationale being that when testing the QTL term in Equation 5, if the mediator contains some of
975 the information of the QTL, its presence in both the alternative and null models will result in a
976 large drop in the LOD score of the detected pQTL. In order for a mediator to be a clear
977 candidate driver of the distal pQTL, we required that the mediator TSS be within 10 Mbp of the
978 pQTL marker. Strong mediators that were not near the pQTL often represent proteins that are
979 correlated with protein t , such as co-regulated members of a protein-complex.

980

981 Sex effects on proteins analysis

982 Proteins that exhibited differential abundance between the sexes, *i.e.*, sex effects, were
983 identified using an LMM similar to the heritability model (Equation 3) for the CC, DO, and
984 founder strains, but instead testing the significance of the sex coefficient:

985 *Equation 6*

$$986 \tilde{y}_i^{\text{prot } j} = \mu + \beta_{\text{Male}} x_{i, \text{Male}} + \mathbf{x}_{i, \text{covar}}^T \boldsymbol{\beta}_{\text{covar}} + u_i^{\text{kinship}} + \varepsilon_i$$

987

988 where β_{Male} is the effect on protein j of being male, $x_{i, \text{Male}}$ is an indicator variable of being
989 male, and all other terms as defined previously. Non-sex covariates and the specification of
990 u_i^{kinship} for the different populations are the same as described for heritability.

991 A p -value for the sex effect was calculated by comparing the model in Equation 6 to a null
992 model without the sex effect through the likelihood ratio test (LRT): $p_{\text{sex}}^{\text{prot } j} = \Pr(X > \hat{\chi}_{\text{prot } j}^2)$
993 where $\Pr(\cdot)$ denotes the $\chi_{(1)}^2$ probability density function and $\hat{\chi}_{\text{prot } j}^2$ is the observed LRT
994 statistic for protein j . The LMM was fit with the `qtl2` R package (Broman et al., 2019), using
995 maximum likelihood estimates (MLE) for parameters rather than REML, which are more
996 appropriate for asymptotic-based significance testing of fixed effects. Proteins with significant
997 sex effects were selected based on FDR < 0.1 using the BH procedure (Benjamini and Hochberg
998 1995).

999 We performed gene set enrichment analysis through the `clusterProfiler` R package (Yu et al.,
1000 2012). We defined gene sets based on having $q_{\text{sex}} < 0.01$ and split them further into subsets
1001 based on having higher abundance in males or higher abundance in females for both the CC
1002 and DO. We used the quantified proteins in each population as the background gene set to
1003 account for biases in the observed proteins. Hypergeometric set tests for enrichment of GO and
1004 KEGG terms were performed with FDR multiple testing control (Storey et al., 2019). Enriched
1005 GO and KEGG terms were selected based on having $q_{\text{set}} < 0.1$.

1006

1007 Protein-complex analysis

1008 We categorized individual proteins as members of protein-complexes as defined by previous
1009 annotations (Ori et al., 2016). For each protein-complex, we quantified how tightly co-
1010 abundant, *i.e.*, cohesive, the members are, by calculating the median pairwise Pearson
1011 correlation for each protein with the other members of the protein-complex. We summarized
1012 cohesiveness within a protein-complex by recording the median and interquartile range across
1013 the median correlations for the individual proteins.

1014 To assess whether genetics or sex regulated protein-complexes as a whole, we estimated the
1015 complex-heritability and complex-sex effect size. We took the PC1 from the set of proteins from
1016 the complex after first removing the effects of key covariates from the individual proteins in
1017 order to keep the PC1 from reflecting other drivers of variation instead of genetic factors or sex.
1018 For complex-heritability, we removed the effect of sex in the CC, and both sex and diet in the
1019 DO. For complex-sex effect size, we only removed the effect of diet from the DO. Complex-
1020 heritability was estimated using the model in Equation 3, with no covariates and the complex

1021 PC1 as the outcome. To estimate the complex-sex effect size: $\phi_{\text{sex}}^2 = 1 -$
1022 $(\sum_i e_i^2 | M_A) / (\sum_i e_i^2 | M_0)$ where $\sum_i e_i^2 | M_A$ is the sum of squared residuals (SSR) under the
1023 alternative model (Equation 6) and $\sum_i e_i^2 | M_0$ is the SSR under the null model (Equation 6
1024 excluding sex effect). Interval estimates for complex-heritability and complex-sex effect size
1025 represent 95% subsample intervals. We randomly sampled without replacement 80 of the CC
1026 and DO data 1,000 times and estimated the complex-heritability and complex-sex effect size for
1027 each subsample as well as the 2.5th and 97.5th quantiles across the subsamples. We estimated
1028 summaries for protein-complexes that had four or more proteins observed in the CC or DO,
1029 after removing proteins with local pQTL (FDR < 0.5) or distal pQTL (FDR < 0.1), limiting the
1030 potential that the PC1 reflects a strong pQTL not shared by other members of the complex.

1031

1032 **Strain-protein outlier analysis**

1033 To identify CC strains with consistent extreme effects in both the female and male, we fit an
1034 LMM:

1035 *Equation 8*

$$1036 \quad \tilde{y}_i^{\text{prot } j} = \mu + \beta_{\text{Male}} x_{i, \text{Male}} + u_{\text{strain}[i]} + \varepsilon_i$$

1037

1038 with all terms as previously defined. Effects for all CC strains for each protein j ($\hat{u}_{\text{strain}}^{\text{prot } j}$) were
1039 estimated as BLUPs, which were then transformed to z-scores per protein ($z_{\text{strain}}^{\text{prot } j}$). We defined
1040 a strain-protein outlier to be a protein j in CC strain i for which $|z_{\text{strain } i}^{\text{prot } j}| > 2.5$. We intersected
1041 the strain-protein outliers with known CC strain-private genetic variants (Srivastava et al.,
1042 2017), identifying CC strain-private variants that likely have local effects on protein abundance.
1043 We permuted the pairings between CC strain and protein for the 6,046 strain-protein outliers
1044 to determine if the observed overlap of 69 privates was significant.

1045 For each CC strain i , we defined sets of proteins that had consistently high, low, and extreme
1046 abundance based on their strain effects: $\Omega_{\text{strain } i}^{\text{high}} = \{\text{prot } j: z_{\text{strain } i}^{\text{prot } j} > 2.5\} \forall j$, $\Omega_{\text{strain } i}^{\text{low}} =$
1047 $\{\text{prot } j: z_{\text{strain } i}^{\text{prot } j} < -2.5\} \forall j$, and $\Omega_{\text{strain } i}^{\text{extreme}} = \{\text{prot } j: |z_{\text{strain } i}^{\text{prot } j}| > 2.5\} \forall j$, respectively. We then
1048 tested whether the CC strain-specific sets were enriched in GO and KEGG terms ($q_{\text{set}} < 0.1$), as
1049 done with proteins with sex effects.

