1 2 3	Genetic, epigenetic, and environmental mechanisms govern allele-specific gene expression
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32 33 34 35 36	Running Title: Metabolic tissues and environments influence allele-specific expression patterns
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#### 39 ABSTRACT

40 Allele-specific expression (ASE) is a phenomenon where one allele is preferentially expressed over the 41 other. Genetic and epigenetic factors cause ASE by altering the final allelic composition of a gene's 42 product, leading to expression imbalances that can have functional consequences on phenotypes. 43 Environmental signals also impact allele-specific gene regulation, but how they contribute to this crosstalk 44 remains understudied. Here, we explored how allelic genotype, parent-of-origin, tissue type, sex, and 45 dietary fat simultaneously influence ASE biases in a  $F_1$  reciprocal cross mouse model. Male and female 46 mice from a F<sub>1</sub> reciprocal cross of the LG/J and SM/J strains were fed a high-fat or low-fat diet. We 47 harnessed strain-specific variants to distinguish between two classes of ASE: parent-of-origin dependent 48 (unequal expression based on an allele's parental origin) and sequence dependent (unequal expression 49 based on an allele's nucleotide identity). We present a comprehensive genome-wide map of ASE 50 patterns across three metabolically-relevant tissues and nine environmental contexts. We find that both 51 ASE classes are highly dependent on tissue type and environmental context. They vary across metabolic 52 tissues, between males and females, and in response to dietary fat levels. Surprisingly, we also find 53 several genes with inconsistent ASE biases that switched direction across tissues and/or contexts (e.g. 54 SM/J biased in one cohort, LG/J biased in another). Together, our results provide novel insights into how 55 genetic, epigenetic, and environmental mechanisms govern allele-specific gene regulation, which is an essential step towards deciphering the genotype to phenotype map. 56

## 57 INTRODUCTION

58 Deciphering the genotype to phenotype map remains a fundamental quest in biology. Gene expression 59 is a promising focal point, as it is an intermediate step between DNA sequence and gross phenotype. 60 Gene expression itself is a complex trait that is regulated by genetic, epigenetic, and environmental 61 factors (Pastinen 2010). Recent efforts to characterize how genes are regulated have often focused on 62 mapping expression quantitative trait loci (eQTLs), which identify genetic variants linked to changes in 63 gene expression at the population level (Cookson et al. 2009). However, these studies can only 64 interrogate total gene expression and assume that genes are biallelically expressed (i.e. both alleles are 65 equally expressed), which may mask underlying regulatory mechanisms. Furthermore, epigenetic 66 changes and gene-by-environment interactions can alter gene expression patterns in a dynamic and 67 tissue-specific manner without modifying the underlying nucleotide sequence; these effects are missed 68 in a typical sequence-based method. An allele-specific approach is required to directly measure how cis-69 regulatory variation impacts gene expression and to tease it apart from *trans*-acting factors that affect 70 both chromosomes (Bonasio et al. 2010).

71 Allele-specific expression (ASE) is a phenomenon where a gene's expression diverges from biallelic. In 72 diploid organisms, one allele is preferentially expressed over the other allele. Previous findings estimate 73 that 30-56% of genes show evidence of allelic imbalance, indicating allele-specific effects have 74 widespread impacts on gene regulation (Ge et al. 2009; Castel et al. 2019; Keane et al. 2011). Depending 75 on a gene's function, these expression imbalances can lead to phenotypic variation with functional 76 consequences. To detect ASE, single nucleotide polymorphisms (SNPs) and other genetic variants are 77 used to distinguish between alleles in heterozygotes and map RNA-sequencing reads to their 78 chromosome of origin (Heap et al. 2010). Allele-specific analyses are a powerful way to exploit a within-79 sample control (the other allele) to gauge how genetic and epigenetic variation in *cis*-regulatory elements 80 shapes gene expression (Pastinen 2010).

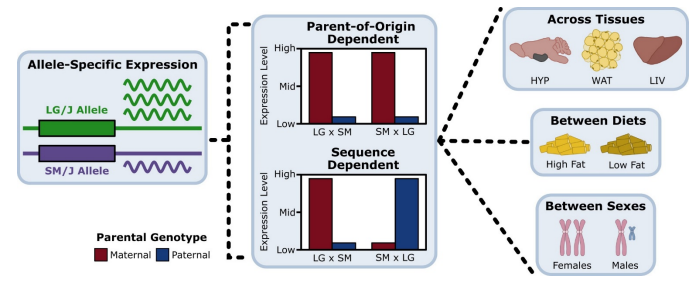
ASE can be divided into two classes: sequence dependent or parent-of-origin dependent (**Figure 1**). Sequence dependent ASE refers to cases where the two alleles are differentially expressed based on

their haplotype or nucleotide identity. These patterns are thought to be largely driven by *cis*-acting genetic
variants in coding and noncoding regions, such as a premature stop codon that truncates one allele's
transcript or a variant in a promoter region that prevents transcription factors from binding (Keane et al.
2011; Rivas et al. 2015). They can also occur further away from the gene yet still impact its expression,
such as motif variations for long-range enhancers or in the sequence context of DNA methyltransferase
substrates (Cavalli et al. 2016; Wienholz et al. 2010).

89 In contrast, parent-of-origin dependent ASE refers to cases where the alleles are differentially expressed 90 based on which parent contributed it, regardless of the underlying sequence. These patterns fall under 91 the umbrella of parent-of-origin effects (**POEs**), a broader class of epigenetic phenomena that manifest 92 as phenotypic differences according to maternal or paternal inheritance (Lawson et al. 2013). The best 93 characterized POE mechanism is genomic imprinting, an extreme case of parent-of-origin dependent 94 ASE where one parent's allele is completely silenced via selective DNA methylation (Barlow and 95 Bartolomei 2014). Known imprinted genes comprise ~1% of the human and mouse genomes, yet play 96 important roles in development, metabolism, cognition, and other complex traits (Reik and Walter 2001).

97 Comprehensive atlases of how both classes of ASE vary between tissues and developmental stages 98 have been generated in human (Leung et al. 2015; Castel et al. 2019) and mouse models (Babak et al. 99 2015; Andergassen et al. 2017). These studies reveal that both parent-of-origin and sequence dependent 100 ASE patterns are not consistent between tissues, indicating that tissue-specific genetic and epigenetic 101 features can mediate allelic imbalances. Additionally, trans-acting environmental factors have been 102 shown to interact with cis-regulatory variants to modulate the magnitude of ASE effects in human (Buil et 103 al. 2015; Knowles et al. 2017; Moverbrailean et al. 2016) as well as in rice models (Shao et al. 2019). 104 Imprinted genes are also known to be responsive to environmental exposures (such as teratogenic 105 agents and maternal nutrition) during fetal development (Kappil et al. 2015). Together, these findings 106 suggest that a complicated crosstalk among genetic variants, epigenetic changes, and environmental 107 signals underlies allele-specific gene regulation, but this model needs to be further investigated.

108 Here, we explored how allelic genotype, parent-of-origin, tissue type, sex, and dietary fat simultaneously 109 work together to influence allele-specific expression patterns in a F<sub>1</sub> reciprocal cross of the LG/J and 110 SM/J inbred mouse strains. These strains have been extensively used in gene-by-environment studies 111 due to their divergent genetic backgrounds and variable responses to dietary nutrition (Ehrich et al. 2003; 112 Nikolskiy et al. 2015; Lawson et al. 2011b, 2010, 2011a; Carson and Lawson 2020; Miranda et al. 2019). 113 We found that both parent-of-origin and sequence dependent ASE patterns are highly dependent on 114 tissue type and environmental context. They vary across metabolic tissues, between males and females, 115 and in response to dietary fat, thus providing novel insights into how gene-by-environmental effects 116 influence complex traits. Untangling the genetic, epigenetic, and environmental mechanisms that govern 117 allele-specific gene regulation is crucial to improving our ability to predict phenotypes from genotypes.



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119 Figure 1: Evaluating parent-of-origin and sequence dependent allele-specific expression across 120 tissues and environmental contexts. We decomposed allele-specific expression (ASE) into its parentof-origin and sequence effects in a F<sub>1</sub> reciprocal cross. An example of parent-of-origin dependent ASE is 121 122 when the maternal allele (red) is preferentially expressed over the paternal allele (blue), regardless of which haplotype contributed it. An example of sequence dependent ASE is when the LG/J allele is 123 124 preferentially expressed over the SM/J allele, regardless of which parent contributed it. Once we identified 125 significant ASE genes, we compared how their expression patterns changed across metabolic tissues 126 (HYP, WAT, LIV), in response to different diets (high fat, low fat), and between sexes (females, males).

## 127 RESULTS

## 128 Allele-specific expression can be decomposed into parent-of-origin and sequence effects

129 We measured ASE in a F<sub>1</sub> reciprocal cross of the LG/J and SM/J inbred mouse strains. Briefly, LG/J

130 mothers were mated with SM/J fathers and vice versa, resulting in F<sub>1</sub> offspring who are genetically

equivalent but differ in the allelic direction of inheritance. Male and female F<sub>1</sub> mice were fed either a highfat or low-fat diet. We obtained RNA-Seq data from three metabolically-relevant tissues: hypothalamus
(HYP), white adipose (WAT), and liver (LIV). To explore how different environmental contexts (dietary fat
and/or sex) impact ASE patterns, we analyzed nine separate cohorts per tissue: high-fat fed diet (H), lowfat fed diet (L), females (F), males (M), high-fat fed females (HF), high-fat fed males (HM), low-fat fed
females (LF), low-fat fed males (LM), and all contexts collapsed (All) (Figure 1).

We harnessed the >6 million SNPs and indels between the LG/J and SM/J genomes (Nikolskiy et al. 2015) to map sequencing reads to their chromosome of origin. Overall, 9,016 protein-coding genes and noncoding RNAs had detectable ASE in at least one tissue-by-context analysis (~31% of all expressed genes, **Supplemental Figure S1**). Next, we identified genes with significant ASE biases and classified them into two patterns: parent-of-origin dependent (unequal expression based on the allele's parental origin) and sequence dependent (unequal expression based on the allele's nucleotide identity) (**Figure** 

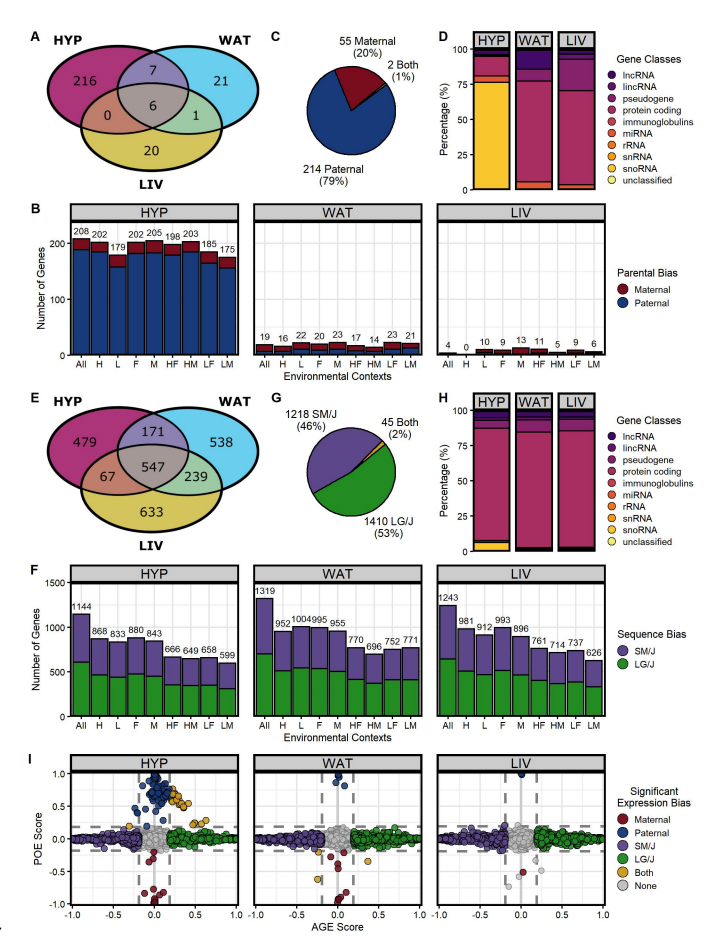
#### 143 **1, Supplemental Figure S2**).

#### 144 Parent-of-origin and sequence dependent ASE patterns are prevalent and distinct

Across our 27 tissue-by-context analyses, we identified 271 genes with significant parent-of-origin 145 146 dependent ASE. HYP had the greatest number of significant genes (n = 229), followed by WAT (n = 35), 147 then LIV (n = 27). 14 genes were expressed in multiple tissues, but the majority were tissue-specific 148 (Figure 2A). In HYP, the most genes were detected in the "All" context. However, in WAT and LIV, more 149 genes were only detected in a specific diet, sex, and/or diet-by-sex context; those expression biases 150 were missed when contexts were collapsed (Figure 2B). 214 genes (79%) were paternally biased, 55 151 genes (20%) were maternally biased, and 2 genes (1%) switched their expression bias direction across 152 the cohorts (Figure 2C). This heavy skew is driven by a 672 kb cluster of 171 paternally-biased small 153 nucleolar RNAs (snoRNAs) located within the Prader-Willi/Angelman syndrome (PWS/AS) orthologous 154 domain on mouse chromosome 7 that are only expressed in HYP (Supplemental Figure S3). Parent-155 of-origin dependent ASE genes also included protein-coding genes, microRNAs, long noncoding RNAs, 156 long interspersed noncoding RNAs, and pseudogenes (Figure 2D).

157 We also identified 2,673 genes with significant sequence dependent ASE across our 27 tissue-by-context 158 analyses. WAT had the greatest number of significant genes (n = 1.495), followed by LIV (n = 1.486), 159 then HYP (n = 1.264). While some genes' biases were tissue-specific, 1.657 genes (62%) were biased 160 in multiple tissues (Figure 2E). In each tissue, the most genes were detected in the "All" context, then 161 the diet- or sex-specific contexts, and finally the diet-by-sex-specific contexts, likely reflecting the sample sizes and power available in each cohort (Figure 2F). 1,218 genes (46%) were SM/J biased, 1,410 genes 162 163 (53%) were LG/J biased, and 45 genes (2%) switched their expression bias direction across the cohorts 164 (Figure 2G). Sequence dependent ASE genes were predominantly classified as protein-coding genes 165 (~80%) in each tissue, but also included pseudogenes, immunoglobulins, and various non-coding RNAs 166 (long non-coding, long interspersed non-coding, micro, ribosomal, small interfering, and small nucleolar) 167 (Figure 2H).

168 Parent-of-origin and sequence dependent ASE patterns were often mutually exclusive. In all three 169 tissues, genes with extreme parental biases typically had weak sequence biases and vice versa (Figure 170 21, Supplemental Figure S4). However, 61 genes in HYP and 3 genes in WAT showed both parental 171 and sequence biases, likely due to epigenetic regulatory mechanisms affected by haplotype 172 variation/allelic identity. Both ASE patterns occurred genome-wide (Supplemental Figure S5). Parent-173 of-origin dependent ASE genes tended to cluster in well-known imprinted domains, such as the 174 Ube3a/Snrpn, Meg3/Gtl2, Peg3/Usp29, and H13/Mcts2 domains (Supplemental Figures S6), Sequence 175 dependent ASE genes were spread more diffusely across chromosomes, but occasionally clustered in 176 regions potentially controlled by the same regulatory element (Supplemental Figures S7).



178 Figure 2: Parent-of-origin and sequence dependent allele-specific expression patterns are 179 prevalent and distinct. (A) Venn diagram of the total parent-of-origin dependent ASE genes across 180 HYP, LIV, and WAT (all contexts collapsed). (B) Number of genes with significant parental biases in each 181 tissue-by-context analysis (maternal = red, paternal = blue): all contexts collapsed (All), high-fat diet (H), low-fat diet (L), females (F), males (M), high-fat fed females (HF), high-fat fed males (HM), low-fat fed 182 183 females (LF), and low-fat fed males (LM). (C) Summary of expression bias directions across all analyses: 184 paternally biased (blue), maternally biased (red), and genes that switch bias direction depending on the 185 cohort (yellow). (D) Gene class proportions of significant parent-of-origin dependent ASE genes in each 186 tissue. (E) Venn diagram of the total sequence dependent ASE genes across HYP, LIV, and WAT (all 187 contexts collapsed). (F) Number of genes with significant sequence biases in each tissue-by-context 188 analysis (SM/J = purple, LG/J = green). (G) Summary of expression bias directions across all analyses: 189 SM/J biased (purple), LG/J biased (green), and genes that switch bias direction depending on the cohort 190 (yellow). (H) Gene class proportions of significant sequence dependent ASE genes in each tissue. (I) 191 Parent-of-Origin Effect (POE) versus Allelic Genotype Effect (AGE) scores in the "All" context of each 192 tissue. Genes with significant parental biases (red = maternal, blue = paternal) have extreme POE scores. 193 but weak AGE scores. Conversely, genes with significant sequence biases (purple = SM/J, green = LG/J) 194 have extreme AGE scores, but weak POE scores. Some genes have both parental and sequence biases 195 (yellow). Most genes have no bias (gray). Dashed lines indicate effect score thresholds.