1050

1051 **Data and software availability**

1052 All analyses were performed using the R statistical programming language (v3.6.1) (R Core
1053 Team, 2018). The scripts and processed data used to generate the results are available at
1054 figshare (<https://doi.org/10.6084/m9.figshare.12818717>). All processed data and pQTL results
1055 are available for download and interactive analysis using the QTLViewer webtool for both the
1056 CC (<https://churchilllab.jax.org/qtlviewer/CC/Ferris>) and DO
1057 (<https://churchilllab.jax.org/qtlviewer/DO/Svenson>). The mass-spec proteomics data for the CC
1058 liver samples have been deposited in ProteomeXchange (<http://www.proteomexchange.org/>)
1059 via the PRIDE partner repository with dataset identifier PXD018886. The mass-spec data for the

1060 DO and founder strain liver samples were previously deposited to ProteomeXchange with
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1062

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1072

1073 **Author contributions**

1074 Conceptualization, M.T.F., S.P.G., and G.A.C.; Methodology, G.R.K., T.Z., S.P.G., and G.A.C.;
1075 Software, G.R.K, D.P., and M.V.; Investigation, G.R.K. and T.Z.; Resources, T.Z., T.A.B., P.H.,
1076 G.D.S., F.P-M.V., M.T.F., and S.P.G.; Data Curation, G.R.K, T.Z., and M.V.; Writing – Original
1077 Draft, G.R.K., T.Z., and G.A.C.; Writing – Review & Editing; Visualization, G.R.K.; Supervision,
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1079

1080 **Declarations of interests**

1081 The authors declare no competing interests.

1082

1083 **Supplemental Information**

1084 Table S1. Heritability and sex effects for the proteins analyzed in the CC, DO, and founder
1085 strains, related to **Figure 1**

1086

1087 Table S2. Detected pQTL (LOD > 6) in the CC and DO, related to **Figures 2 and S1**

1088

1089 Table S3. Consistency of haplotype effects for pQTL detected (FDR < 0.5) in the CC and DO,
1090 related to **Figures 2 and S1**

1091

1092 Table S4. Mediation results for distal pQTL detected (LOD > 6) in the CC and DO, related to
1093 **Figures 2 and S1**

1094

1095 Table S5. Annotated protein-complexes (Ori et al., 2016) used in this study, with added mouse
1096 ENSEMBL gene IDs and the numbers of member proteins observed in each mouse population,
1097 related to **Figures 3 and S4**

1098

1099 Table S6. Summaries on heritability, pQTL, and sex effect size for individual proteins annotated
1100 to protein-complexes, related to **Figures 3 and S4**

1101

- 1102 Table S7. Summaries of cohesiveness, heritability, and sex effect size for protein-complexes,
1103 related to **Figures 3** and **S4**
1104
- 1105 Table S8. Gene ontology enrichment results of proteins with extreme abundances specific to
1106 individual CC strains, related **Figures 7** and **S7**
1107
- 1108 Table S9. KEGG enrichment results of proteins with extreme abundances specific to individual
1109 CC strains, related **Figures 7** and **S7**
1110
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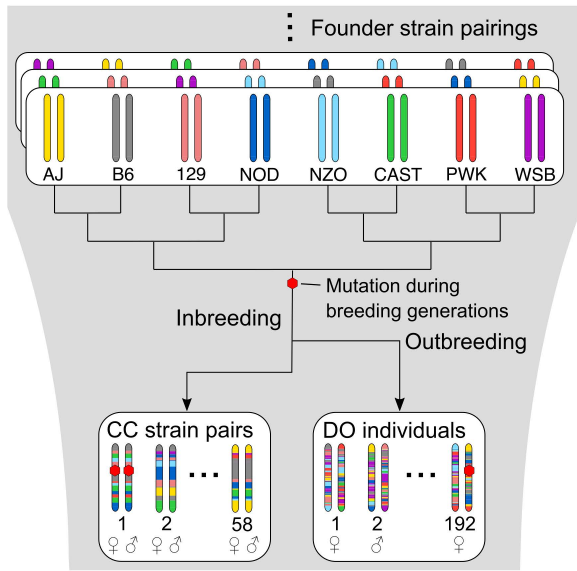
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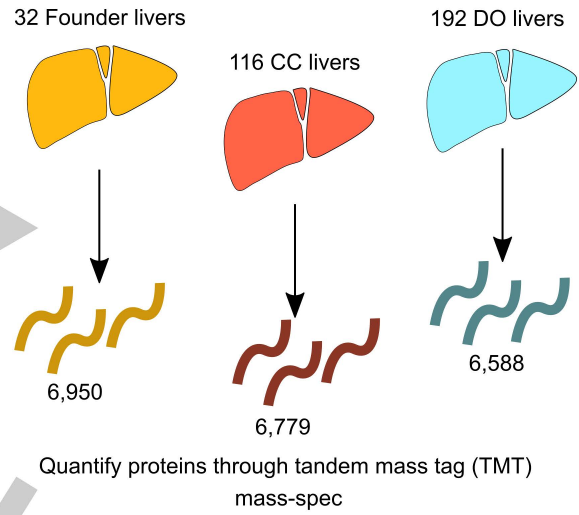
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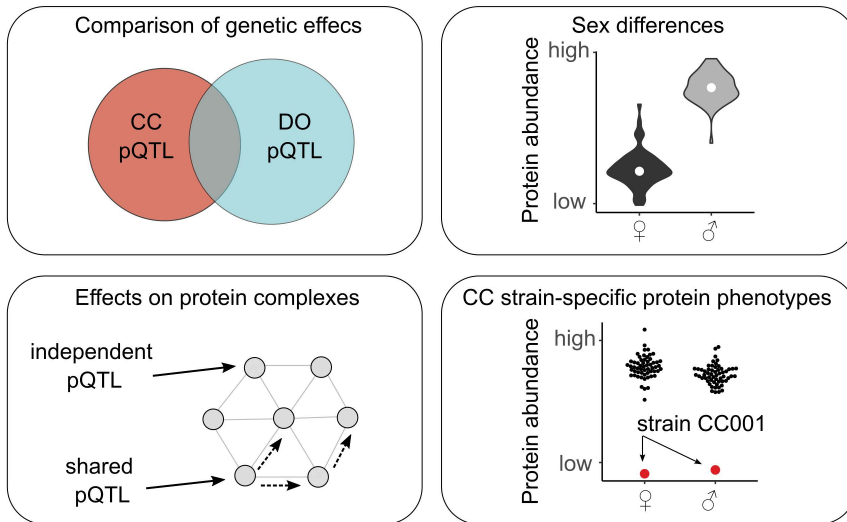
Genetically diverse mouse populations