196 Parent-of-origin dependent ASE recapitulates canonical imprinting patterns

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197 To evaluate whether parent-of-origin dependent ASE is influenced by tissue or environmental context 198 (diet and/or sex), we characterized the expression profiles of the 271 parentally biased genes across our 199 27 tissue-by-context analyses (3 tissues x 9 diet-by-sex contexts). For each analysis, we quantified the 200 direction and magnitude of each gene's parental expression bias by calculating a Parent-of-Origin Effect 201 (**POE**) score from the mean allelic bias of each  $F_1$  reciprocal cross. POE scores range from -1 (completely 202 maternally expressed) to +1 (completely paternally expressed); a score of 0 indicates biallelic expression (see Methods). In each tissue-by-context analysis, a gene could be expressed in one of three ways: 203 204 significant parental bias, biallelic (expressed but with no allele-specific bias), or not expressed. We sorted 205 the 271 genes with significant parent-of-origin dependent ASE into three expression profiles: tissue-206 independent, tissue-dependent, and context-dependent (Figure 3A-3D). 207 We identified 183 tissue-independent genes (67%), defined as a consistent parental bias in every tissue

they were expressed. Within a given tissue, they were parentally biased in most environmental contexts

(Figure 3B). Next, we identified 15 genes (6%) as tissue-dependent. These genes showed a parental

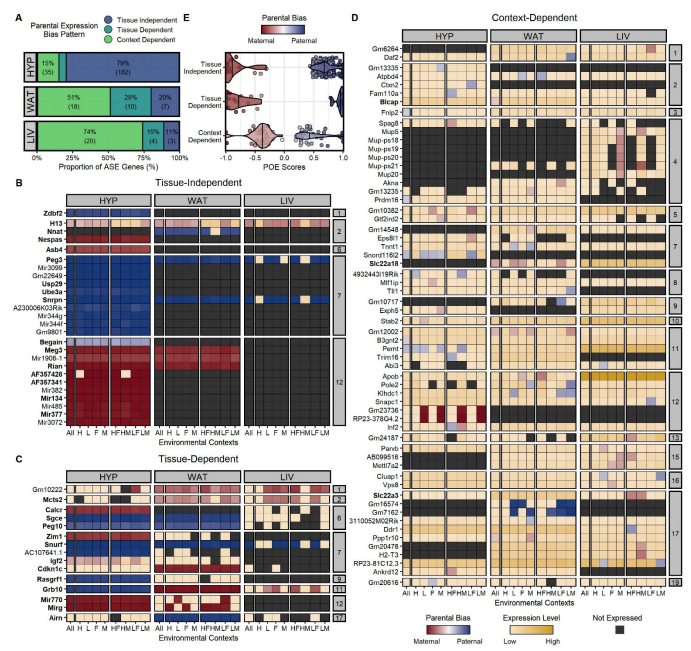
bias across most contexts in one or two tissues, but were biallelically expressed (no bias) across contexts

in the other tissue(s) (Figure 3C). Here, we distinguish between tissue-specific gene expression (not

expressed in certain tissues) and tissue-dependent ASE (biased expression only in certain tissues).

213 All 198 tissue-independent/dependent genes were related to genomic imprinting, the best characterized 214 mechanism of parent-of-origin dependent ASE. 29 genes were canonically imprinted (bolded in Figure 215 3) and the other 169 genes were various non-coding RNAs located within well-known imprinted domains. 216 For example, 156 genes were part of a cluster of 171 paternally-biased snoRNAs in the PWS/AS domain 217 on mouse chromosome 7 (Supplemental Figure S3). As expected with imprinting, these genes were 218 extremely biased towards one parent's allele; their mean POE scores were -0.82 and 0.71 for maternally 219 and paternally biased genes, respectively (Figure 3E). HYP had the highest proportion of these tissue-220 independent/dependent genes among our three adult tissues (85%). largely due to the snoRNA cluster 221 in the PWS/AS domain being HYP-specific. WAT had the second highest proportion (49%), but very few 222 of these genes were expressed in LIV (7%) (Figure 3A). These findings are consistent with the 223 previously-reported dichotomy of imprinting levels between neural and non-neural adult tissues (Babak 224 et al. 2015) as well as imprinting's role in development and cognition (Barlow and Bartolomei 2014).

225 We validated the expression profiles of two canonically imprinted genes (Peg3 and Grb10) by 226 pyrosequencing (Figure 3B-3C, Supplemental Figure S8). Peg3 (paternally expressed gene 3) had a 227 significant paternal bias in all three tissues, regardless of context. Its locus has 60 variants between the 228 LG/J and SM/J backgrounds, though none are predicted to be functional. Peg3 functions as a DNA-229 binding transcriptional repressor to control fetal growth rates, maternal caring behaviors, and tumor 230 growth. It is only expressed from the paternal allele in most tissues, especially the placenta and brain 231 (Thiaville et al. 2013; He and Kim 2014). Grb10 (growth factor receptor bound protein 10) had a tissue-232 dependent pattern of significant paternal bias in HYP but a maternal bias in WAT and LIV. Its locus has 233 154 variants between the LG/J and SM/J backgrounds, but none are predicted to be functional. Grb10 234 encodes an adapter protein that interacts with receptor tyrosine kinases to impact insulin signaling and 235 growth hormone pathways (He et al. 1998). It has a documented pattern of maternal expression in most 236 adult mouse tissues, but paternal expression in the brain (Plasschaert and Bartolomei 2015).



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238 Figure 3: Atlas of parent-of-origin dependent ASE patterns across tissues and environmental 239 contexts. (A) Number and proportion of ASE genes per tissue with each parental expression bias profile: 240 tissue-independent (dark blue), tissue-dependent (teal), and context-dependent (green). Heatmaps of ASE profiles across tissues and contexts: (B) tissue-independent (parental bias wherever expressed). 241 242 (C) tissue-dependent (parental bias in some tissues, no bias in others), and (D) context-dependent 243 (parental bias only in certain diet-by-sex contexts, no bias elsewhere). All genes with significant parent-244 of-origin dependent ASE are shown, except the cluster of 171 snoRNAs on chr 7 (Supplemental Figure 245 S3). Genes are color-coded by their expression pattern in each tissue-by-context analysis. Shades of red 246 and blue indicate their degree of maternal or paternal bias, respectively (POE scores). Where genes are 247 not significantly biased, shades of yellow indicate their biallelic gene expression levels (log-transformed 248 total counts). Black indicates genes are not expressed in that context/tissue. Bolded genes are 249 canonically imprinted. The v-axis is grouped and sorted by chromosomal position. Each supercolumn 250 denotes a tissue: hypothalamus (HYP), white adipose (WAT), and liver (LIV). Each subcolumn denotes 251 an environmental context: all contexts collapsed (All), high-fat diet (H), low-fat diet (L), females (F), males 252 (M), high-fat fed females (HF), high-fat fed males (HM), low-fat fed females (LF), and low-fat fed males

(LM). (E) Parent-of-Origin Effect (POE) score distributions for each parental expression bias profile.
 Vertical lines indicate that profile's mean POE score. Dots represent individual significant ASE genes.

255 Dietary environment and sex influence parent-of-origin dependent ASE in a partially imprinted manner

256 Finally, we classified 73 genes (27%) as context-dependent, meaning they had a parental bias only in 257 certain environmental contexts within a tissue, but biallelic expression (no bias) in other contexts and/or 258 tissues (Figure 3D). Only three of these genes were canonically imprinted and 18 genes were non-coding 259 RNAs located in known imprinted domains (including 14 snoRNAs in the PWS/AS domain). The 260 remaining 52 genes had no clear connection to genomic imprinting, yet they showed significant parent-261 of-origin dependent ASE in certain context(s). These context-dependent genes had more subtle allelic 262 biases than the tissue-independent/dependent genes; their mean POE scores were -0.39 and 0.39 for 263 maternally and paternally biased genes, respectively (Figure 3E). These patterns are consistent with 264 partial imprinting, where the two parental alleles are differentially expressed in a less extreme manner than the uniparental expression associated with genomic imprinting (Wolf et al. 2008; Morcos et al. 2011). 265 266 Interestingly, LIV had the highest proportion of context-dependent genes among its total parent-of-origin 267 dependent ASE genes (74%) while HYP had the lowest proportion (15%) (Figure 3A). LIV is also a tissue 268 where more parent-of-origin dependent ASE genes were detected in the diet- and/or sex-specific 269 contexts than the "All" context (Figure 2B). LIV is incredibly responsive to environmental factors 270 (especially diet), given its central roles in digestion and detoxification (Trefts et al. 2017). Taken together, 271 these findings suggest a mechanism of parent-of-origin dependent ASE outside of traditional imprinting 272 that is sensitive to environmental perturbations.

To further explore how intrinsic (sex) and/or extrinsic (dietary fat) environments can alter parental ASE biases, we calculated individualized POE scores for each context-dependent gene and modeled how they vary across diet, sex, and diet-by-sex contexts. These categories were not mutually exclusive; across all three tissues, most context-dependent genes were significant for more than one effect. Nonetheless, when we intersected these significant gene lists, we found that each tissue showed a distinct pattern of context-dependent parental ASE biases (**Supplemental Figure S9**). For example, LIV had similar proportions of significant sex, diet, and diet-by-sex effects (each 75 – 80% of its context-

dependent genes); 60% of its genes were significant for all three effects. Most of WAT's genes had a
significant sex effect (83%), but diet or diet-by-sex effects were much less common (44% and 28%,
respectively). Finally, 63% of HYP's genes had a significant sex effect, while 40 – 45% had significant
diet or diet-by-sex effects.

284 We validated the expression profiles of two context-dependent genes (Apob and SIc22a3) by pyrosequencing (Figure 3D, Supplemental Figure S10). Apob (apolipoprotein B) had a significant diet 285 286 effect in WAT, reflected in significant maternal biases in the HF and F contexts. It also had a maternal 287 bias in the HM context, but low sample sizes excluded it from further analysis, Apob was biallelically 288 expressed in the remaining contexts in WAT and all contexts in LIV and HYP. Its locus has 126 variants 289 between the LG/J and SM/J backgrounds, including seven non-synonymous SNPs in the coding region 290 (six = LG/J genome, one = SM/J genome). Apob produces the main component of lipoproteins, which 291 transport lipids (including cholesterol) in the blood (Olofsson and Borèn 2005). Maternal-specific 292 associations between APOB variants and adiposity traits have been found in humans (Hochner et al. 293 2015). Apob expression levels also differ between high and low fructose diets in mice livers, suggesting 294 its function is susceptible to dietary environment (Sud et al. 2017). Slc22a3 (solute carrier family 22 295 member 3) had significant diet and diet-by-sex effects in LIV, reflected in strong maternal biases in the 296 HF and HM contexts. SIc22a3 was biallelically expressed in the remaining contexts in LIV and all contexts 297 in WAT and HYP. Its locus has 285 variants between the LG/J and SM/J backgrounds, but none are 298 predicted to be functional. SIc22a3 is a transporter that eliminates organic cations from cells, such as 299 monoamine neurotransmitters, cationic drugs, and xenobiotics (Kekuda et al. 1998). It has been reported 300 to be maternally expressed in liver and extraembryonic tissues, but biallelically expressed elsewhere 301 (Babak et al. 2015). Slc22a3 is also significantly differentially expressed in kidneys between high-fat diet-302 and chow-fed mice (Gai et al. 2016).

303 Sequence dependent ASE arises from haplotype-specific genetic variation

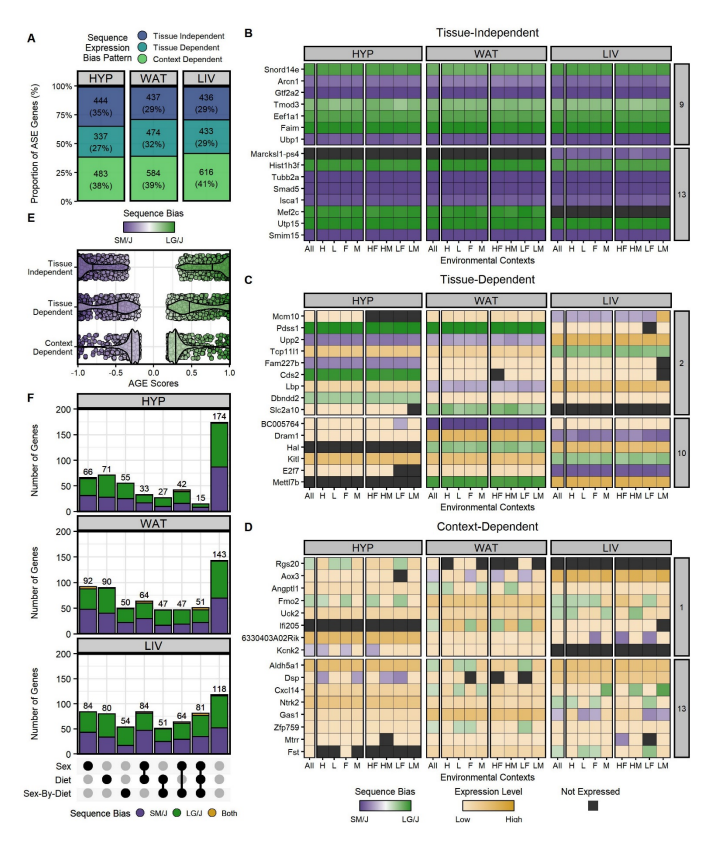
Next, we similarly characterized the expression profiles of the 2,673 sequence biased genes across our
 27 tissue-by-context analyses (3 tissues x 9 diet-by-sex contexts) to evaluate whether sequence

306 dependent ASE is also influenced by tissue or environmental context. For each analysis, we quantified 307 the direction and magnitude of each gene's sequence expression bias by calculating an Allelic Genotype 308 Effect (AGE) score from the mean allelic bias of each  $F_1$  reciprocal cross. AGE scores range from -1 309 (completely SM/J expressed) to +1 (completely LG/J expressed); a score of 0 indicates biallelic 310 expression (see Methods). In each analysis, a gene could be expressed in one of three ways: significant 311 sequence/allelic genotype bias, biallelic (expressed but with no allele-specific bias), or not expressed. 312 We sorted the 2,673 genes with significant sequence dependent ASE into three expression profiles: 313 tissue-independent, tissue-dependent, and context-dependent (Figure 4A-4D).

314 We identified 605 genes (23%) as tissue-independent, meaning they had a consistent sequence bias in 315 every tissue they were expressed. Within a given tissue, they had a sequence bias in most environmental 316 contexts (Figure 4B). These tissue-independent genes were strongly biased towards one strain's allele; 317 their mean AGE scores were -0.79 and 0.78 for SM/J and LG/J biased genes, respectively (Figure 4E). 318 They also comprised a similar proportion of the total sequence dependent ASE genes in all three tissues 319 (29-35%, Figure 4A). These patterns likely reflect the vast genetic variation that has accumulated 320 between the LG/J and SM/J backgrounds over the decades, whereby a cis-acting variant impacts one 321 strain's allelic function wherever that gene is expressed, regardless of tissue type.

322 We validated the expression profiles of two tissue-independent genes (*Eef1a1* and *Tubb2a*) by 323 pyrosequencing (Figure 4B, Supplemental Figure S11). Eef1a1 (eukaryotic translation elongation 324 factor 1 alpha 1) had a significant LG/J allelic bias in all three tissues, regardless of context. Its locus has 325 30 variants between the LG/J and SM/J backgrounds, including two non-synonymous SNPs in the coding 326 region of the SM/J genome. Eef1a1 delivers aminoacylated transfer RNAs to the elongating ribosome 327 during protein synthesis and has crucial roles in protein degradation, RNA virus replication, and other 328 cellular processes. It is abundantly and ubiquitously expressed in most tissues (Mateyak and Kinzy 2010; 329 Li et al. 2013). Tubb2a (tubulin beta-2A Class IIa) had a significant SM/J allelic bias in all three tissues, 330 regardless of context. Mutations in this gene are rare and often non-viable; however, its locus has one 331 SNP in the LG/J genome. Tubb2a is a tubulin isoform that binds GTP to create microtubules, which are

- 332 critical for the cytoskeleton organization, intracellular trafficking, and mitotic cell division. It is also
- 333 ubiquitously expressed in most tissues, especially the brain (Hammond et al. 2008; Rice et al. 2008).



335 Figure 4: Atlas of sequence dependent ASE patterns across tissues and environmental contexts. 336 (A) Number and proportion of ASE genes per tissue with each sequence expression bias profile: tissue-337 independent (dark blue), tissue-dependent (teal), and context-dependent (green). Heatmaps of ASE 338 profiles across tissues and contexts: (B) tissue-independent (sequence bias wherever expressed), (C) 339 tissue-dependent (sequence bias in some tissues, no bias in others), and (D) context-dependent 340 (sequence bias only in certain diet-by-sex contexts, no bias elsewhere). A subset of the 2,673 genes with 341 significant sequence dependent ASE are shown, including those validated with pyrosequencing. Genes 342 are color-coded by their expression pattern in each tissue-by-context analysis. Shades of purple and 343 green indicate their degree of SM/J or LG/J allelic bias, respectively (AGE scores). Where genes are not 344 significantly biased, shades of yellow indicate their biallelic gene expression levels (log-transformed total 345 counts). Black indicates genes are not expressed in that context/tissue. The y-axis is grouped and sorted 346 by chromosomal position. Each supercolumn denotes a tissue: hypothalamus (HYP), white adipose 347 (WAT), and liver (LIV). Each subcolumn denotes an environmental context: all contexts collapsed (All), 348 high-fat diet (H), low-fat diet (L), females (F), males (M), high-fat fed females (HF), high-fat fed males 349 (HM), low-fat fed females (LF), and low-fat fed males (LM). (E) Allelic Genotype Effect (AGE) score 350 distributions for each sequence expression bias profile. Vertical lines indicate that profile's mean AGE 351 score. Dots represent individual significant ASE genes. (F) UpSet plots for each tissue summarizing the 352 set intersections of context-dependent genes with significant sex, diet, and/or sex-by-diet effects. Bar 353 height and color indicate the number of genes with each sequence bias direction: SM/J biased (purple), 354 LG/J biased (green), and those that switch expression bias direction depending on the cohort (yellow).