Liver proteomics data

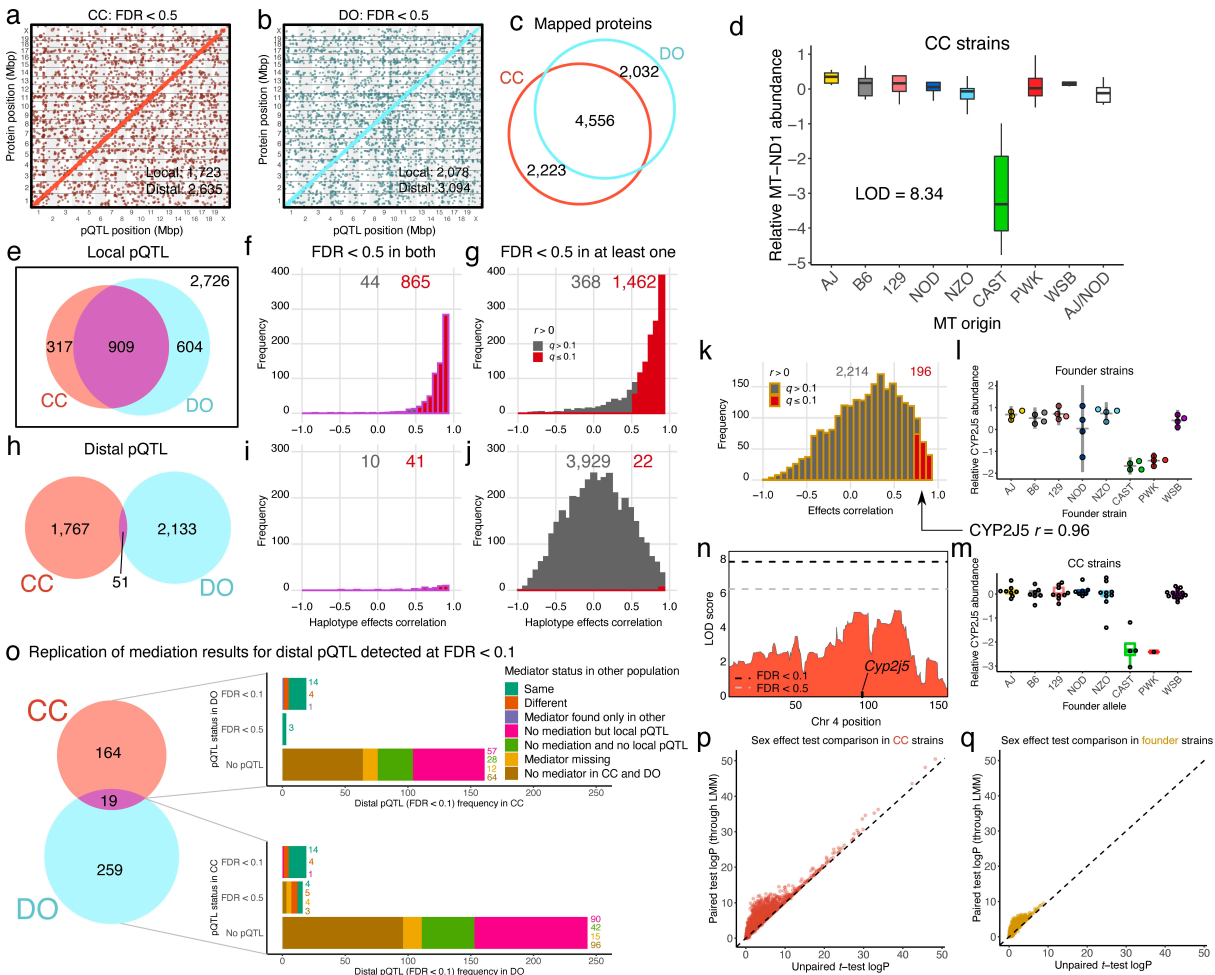


Proteomics/genetics analysis



Graphical abstract

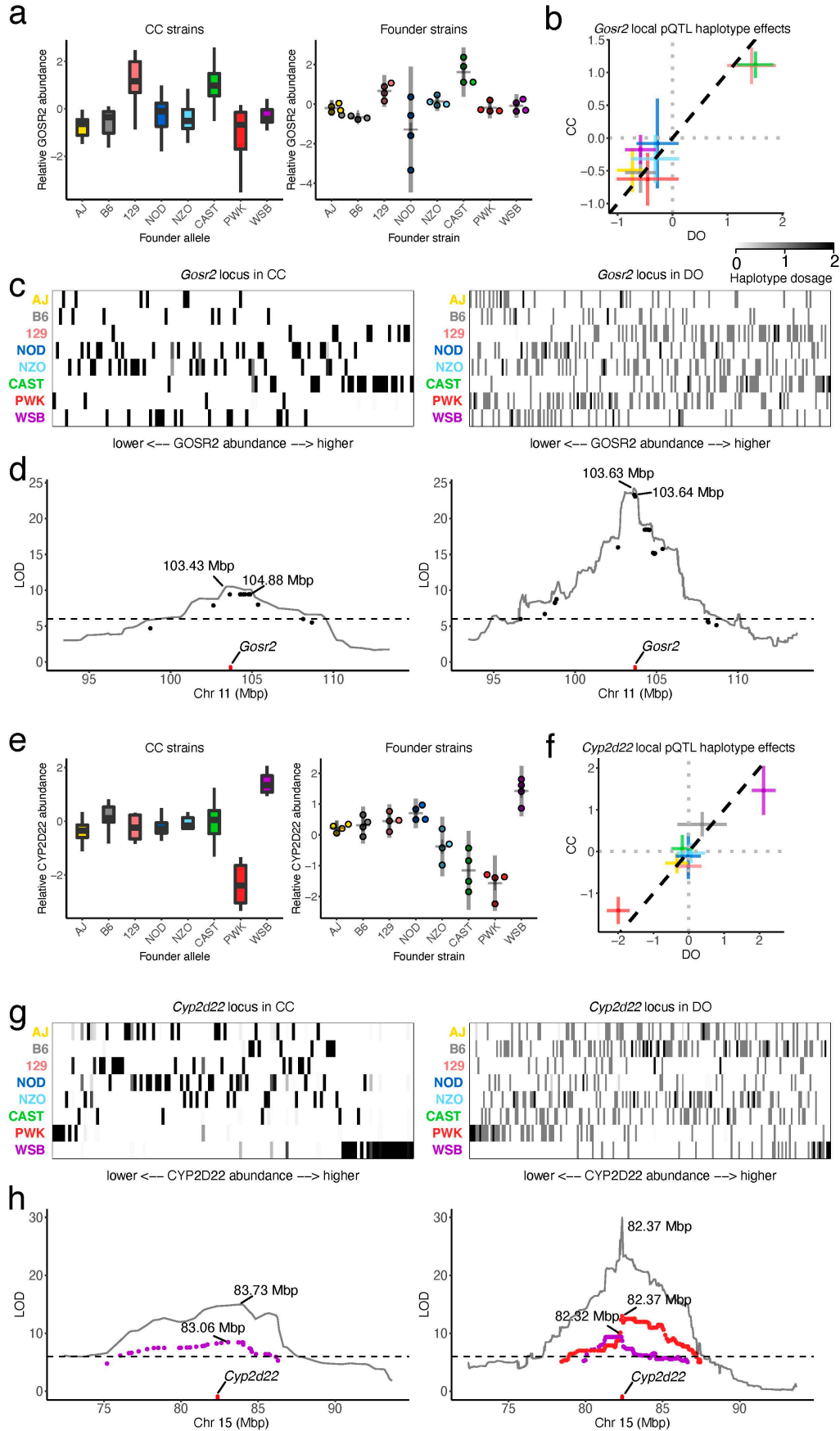
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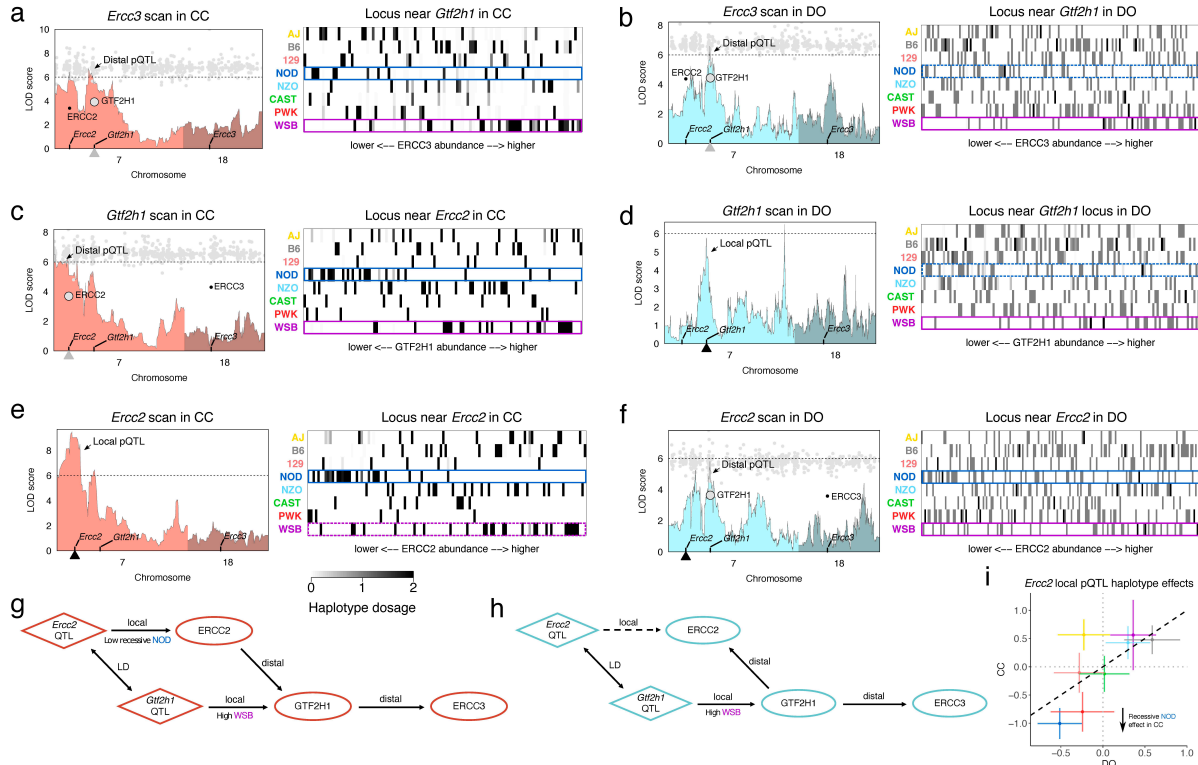
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Figure S1. Comparison of lenient mapping results, mediation candidates, and sex effects among genetically diverse mouse populations, related to Figures 1 and 2. Leniently detected (FDR < 0.5) pQTL are plotted by the genomic position of protein against their coordinate. Dot size is proportional to association strength (LOD score). (c) Venn diagram of the overlap of analyzed proteins between the CC and DO. (d) A local pQTL was detected in the CC for *mt-Nd1*, a gene encoded on the mitochondrial genome. The pQTL is characterized by low MT-ND1 in CC strains with the CAST mitochondrial genome. For six CC strains, mitochondrial inheritance was ambiguous between AJ and NOD (white boxplot). (e) Venn diagram of the overlap in local pQTL detected in the CC and DO. (f) The haplotype effects of local pQTL detected in both populations are highly consistent, as measured by the correlation coefficient comparing the effects in the CC and DO. (g) More local pQTL are consistent between the populations when also considering pQTL detected in only one of them. Red bars represent pQTL that had significantly correlated effects (FDR < 0.1). (h) Venn diagram of overlap of distal pQTL detected in the CC and DO. (i) Using lenient detection resulted in more distal pQTL in both populations with consistent haplotype effects (41 out of 51). (j) In contrast to local genetic effects, considering distal pQTL that were