355 Sequence dependent ASE can be mediated by tissue-specific features

356 Next, we identified 684 genes (25%) as tissue-dependent. These genes showed a sequence bias across 357 most contexts in one or two tissues, but were biallelically expressed (no bias) across contexts in the other 358 tissue(s) (Figure 4C). Here, we again distinguish between tissue-dependent ASE (biallelic expression in 359 some tissues) and tissue-specific gene expression (simply not expressed in some tissues). These tissue-360 dependent genes were moderately biased towards one strain's allele; their mean AGE scores were -0.57 and 0.55 for SM/J and LG/J biased genes, respectively (Figure 4E). They also comprised a similar 361 362 proportion of the total sequence dependent ASE genes in all three tissues (27-32%, Figure 4A). These 363 patterns demonstrate that sequence dependent ASE is not solely due to genetic variation; here, tissue-364 specific epigenetic factors likely interact with *cis*-acting variants to influence allelic frequency. 365 We validated the expression profiles of two tissue-dependent genes (Upp2 and Mettl7b) by

366 pyrosequencing (**Figure 4C, Supplemental Figure S12**). *Upp2* (uridine phosphorylase 2) had a strong

367 SM/J allelic bias across contexts in HYP, a weaker SM/J bias in WAT, and biallelic expression in LIV. Its

368 locus has 1,043 variants between the LG/J and SM/J backgrounds, including five non-synonymous SNPs

369 in the coding region of the LG/J genome. *Upp2* catalyzes the phosphorolysis of uridine into uracil and

370 ribose-1-phosphate, which are used as carbon and energy sources during catabolic metabolism. In mice,

it is predominantly expressed in liver and weakly expressed in brain (Johansson 2003; Roosild et al. 2011). *Mettl7b* (methyltransferase-like 7b) had a LG/J allelic bias across contexts in WAT, biallelic expression in LIV, and was not expressed in HYP. Its locus has 77 variants between the LG/J and SM/J backgrounds, including one non-synonymous SNP in the coding region of the SM/J genome. *Mettl7b* is an akyl thiol methyltransferase that is implicated in several cancers. It is highly expressed in lipid droplets, particularly in liver and adipose tissue (Maldonato et al. 2021; Turró et al. 2006).

377 Sequence dependent ASE is sensitive to sex and dietary environments

378 Finally, we classified 1,384 genes (52%) as context-dependent. These genes had a sequence bias only 379 in certain environmental contexts within a tissue, but biallelic expression (no bias) in other contexts and/or 380 tissues (Figure 4D). These context-dependent genes had more subtle allelic biases than the other two 381 profiles; their mean AGE scores were -0.32 and 0.34 for SM/J and LG/J biased genes, respectively 382 (Figure 4E). LIV had the most context-dependent genes (n = 616) and HYP had the fewest (n = 483), 383 but overall they comprised a similar proportion of the total ASE genes in all three tissues (38-41%, Figure 384 4A). These patterns suggest that environmental factors can interact with genetic variation to influence 385 the final allelic composition of a gene's product, resulting in sequence dependent ASE.

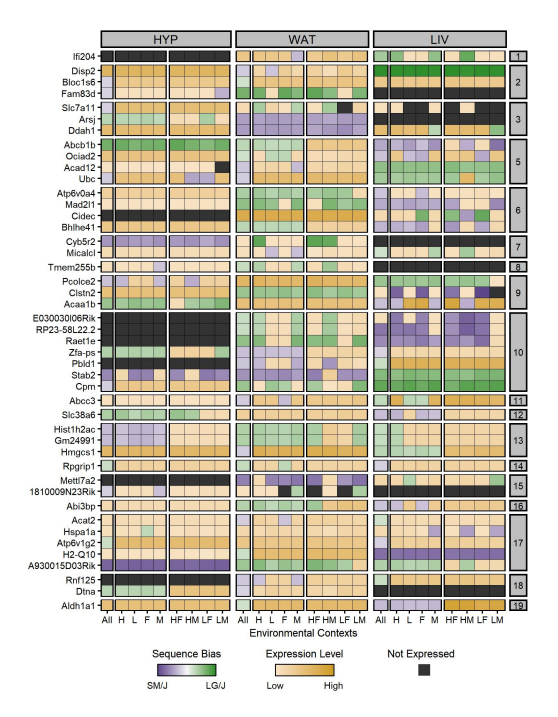
386 To further explore how sex and/or dietary fat can alter sequence/allelic genotype ASE biases, we 387 calculated individualized AGE scores for each context-dependent gene and modeled how they vary 388 across diet, sex, and diet-by-sex contexts (Supplemental Table S13). When we intersected these gene 389 lists, we found that each tissue showed a similar pattern of context-dependent sequence ASE biases 390 (Figure 4F). Significant sex effects were the most prevalent in each tissue, ranging from 156 genes in 391 HYP (32%), to 254 genes in WAT (43%), and 313 genes in LIV (51%). Diet effects were slightly less 392 common but still widespread: 146 genes in HYP (30%), 252 genes in WAT (43%), and 296 genes in LIV 393 (48%). Finally, diet-by-sex effects were the least frequent yet incredibly rampant, comprising 139 genes 394 in HYP (29%), 195 genes in WAT (33%), and 250 genes in LIV (41%). These categories were not 395 mutually exclusive; most context-dependent genes were significant for more than one effect but in 396 different combinations across the three tissues.

397 We validated the expression profiles of two context-dependent genes (Ifi205 and Gas1) by 398 pyrosequencing (Figure 4D, Supplemental Figure S14). Ifi205 (interferon activated gene 205) had 399 significant sex and diet-by-sex effects in WAT, reflected in strong LG/J allelic biases in the three female-400 related (HF, LF, F) and "All" contexts. Ifi205 was biallelically expressed in the remaining contexts in WAT 401 and all contexts in LIV. Its locus has 254 variants between the LG/J and SM/J backgrounds, including 402 ten non-synonymous SNPs in the coding region of the SM/J genome. If 205 binds DNA in response to 403 interferon signaling, a cytokine family that activates the immune system. Significant sex differences have 404 been reported in mouse Ifi205 expression levels, where females have higher total expression levels than 405 males (Albrecht et al. 2005; Cao et al. 2018). Gas1 (growth arrest specific 1) had significant diet and sex 406 effects in LIV, reflected in strong SM/J biases in the three low-fat fed diet-related (LF, LM, L), female, and 407 "All" contexts. Gas1 was biallelically expressed in the remaining high-fat fed diet and male contexts in 408 LIV and all contexts in HYP and WAT. Its locus has 57 variants between the LG/J and SM/J backgrounds, 409 though none are predicted to have functional impacts. Gas1 encodes a membrane glycoprotein that binds 410 and regulates sonic hedgehog during development. Gas1 has been found to be differentially expressed 411 between high and low selenium diets in mice ovaries, suggesting its expression may be sensitive to 412 dietary environment (Lee et al. 2001; Qazi et al. 2021).

413 Sequence dependent ASE genes can switch the direction of their allelic biases

414 Surprisingly, we found 45 sequence dependent ASE genes with inconsistent patterns of allelic biases. 415 These genes showed significant ASE in opposite directions among the tissues and/or environmental 416 contexts (Figure 5). For 44 of the 45 genes (98%), such direction-switching occurred at the tissue level: 417 for example, a gene may have a LG/J bias in one tissue, a SM/J bias in another tissue, and sometimes 418 even no bias (biallelic) in the third tissue. Four genes had tissue-independent ASE, or a sequence bias 419 across contexts in every expressed tissue (albeit in different directions). 19 genes had context-dependent 420 ASE, or a sequence bias only in certain diet and/or sex contexts that switched direction across tissues. 421 One of these genes (Cidec) had a sex-dependent switch in ASE direction within the same tissue, 422 discussed further below. The remaining 22 genes had a combination of context- and tissue-dependent 423 ASE patterns: such genes had a sequence bias in one direction across most contexts in one or two

tissues, but a context-dependent sequence bias in the opposite direction in another tissue. These dynamic direction-switching patterns confirm that sequence dependent ASE is not solely due to genetic variation; otherwise, the same variant causing biased expression in one tissue would also be present in the cells of any other tissue expressing that gene. These patterns hint at epigenetic regulatory elements or post-transcriptional modifications that interact with genetic variation in a tissue-specific manner to influence the final allelic frequency.