leniently detected in only one of the populations resulted in many inconsistent effects comparisons, likely representing false positives or subtle distal effects specific to one population. The founder strains can reveal under-powered local pQTL in the CC (and DO), as revealed by the correlations between haplotype effects at putative local pQTL for genes with rare alleles in the CC (≤ 3 CC strains) and that did not have a local pQTL detected (FDR < 0.5) and strain effects in mice from the founder strains (k). The enrichment in positive correlations suggests the founders can provide additional information for detecting local pQTL in the CC. The gene *Cyp2j5* had significant correlation ($r = 0.96$) between the strain effects in the (l) founder strains with the (m) local haplotype effects in the CC, characterized by low CAST and PWK effects. Mean and ± 2 standard deviation bars are shown for the founder strains, and boxplots for CC strains. Four CC strains possessed the CAST allele and only one had the PWK allele, resulting in (n) poor power to detect the local pQTL of *Cyp2j5*. Stringent and lenient significance thresholds are included as horizontal black and gray dashed lines, respectively. (o) Comparison of mediation results for distal pQTL stringently detected (FDR < 0.1) in at least one of the populations. For distal pQTL detected stringently in both populations, mediation results were strongly consistent (14 out of 19). If the distal pQTL were detected leniently in the other population, mediation was still somewhat consistent (7 out of 19). When the distal pQTL was not detected in the other population, often no mediator was detected, or the candidate mediator was missing from the other

1371 population. Observance of both sexes within the (p) CC and (q) founder strains improves power to detect sex effects. The -
1372 \log_{10} (p-value), or logP, from a linear mixed effect model (LMM) that takes into account strain replicates compared to the logP
1373 from an unpaired t-test. The lower logP in the founder strains reflects the overall smaller sample size (32 mice compared to 116
1374 in the CC). Dashed diagonal lines included for reference.

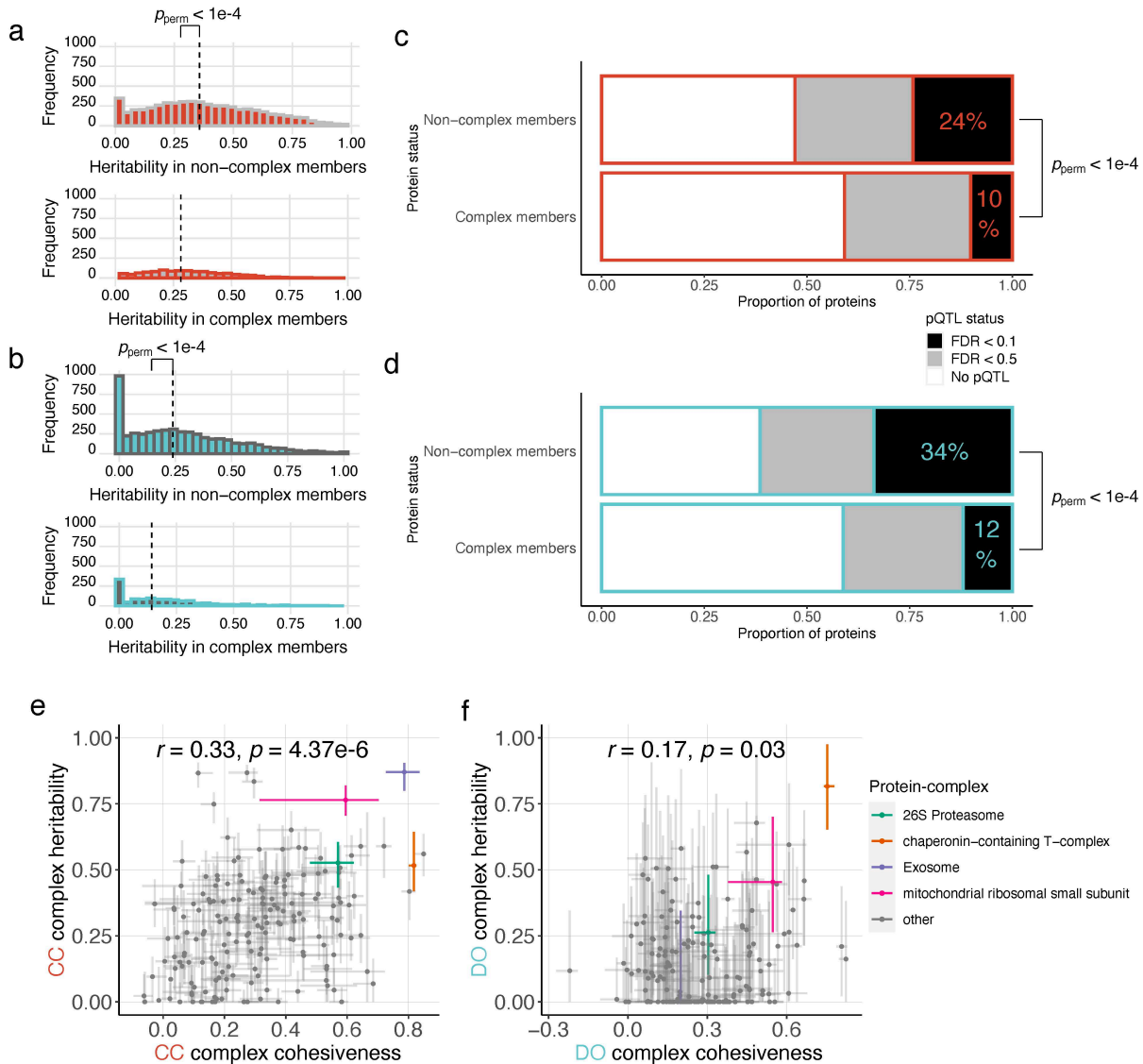


1376 **Figure S2. Highly consistent local pQTL effects across the CC, DO, and founder strains, related to Figure 2.** Local pQTL for (a)
1377 *Gosr2* and (e) *Cyp2d22* have highly consistent effects across the CC (left) and founder strains (right). Boxplots are shown for the
1378 CC strains and mean \pm 2 standard deviation bars for the founder strains. The local pQTL effects are also highly consistent
1379 between the CC and DO, both in terms of modeled effects and the actual data – for (b, c) *Gosr2* and (f, g) *Cyp2d22*. For
1380 comparisons of modeled effects (b, f), intervals represent \pm standard error bars. Dashed diagonal lines included for reference.
1381 To visualize the effects in the actual data (c, g), the CC and DO data at the pQTL are represented as heatmaps with rows
1382 indicating founder allele dosage at the local pQTL (expected allele counts) and columns indicating individual mice, ordered by
1383 protein abundance. Clusters in rows towards the left or right sides represent founder haplotype effects, such as the high WSB
1384 effect in *Cyp2d22*. Haplotype-based association scans at the pQTL are overlaid with variant associations for (d) *Gosr2* and (h)
1385 *Cyp2d22*. When the pQTL effects are approximately bi-allelic, as with *Gosr2* (d), the peak variant association and haplotype-
1386 based association are very close, consistent with a single variant largely driving the pQTL. Variants with an allele shared by only
1387 129 and CAST, matching the effects pattern, are shown. When the effects were more complex than bi-allelic, as with *Cyp2d22*
1388 (h), there are likely multiple causal variants, and haplotype-based association produces higher scores of association. WSB- and
1389 PWK-private variants with LOD > 6 were included for *Cyp2d22*, highlighting LD blocks that potentially carry founder-specific
1390 variants driving their extreme effects. The larger sample size and finer mapping resolution of the DO sample are evident in the
1391 high LOD scores and narrower association peaks. Genomic positions of peak associations from variant- and haplotype-based
1392 mapping are marked. Horizontal dashed lines at LOD of 6 included as reference.