431 Figure 5: 45 sequence dependent ASE genes switch their expression bias directions across 432 tissues or environmental contexts. Heatmap of ASE profiles for the 45 genes with significant sequence 433 biases in opposite directions across tissues and/or environmental contexts, including those validated with 434 pyrosequencing. Genes are color-coded by their expression pattern in each tissue-by-context analysis. 435 Shades of purple and green indicate their degree of SM/J or LG/J allelic bias, respectively (AGE scores). 436 Where genes are not significantly biased, shades of yellow indicate their biallelic gene expression levels 437 (log-transformed total counts). Black indicates genes are not expressed in that context/tissue. The y-axis 438 is grouped and sorted by chromosomal position. Each supercolumn denotes a tissue: hypothalamus 439 (HYP), white adipose (WAT), and liver (LIV). Each subcolumn denotes an environmental context: all 440 contexts collapsed (All), high-fat diet (H), low-fat diet (L), females (F), males (M), high-fat fed females 441 (HF), high-fat fed males (HM), low-fat fed females (LF), and low-fat fed males (LM).

442 We validated the expression profiles of three direction-switching genes (*Stab2*, *Ociad2*, and *Cidec*) with

pyrosequencing (Figure 5, Supplemental Figure S15). *Stab2* (stabilin 2) had a significant SM/J bias
across most contexts in HYP and WAT, but a LG/J bias in all contexts in LIV. Its locus has 1,019 variants

between the LG/J and SM/J backgrounds, including 14 non-synonymous SNPs in the coding region of

the SM/J genome. Stab2 binds to hyaluronic acid and mediates its transportation inside the cell (Zhou et

447 al. 2002). It is predominantly expressed in two isoforms in the liver and spleen, while weakly expressed

448 in the brain and adipose tissue (Falkowski et al. 2003). Together with the non-synonymous variants in

the SM/J allele, this may explain Stab2's LG/J bias in the LIV but SM/J bias in the other two tissues.

450 Ociad2 (ovarian cancer immunoreactive antigen domain-containing protein 2) had significant sex and

451 sex-by-diet effects in WAT and LIV. These are reflected in strong LG/J biases in the L, F, and "All"

452 contexts in WAT, but strong SM/J biases in the LF, L, F, and "All" contexts in LIV. Ociad2 was biallelically

453 expressed in the remaining contexts of both tissues and in HYP. Its locus has 118 variants between the

454 LG/J and SM/J backgrounds, though none are predicted to be functional. *Ociad2* activates STAT3,

455 regulates cell migration, and is implicated in several cancers. It is moderately expressed in several non-

456 cancerous tissues, including the brain and liver (Sinha et al. 2018). Sex or diet differences in expression
457 levels have not been explored in the literature, but *Ociad2* is highly expressed in female ovarian tumors
458 (Nagata et al. 2012). Low-fat fed female WAT and LIV tissues have strong sequence biases in *Ociad2*

459 (but in different directions), suggesting a tissue- and sex-specific expression pattern that could have 460 functional consequences.

An interesting exception to the tissue-level switch in bias direction is the gene *Cidec* (cell death-inducing
 DNA fragmentation effector C), which showed a sex-dependent switch in ASE direction within the same

463 tissue validated by pyrosequencing (Figure 5, Supplemental Figure S15). In LIV, Cidec had a strong 464 LG/J bias in the HF, LF, and F contexts, yet a SM/J bias in the HM context. Cidec was biallelically 465 expressed in the remaining LIV contexts and across all WAT contexts, but was not expressed in HYP. 466 Its locus has 46 variants between the LG/J and SM/J backgrounds, though none are predicted to be 467 functional, Cidec promotes lipid droplet formation in adipocytes and regulates low-density lipoprotein 468 maturation in hepatocytes (Xu et al. 2012). Hepatic Cidec expression is very sensitive to diet composition 469 and sex hormones; in particular, female mice have lower total expression levels than males when fed a 470 Western diet (high cholesterol and saturated fats) (Herrera-Marcos et al. 2020).

## 471 **DISCUSSION**

472 Allele-specific expression imbalances due to genetic and epigenetic variation have widespread functional 473 consequences on complex traits, but how environmental signals contribute to this crosstalk remains 474 understudied. Here, we explored how allelic genotype, parent-of-origin, tissue type, sex, and dietary fat 475 simultaneously impact ASE biases in an adult mouse F<sub>1</sub> reciprocal cross. We present a comprehensive 476 genome-wide map of both parent-of-origin and sequence dependent ASE patterns across three 477 metabolically-relevant tissues and nine environmental contexts. The granularity of our analyses revealed 478 that both types of ASE are highly dependent on tissue type and environmental context. We identified 479 2.853 genes with significant parental and/or sequence biases and sorted them into three major 480 expression profiles: tissue-independent, tissue-dependent, and context-dependent. Interestingly, we also 481 found several genes with inconsistent ASE biases that switched direction across tissues and/or contexts 482 (e.g. SM/J biased in one cohort, LG/J biased in another). Although the breadth of these patterns 483 precludes a detailed discussion of each gene, we have validated examples of each expression profile to 484 show how these allelic imbalances could manifest and lead to potential functional consequences.

485 "Tissue-independent" genes are strongly biased wherever they are expressed. Most parent-of-origin 486 dependent ASE genes (67%) have this expression profile – all of which are related to genomic imprinting, 487 a well-characterized epigenetic phenomenon that results in uniparental expression and is often 488 conserved across tissues (Barlow and Bartolomei 2014). In contrast, only 23% of sequence dependent

ASE genes have this expression profile, likely due to strain-specific *cis*-acting genetic variants in coding regions that severely affect one allele's function whenever that gene is expressed. Both patterns conform to the conventional mechanism for ASE whereby the genetic and epigenetic processes that influence the allelic composition will always do so wherever a gene is expressed, in a manner impervious to tissue type or environmental signals (Lo et al. 2003; Babak et al. 2015). However, we found this is not always the case.

495 "Tissue-dependent" genes are moderately biased in one direction in some tissues, but are biallelically 496 expressed (no bias) or biased in the opposite direction in other tissues. Tissue-specific gene expression (simply not expressed in certain tissues) is not considered here, as we are specifically interested in cases 497 498 where a gene has one allelic ratio in one tissue (e.g. 80% SM/J, 20% LG/J) and a different allelic ratio in 499 another tissue (e.g. 50% SM/J, 50% LG/J). Merely 6% of parent-of-origin dependent ASE genes have 500 this expression profile and all are located in known imprinted domains. 25% of sequence dependent ASE 501 genes also have this expression profile, demonstrating that sequence biases are often not solely 502 attributable to genetic variation in coding regions. Instead, both patterns may be explained by variation 503 in tissue-specific regulatory features, such as enhancers or RNA binding proteins. Such tissue-specific 504 factors can then interact with *cis*-acting variants to produce a tissue-dependent allelic imbalance. This is 505 likely the case for genes with inconsistent biases between tissues, where the variation in one allele may 506 be favorable in certain tissues' regulatory landscape but the opposite allele is more favorable elsewhere 507 (Andergassen et al. 2017; Leung et al. 2015).

508 Finally, "context-dependent" genes are subtly biased in one direction in certain environmental contexts, 509 but are biallelically expressed or biased in the opposite direction in other contexts and tissues. We found 510 this expression profile is more prevalent than expected in both classes of ASE. 27% of parent-of-origin 511 dependent ASE genes have diet- and/or sex-specific biases in a less extreme manner than the other 512 profiles. Most genes have no clear connection to imprinting, suggesting another mechanism for parental 513 ASE outside of traditional imprinting that is sensitive to environmental perturbations (Wolf et al. 2008; 514 Morcos et al. 2011; Macias-Velasco et al. 2021). Surprisingly, over half of sequence dependent ASE 515 genes (52%) have diet- and/or sex-specific biases, indicating these intrinsic and extrinsic environmental

factors interact with strain-specific genetic variation to cause a sequence bias. Both patterns suggest a model where environmental signals may interact more efficiently with one allele over the other, leading to shifting and inconsistent allelic proportions in response to environmental cues (Shao et al. 2019). Often, these genes are not significantly biased in our "all contexts" analysis, only in a specific diet and/or sex cohort. This highlights the necessity of studying gene-by-environment interactions, as such effects are obscured when multiple contexts are collapsed together.