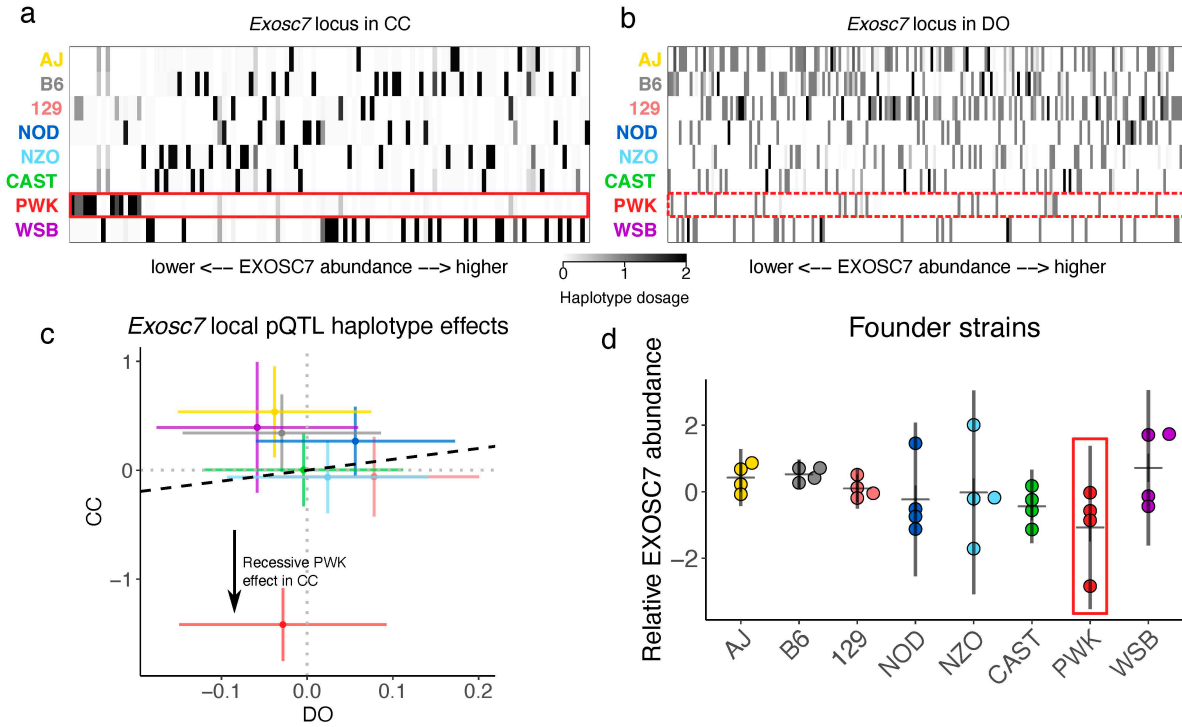


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Figure S3. Similarity and differences in the genetic effects on *Ercc3* between the CC and DO, related to Figure 2. Suggestive distal pQTL for *Ercc3* map to chromosome 7 in both the (a) CC and (b) DO, consisting of two peaks above *Ercc2* and *Gtf2h1*, genes that are approximately 30 Mbp apart and known to interact with ERCC3. Mediation analysis identified GTF2H1 and ERCC2 as candidate drivers of the *Ercc3* pQTL. Gray dots represent mediation scores for all quantified proteins on chromosomes 7 and 18. Horizontal dashed lines at LOD of 6 included for reference. To better understand the pQTL effects, the founder haplotype inheritance was plotted as heatmaps with founder allele dosage (expected allele counts) as rows and individual mice as columns, ordered by ERCC3 abundance, at the chromosome 7 locus near *Gtf2h1*. A low NOD effect was observed, indicated by the dark blue box, which was weak to non-existent in the DO (dotted dark blue box), as well as a high WSB effect in both the CC and DO (purple boxes). (c) In the CC, *Gtf2h1* has a weak distal pQTL that mapped nearby *Ercc2* and was mediated by ERCC2 with similar low NOD and high WSB effects. (d) In the DO, *Gtf2h1* has a suggestive local pQTL (LOD < 6) and notably no suggestive association near *Ercc2*. (e) *Ercc2* has a strong local pQTL in the CC (FDR < 0.1), driven by a low NOD effect, whereas in the (f) DO, a suggestive distal pQTL is observed near *Gtf2h1*. Diagrams for the relationships defined by pQTL and mediation in the (g) CC and (h) DO. In both populations, *Ercc2* and/or *Gtf2h1* affect ERCC3 abundance, and potentially each other as well. Teasing apart the directionality of effects is further complicated by the linkage disequilibrium (LD) between *Ercc2* and *Gtf2h1*. The low NOD effect at the *Ercc2* locus is stronger in the CC, and the high WSB effect is strongest at the *Gtf2h1* locus. (i) The haplotype effects at the locus near *Ercc2* are relatively similar, though the NOD effect is more extreme in the inbred CC background. Intervals represent \pm standard error bars. Dashed diagonal line included for reference.



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 1412 **Figure S4. Comparatively fewer detected genetic effects on protein-complex members, and the correlation between protein-**
 1413 **complex heritability and cohesiveness, related to Figures 3, 4, 5, and 6.** Histograms of the heritability of proteins that are not
 1414 annotated as members of a protein-complex (top) and that are members (bottom) in the (a) CC and (b) DO. Vertical dashed line
 1415 represents the median heritability, which were significantly different ($p_{perm} < 1e-4$). A lower proportion of proteins that are
 1416 members of protein-complexes possess pQTL ($p_{perm} < 1e-4$) in both the (c) CC and (d) DO. Black boxes represent stringently
 1417 detected (FDR < 0.1) pQTL, gray boxes represent leniently detected (FDR < 0.5) pQTL, and white boxes represent proteins that
 1418 had no pQTL detected. Numbers represent the percentage of proteins with detected pQTL at FDR < 0.1 within each category.
 1419 The correlation between protein-complex heritability and cohesiveness is stronger in the (e) CC ($r = 0.33, p = 4.37e-6$) than the
 1420 (f) DO ($r = 0.17, p = 0.03$). For protein-complex cohesiveness, points and bars represent medians and interquartile ranges,
 1421 respectively. Protein-complex heritability was estimated from the first principal component (PC1) of the complex members, and
 1422 bars represent 95% subsample intervals around the estimate. Exosome, chaperonin-containing T-complex, 26S Proteasome,
 1423 and the mitochondrial ribosomal small subunit are highlighted and are examined in detail (Figures 4, 5, 6, S5, & S6).