522 It is important to note that we can only detect ASE in genes with strain-specific variants in transcribed 523 regions, which is a fraction of the total set of expressed genes. In particular, imprinted genes that are crucial for development/survival may be under tight evolutionary control and not have variants, thus we 524 525 are unable to assess their ASE status here. Although our exact findings are limited to this F<sub>1</sub> reciprocal 526 cross mouse model, the broad patterns nevertheless demonstrate the complexity of allele-specific gene 527 regulation and its contribution to complex traits. Traditional mapping studies connect genotypes to 528 phenotypes, but are agnostic to tissue and often do not consider environment. eQTL studies connect 529 genotypes to total gene expression levels, but assume biallelic expression. Tissue- and context-530 dependent ASE of both classes (parent-of-origin and sequence dependent) can bridge these approaches 531 and pinpoint what tissues and/or environments are relevant for a phenotype. A gene could be expressed 532 at the same level in two cohorts, but its allelic composition could differ due to ASE. If there is a functional 533 difference between the two alleles, then an expression imbalance in the right tissue and/or environment 534 could lead to phenotypic consequences. Incorporating these dynamic ASE patterns into our frameworks 535 will help us decipher the genotype to phenotype map.

#### 536 METHODS

#### 537 *F*<sup>1</sup> Reciprocal Cross Mouse Model

We obtained LG/J and SM/J founders from The Jackson Laboratory (Bar Harbor, ME) and generated F<sub>1</sub> reciprocal crosses by mating LG/J mothers with SM/J fathers (**LxS**) and vice versa (**SxL**). F<sub>1</sub> offspring were weaned into sex-specific cages at three weeks of age and randomly placed on either a high-fat diet (42% kcal from fat; Teklad TD88137) or an isocaloric low-fat diet (15% kcal from fat; Research Diets

542 D12284). They were fed *ad libitum*. F<sub>1</sub> mice were euthanized at 20 weeks of age with a sodium 543 pentobarbital injection followed by cardiac perfusion with phosphate-buffered saline. We harvested 544 hypothalamus (**HYP**), liver (**LIV**), and reproductive white adipose (**WAT**) tissue, which were flash frozen 545 in liquid nitrogen and stored at -80°C until RNA extraction. All procedures were approved by the 546 Institutional Animal Care and Use Committee at Washington University School of Medicine.

547 RNA-Sequencing and Allele-Specific Mapping

548 We sequenced 32 samples per tissue, representing 4 mice from each sex (male or female), diet (high or 549 low fat), and F1 reciprocal cross (LxS or SxL) cohort. We extracted total RNA from WAT and HYP using 550 the RNeasy Lipid Tissue Mini Kit (QIAGEN) and from LIV using a standard TRIzol-chloroform procedure. 551 Samples were selected based on sufficient NanoDrop RNA concentrations (ThermoFisher) and RNA 552 integrity scores ≥8.0 (Agilent). We constructed RNA-Seg libraries with the RiboZero rRNA Removal Kit 553 (Illumina), checked their quality with the BioAnalyzer DNA 1000 assay (Agilent), and sequenced them at 100bp, paired-end reads on an Illumina HiSeg 400. After sequencing, reads were de-multiplexed and 554 555 assigned to individual samples.

556 Quantifying ASE in a  $F_1$  reciprocal cross is vulnerable to reference genome alignment bias. If one parental 557 strain is more closely related to the reference genome, then their reads tend to have a higher mapping guality than reads from the other parental strain (Wang and Clark 2014; Degner et al. 2009). We mitigated 558 559 this concern by aligning RNA-Seg reads to a custom LG/J x SM/J "merged genome". Previously, LG/J 560 and SM/J reference genomes were created by combining strain-specific SNVs and indels with the 561 GRC38.72-mm10 reference template (Nikolskiy et al. 2015). Customized gene annotations were also 562 created by adjusting Ensembl definitions (Mus musculus.GRCm38.72) for indexing differences due to 563 strain-specific indels. Here, those LG/J and SM/J genomes were combined into one pseudogenome so 564 we could align reads to both parental strains simultaneously.

565 We mapped reads uniquely using the two-pass mapping strategy in STAR v.2.7.2b (Dobin et al. 2013). 566 Briefly, splice junctions are collected during the first round and used to inform a second round of mapping, 567 thus detecting more reads that span novel junctions. By not allowing multi-mapping, we only retained

reads uniquely covering strain-specific variants so we could assign reads to their allelic origin; reads covering identical regions between the parental strains were discarded. STAR alignment summaries are provided in **Supplemental Table S1** and **Supplemental Figure S16**.

571 Next, we assigned each aligned read to a gene using bedtools v.2.27.1 (Quinlan and Hall 2010) and our 572 strain-specific Ensembl annotations. Gene-level allele-specific read counts were then upper quartile 573 normalized (**Supplemental Figure S17**) and filtered to remove lowly-expressed genes (total normalized 574 read counts <20). We retained a total of 9171 genes with detectable allele-specific expression in HYP, 575 9761 genes in WAT, and 8082 genes in LIV.

576 Library Complexity

577 Insufficient library complexity can also hamper detecting ASE in a F<sub>1</sub> reciprocal cross. In lowly or 578 moderately expressed genes, if a read fragment from only one of the two alleles is randomly subjected 579 to a duplicating event (i.e. PCR jackpotting) during RNA-Seq library construction, then that gene may 580 spuriously appear as monoallelically expressed, even though it is a false positive (Wang and Clark 2014). 581 To check for this, we measured each library's complexity by fitting the distribution of LG/J allele 582 expression biases (the proportion of total allele-specific read counts with the LG/J haplotype) to a beta-583 binomial distribution using the VGAM package (Yee 2010). We estimated the shape parameters ( $\alpha$ ,  $\beta$ ) of 584 the beta-binomial distribution and calculated the overdispersion parameter (p) as  $\rho = 1 / (1 + \alpha + \beta)$ . 585 Lower values of  $\rho$  (< 0.075) indicate a library is sufficiently complex. One WAT library (CCGGACC) and 586 one LIV library (TGATTAC) were deemed to have poor complexity and were removed from further 587 analyses (Supplemental Figure S18).

588 Determining Biased Allele-Specific Expression

To explore how environmental context (diet and/or sex) impacts ASE patterns, we analyzed nine separate cohorts per tissue: high fat diet (**H**), low fat diet (**L**), females (**F**), males (**M**), high-fat fed females (**HF**), high-fat fed males (**HM**), low-fat fed females (**LF**), low-fat fed males (**LM**), and all contexts together (**AII**) (**Figure 1A**) (3 tissues x 9 contexts = 27 tissue-by-context cohorts). For each tissue-by-context analysis, we required a gene to be expressed in  $\geq$ 75% of the total sample size per F<sub>1</sub> reciprocal cross: 3 mice per

594 cross for diet-by-sex-specific contexts (N = 4), 6 mice per cross for diet- and sex-specific contexts (N = 595 8), and 12 mice per cross for the "All" context (N = 16).

596 We adapted a previously-published model (Takada et al. 2017) for jointly estimating parent-of-origin (PO) 597 and allelic genotype (AG) effects on ASE. First, we assigned two binary variables to each gene's allele-598 specific counts based on their allelic origin. For the PO term, maternal alleles received a 0 and paternal alleles received a 1; for the AG term, LG/J alleles received a 0 and SM/J alleles received a 1 599 600 (Supplemental Figure S2). Next, we implemented a paired-sample design to handle missing data in 601 edgeR, which converts NAs to zero counts. There is a fundamental difference between a gene not being 602 expressed in a particular sample (both allele counts are zero) and an extreme allelic expression bias 603 (only one allele's counts are zero). Thus, we added a series of n-1 dummy variables (indicating library 604 barcodes) to the GLM so that both allele-specific counts for each sample are treated as a pair. If a gene 605 is not expressed in a library, then the coefficient corresponding to that sample will approach negative 606 infinity in the fitted GLM. The missing sample is effectively removed from consideration and does not 607 affect the PO and AG coefficient estimates for the other samples. Finally, we fit a negative binomial 608 generalized linear model (GLM) and conducted likelihood ratio tests in edgeR (Robinson et al. 2010; 609 McCarthy et al. 2012) to estimate the PO and AG effects on allele-specific gene expression levels.

Next, we quantified the direction and magnitude of each gene's expression biases. For each sample, we calculated a gene's allelic bias as the proportion of total read counts with the LG/J haplotype ( $L_{bias}$ ) or the SM/J haplotype ( $S_{bias}$ ). Using the mean allelic biases of each F<sub>1</sub> reciprocal cross, we constructed Parentof-Origin Effect (**POE**) and Allelic Genotype Effect (**AGE**) scores per gene as follows:

$$POE = mean(SxL L_{bias}) - mean(LxS L_{bias})$$

615 
$$AGE = \frac{(mean(LxS L_{bias}) + mean(SxL L_{bias})) - (mean(LxS S_{bias}) + mean(SxL S_{bias}))}{2}$$

POE scores range from -1 (completely maternally expressed) to +1 (completely paternally expressed).
Similarly, AGE scores range from -1 (completely SM/J expressed) to +1 (completely LG/J expressed).