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Figure S5. A local recessive effect on EXOSC7 that impacts the whole exosome is exposed by the inbred genomes of the CC strains, related to Figure 4. The founder haplotype inheritance at the *Exosc7* local pQTL represented as a heatmap with founder allele dosages (expected allele counts) as rows and individuals as the columns, ordered by EXOSC7 abundance, for the (a) CC and (b) DO. The cluster on the left side in the PWK row (red box) for the CC reflects the low EXOSC7 abundance observed in CC mice that are homozygous for the PWK allele at the locus. No homozygotes for PWK were observed in the DO, and the heterozygous PWK carriers do not have low EXOSC7 abundance (dotted red box). (c) Comparison of the modeled allele effects at the *Exosc7* locus reveal a marked lower PWK effect specific to the CC, which is consistent with the effects of a recessive allele. Intervals represent \pm standard error bars. Dashed diagonal line included for reference. (d) Mice from the PWK strain with low EXOSC7 abundance were observed, though the effect is not as distinct as in the CC, possibly due to decreased accuracy without a bridge sample and the relative nature of the mass-spec quantification. Founder strain mice are summarized with mean \pm 2 standard deviation bars.

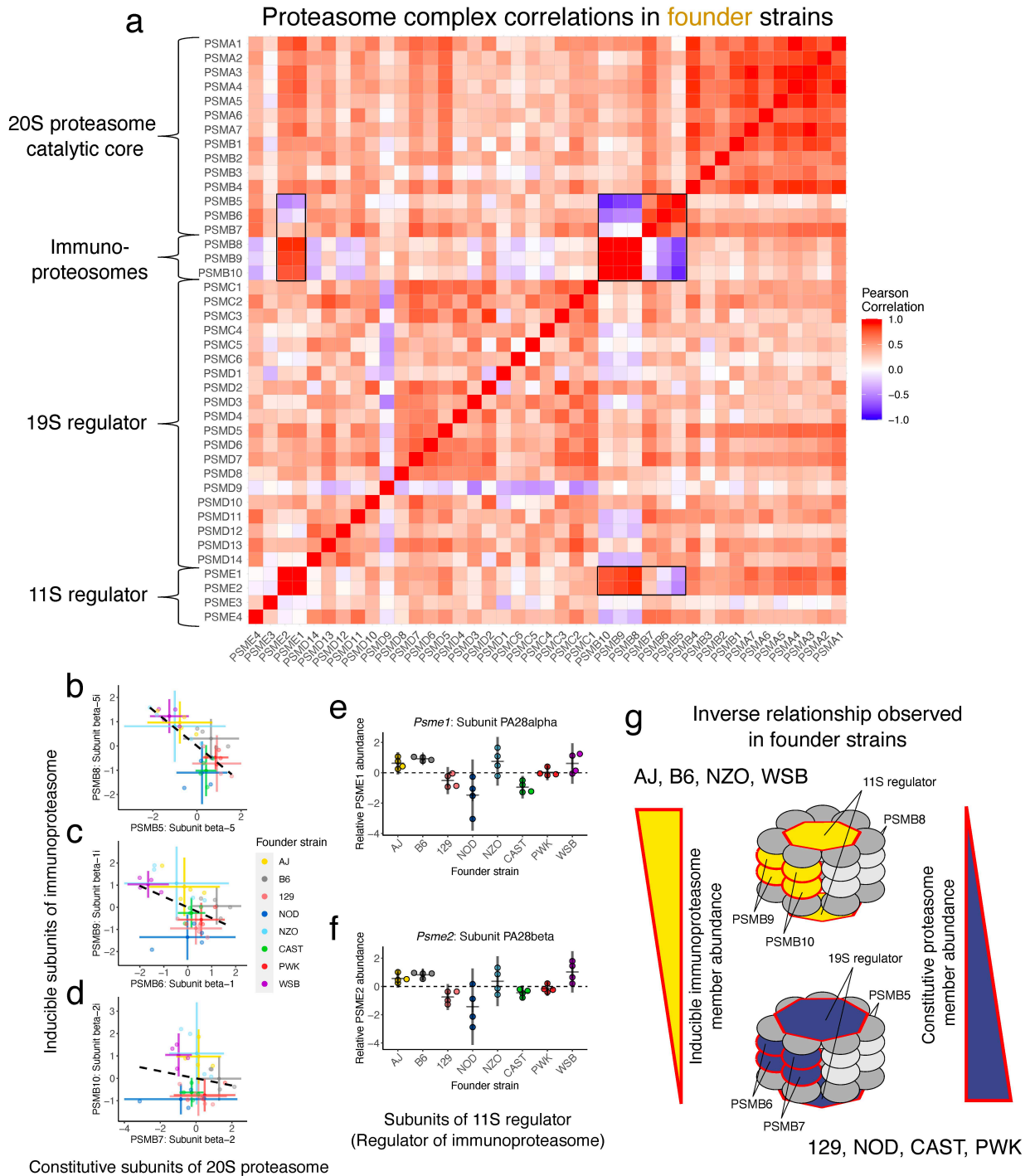
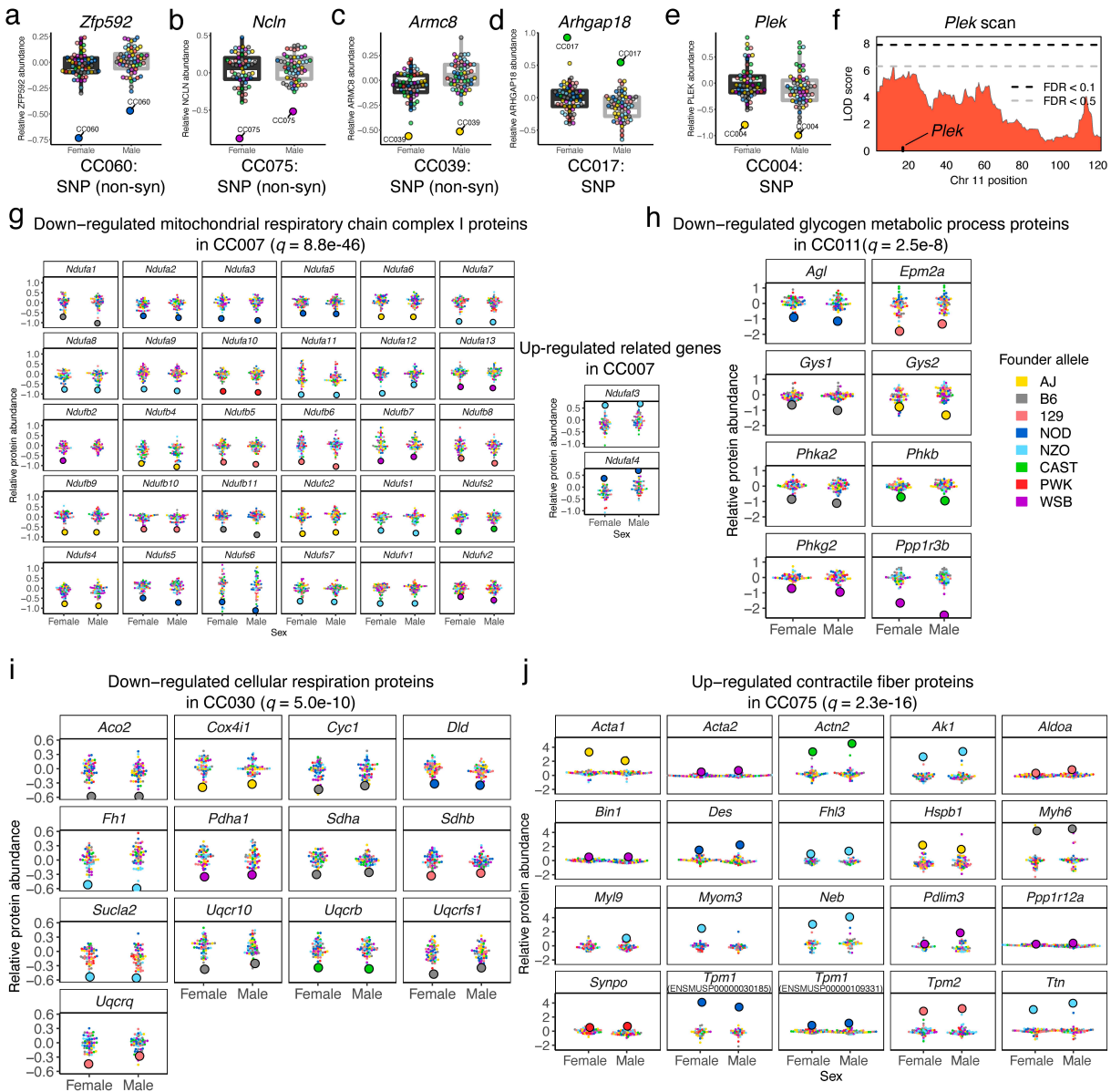


Figure S6. The founder strains differ in the balance between the constitutive proteasome and the inducible immunoproteasome, related to Figure 6. The correlation pattern among members of the 26S proteasome in the founder strains matches closely with those observed in the CC and DO (Figure 6a & b). The constitutive members of the 20S proteasome (PSMB5, PSMB6, and PSMB7) are anti-correlated with the corresponding inducible immunoproteasome components (PSMB8, PSMB9, and PSMB10) and the subunits from the 11S regulator – the regulator complex specific to the immunoproteasome – highlighted with black boxes. Founder strains with high abundance of the constitutive proteasome members have low levels of immunoproteasome proteins, and vice versa. The direction of this relationship depends on genetic factors, as revealed by the founder strains, with WSB and AJ mice consistently possessing relatively high abundance of immunoproteasome members and

1445 low abundance of the corresponding constitutive members: (b) PSMB8 vs PSMB5, (c) PSMB9 vs PSMB6, and (d) PSMB10 vs
1446 PSMB7. Intervals represent mean \pm 2 standard deviation bars. Dashed lines represent the best fit line between corresponding
1447 inducible and constitutive subunits. Two of the proteins from the 11S regulator of the immunoproteasome, (e) PSME1 and (f)
1448 PSME2, have matching abundance in AJ, B6, NZO, and WSB mice as in the inducible immunoproteasome components, which is
1449 largely consistent with joint regulation of constitutive/inducible members (g). The WSB mice are consistent with genetic
1450 variation identified through QTL mapping near *Psmb9* that drives PSMB9 and indirectly PSMB6 abundance relative to each
1451 other in the recombinant CC and DO populations (**Figure 6i-l**). Founder strain mice are summarized with mean \pm 2 standard
1452 deviation bars. Horizontal dashed lines provide reference at 0 for 11S regulator proteins.



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Figure S7. Proteins with extreme abundance in CC strains that possess unique local variants, and CC strains with extreme protein abundance patterns in functional pathways, related to Figure 7. Mutations became fixed in CC strains and were previously identified (Srivastava et al., 2017). These private variants of CC strains potentially have local effects on protein abundance. Examples include (a) *Zfp952*, (b) *Ncln*, (c) *Armc8*, (d) *Arhgap18*, and (e) *Plek*. *Zfp952*, *Ncln*, and *Armc8* each possess a non-synonymous SNP variant in a CC strain which had strikingly low protein abundance. These variants may cause reduced abundance, but alternatively, they may represent a coding polymorphism in the quantified peptides, resulting in false low abundance due to the allele-specific nature of mass-spec quantification. The color or the dots indicates the founder allele at the gene. Strain-specific variants associated with high abundance at the local protein were also observed, such as high ARHGAP18 abundance in CC017. (e) A novel SNP allele in CC004 at *Plek* was associated with low abundance, potentially representing a new allele at a gene that already possessed local genetic variation from the founder strains that drove a (f) suggestive local pQTL. Stringent and lenient significance thresholds are included as horizontal black and gray dashed lines, respectively. CC strain-specific sets of proteins with extreme abundance were identified that were significantly enriched in GO and KEGG functional terms. (g) CC007 possessed distinctly low abundance of mitochondrial respiratory chain complex I proteins, as well as high abundance in two assembly factor proteins from the complex, NDUFAF3 and NDUFAF4, suggesting compensatory mechanisms may be occurring at the complex in the strain. Large dots represent the CC strain with extreme protein abundance. Other detected CC strain-specific protein dynamics include low abundance of (h) glycogen metabolic process proteins for CC0011, (i) cellular respiration proteins for CC030, and (j) high abundance of contractile fiber proteins for CC075.