618 Scores of 0 for both indicate biallelic expression. Full GLM summary statistics and POE/AGE scores for

each tissue-by-context analysis are provided in **Supplemental Tables S2 - S4**.

## 620 Significance Thresholds

A crucial consideration for ASE analyses is how to best adjust for multiple tests. Allelic expression biases are often correlated for genes within and between imprinted domains (Edwards and Ferguson-Smith 2007) and for genes controlled by the same regulatory element with a functional variant (Cavalli et al. 2016). This ensures any tests performed on those genes are also correlated, breaking independence assumptions. We addressed this challenge by using a permutation approach (Westfall and Young 1993) to estimate our statistical and biological significance thresholds.

627 For each tissue-by-context cohort, we generated a stable null distribution of likelihood ratios (LR) for both 628 AG and PO terms. We randomly shuffled the allele-specific read counts for all genes and reran our 629 analyses over several iterations. After each iteration, we calculated the change in mean LR quantiles for 630 both terms in the full permuted dataset. Quantiles corresponded to one percent increments of the LR 631 distribution. We added new iterations until the null model met our "stability" criteria: the mean LR quantile 632 differences for both AG and PO terms fluctuated by less than [0.001] for 10 consecutive iterations (mean 633 = 51 iterations) (Supplemental Figure S19). Comparisons between the real and permuted datasets are 634 provided for p-values, POE/AGE scores, and likelihood ratios (Supplemental Figures S20 – S22).

Next, we built an empirical cumulative distribution function (ECDF) from each term's permuted p-values for each tissue-by-context analysis (**Supplemental Figures S23**). We fit each term's raw p-values to its respective ECDF to compute the adjusted p-values, i.e. the proportion of tests from the permuted null model that are more extreme (smaller p-values) than the test from the real model. We also calculated the 5<sup>th</sup> and 95<sup>th</sup> quantiles of the permuted POE/AGE scores for each term in each tissue-by-context analysis. We set our critical threshold as the more extreme value. We deemed an adjusted p-value  $\leq$  0.05 as statistically significant and real POE/AGE scores beyond the critical threshold as biologically significant.

642	Genes with significant PO term p-values and POE scores were considered to have parent-of-origin
643	dependent ASE. Similarly, genes with significant AG term p-values and AGE scores were considered to

#### 644 have sequence dependent ASE (**Supplemental Figures S24, Supplemental Table S5**).

#### 645 Characterizing ASE profiles: tissue-independent, tissue-dependent, and context-dependent

To examine how ASE is influenced by tissue type and/or environmental context, we characterized the expression profiles of the significant ASE genes across our 27 tissue-by-context analyses (3 tissues x 9 diet-by-sex contexts). In each analysis, a gene could be expressed in one of three ways: significantly biased, expressed with no allele-specific bias (biallelic), or simply not expressed. We sorted the significant ASE genes of both classes into three expression profiles based on the following criteria.

651 Tissue-independent ASE genes have a significant bias in every tissue where they are expressed. Within 652 a tissue, the gene must have a significant bias in  $\geq 5$  of the 9 diet-by-sex contexts (i.e. most, but not all, 653 environmental contexts). This flexibility allows for genes that may have true ASE but were excluded due 654 to our stringent minimum sample size requirements; these genes could appear as biallelically expressed 655 since they still pass our minimum read depth requirements. Tissue-dependent ASE genes have a 656 significant bias in some tissues and no bias in others. The gene must have a significant bias in >5 of the 657 9 contexts in one or two tissues, but biallelic expression in >5 of the 9 contexts in the other tissue(s). Both 658 profiles allowed for genes to not be expressed in some tissues, as we wanted to deliberately distinguish 659 between tissue-specific gene expression (not expressed in certain tissues) and tissue-dependent ASE 660 (biased in certain tissues). Finally, context-dependent ASE genes have a significant bias only in certain 661 environmental contexts and no bias elsewhere. The gene must have a significant bias in ≤5 of the 9 662 contexts within a tissue, but biallelic expression in the other contexts and/or tissues. All significant ASE 663 genes of both classes fit into one of these expression profiles; no genes were unclassified.

#### 664 *Evaluating environmental context-dependency*

665 Once we identified context-dependent genes, we evaluated how sex and/or dietary environment shape 666 their ASE patterns. For each context-dependent gene, we calculated each sample's allelic bias as the 667 proportion of total read counts with the LG/J haplotype (L<sub>bias</sub>) or the SM/J haplotype (S<sub>bias</sub>). We

668 constructed individualized Parent-of-Origin Effect (POE) or Allelic Genotype Effect (AGE) scores per 669 sample per gene by modifying the above equations as follows:

670 
$$Individual POE = \begin{cases} mean(SxL L_{bias}) - L_{bias}, & cross = LxS \\ L_{bias} - mean(LxS L_{bias}), & cross = SxL \end{cases}$$

671 Individual AGE = 
$$\begin{cases} \frac{(L_{bias} + mean(SxL L_{bias})) - (S_{bias} + mean(SxL S_{bias}))}{2}, & cross = LxS\\ \frac{(mean(LxS L_{bias}) + L_{bias}) - (mean(LxS S_{bias}) + S_{bias})}{2}, & cross = SxL \end{cases}$$

Individualized POE scores range from -1 (maternally expressed) to +1 (paternally expressed). Similarly,
individualized AGE scores range from -1 (SM/J expressed) to +1 (LG/J expressed). Scores of 0 for both
scores indicate biallelic expression or that the gene was not expressed in that sample.

Next, we used ANOVA models to test whether a gene's allelic biases (individualized POE/AGE scores) were influenced by sex, diet, and/or their interaction. We considered FDR-corrected p-values  $\leq 0.1$  to be significant (**Supplemental Table S6**). For genes with significant diet-by-sex interactions, we conducted Tukey's post-hoc tests to identify significant differences among diet-by-sex cohorts (adjusted p  $\leq 0.05$ ) (**Supplemental Table S7**).

# 680 Imprinted gene list

We defined a significant ASE gene as "canonically imprinted" if it appeared in the GeneImprint mouse database (https://www.geneimprint.com, as of May 2020) and/or a PubMed search of the gene name (or synonyms) and imprinting-related terms.

684 Pyrosequencing

We randomly selected 13 genes to validate based on their allele-specific expression profiles (i.e. tissueindependent, tissue-dependent, context-dependent, switches bias direction). We prioritized genes that were highly expressed and statistically significant (for context-dependent examples), but excluded those with high expression level variance between biological replicates. For each gene, we identified the strainspecific SNPs within exons and designed primer sets to flank these variants using Geneious Prime 2020.0.4 (https://www.geneious.com) (Kearse et al. 2012). Wherever possible, target regions were 150-

691 200bp long and spanned an exon-exon junction to avoid genomic DNA contamination. We verified the 692 specificity of each primer set *in silico* with Geneious and *in vitro* with PCR and Sanger sequencing. All 693 primer sequences are provided in **Supplemental Table S8**.

694 We extracted total RNA from the HYP, WAT, and LIV of one mouse per F<sub>1</sub> reciprocal cross (LxS and 695 SxL) in each diet-by-sex cohort using the RNeasy Lipid Tissue Mini Kit (QIAGEN). Next, cDNA of each 696 gene target was reverse-transcribed and PCR amplified with the PyroMark OneStep RT-PCR kit 697 (QIAGEN) using one biotinylated (reverse) and one non-biotinylated (forward) primer. The biotinylated 698 single-stranded PCR products were then purified with Streptavidin Sepharose High Performance beads (Cytiva) and hybridized to sequencing primers (same as forward) on the PyroMark vacuum prep 699 700 workstation (QIAGEN). Finally, we performed pyrosequencing with the Allele Quantification program on 701 the PyroMark Q24 system (QIAGEN). The pyrosequencing reaction emits a light signal as it builds the 702 DNA fragment, which appears on the pyrogram output as a peak whose height is proportional to how 703 many nucleotides were incorporated at that base. From these peaks, the PyroMark Q24 software 704 quantified the allelic ratio of the variable position(s) in each gene's assay. We calculated the mean allelic 705 ratios of each variant in each tissue-by-context cohort.

## 706 DATA ACCESS

All raw RNA-Sequencing data generated in this study have been submitted to the NCBI BioProject
 database (https://www.ncbi.nlm.nih.gov/bioproject/) under accession number PRJNA753198.

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