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X chromosome-dependent disruption of placental regulatory networks in hybrid dwarf hamsters

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Abstract

Embryonic development in mammals is highly sensitive to changes in gene expression within the placenta. The placenta is also highly enriched for genes showing parent-of-origin or imprinted expression, which is predicted to evolve rapidly in response to parental conflict. However, little is known about the evolution of placental gene expression, or if divergence of placental gene expression plays an important role in mammalian speciation. We used crosses between two species of dwarf hamsters (*Phodopus sungorus* and *Phodopus campbelli*) to examine the genetic and regulatory underpinnings of severe placental overgrowth in their hybrids. Using quantitative genetic mapping and mitochondrial substitution lines, we show that overgrowth of hybrid placentas was primarily caused by genetic differences on the maternally inherited *P. sungorus* X chromosome. Mitochondrial interactions did not contribute to abnormal hybrid placental development, and there was only weak correspondence between placental disruption and embryonic growth. Genome-wide analyses of placental transcriptomes from the parental species and first- and second-generation hybrids revealed a central group of co-expressed X-linked and autosomal genes that were highly enriched for maternally biased expression. Expression of this gene network was strongly correlated with placental size and showed widespread misexpression dependent on epistatic interactions with X-linked hybrid incompatibilities. Collectively, our results indicate that the X chromosome is likely to play a prominent role in the evolution of placental gene expression and the accumulation of hybrid developmental barriers between mammalian species.

Keywords: gene expression; reproductive isolation; genomic imprinting; Phodopus

Introduction

Developing mammalian embryos depend on the extra-embryonic placenta for a broad array of functions including hormone production, immunity, and as a conduit for maternal nutrients and gas exchange (Reik et al. 2003; Levy 2007). Normal intrauterine development in humans and mice depends on the tightly controlled placental expression of a diverse set of genes (Constancia et al. 2002; Levy 2007; Plasschaert and Bartolomei 2014). Placental gene expression has also likely played an important role in the evolution of mammalian development (Haig 1996; Capellini et al. 2011; Kaneko-Ishino and Ishino 2019). Indeed, much of the phenotypic diversity across mammalian species is thought to have evolved by changes in gene expression during critical stages of development (King and Wilson 1975; Carroll 2008; Sears et al. 2015). However, relatively little is known about the broader genomic organization, functional integration, and evolution of placental gene expression networks across species (Al Adhami et al. 2015).

The placenta is characterized by two unusual regulatory phenomena that likely play critical roles in its evolution. First, the placenta is highly enriched for genes showing monoallelic expression due to epigenetic silencing of one parental allele (i.e., genomic imprinting, Morison et al. 2005; Hudson et al. 2010; Babak et al. 2015). Genomic imprinting is thought to have evolved to help resolve fitness conflicts between maternally and paternally inherited alleles (i.e., kinship or parental conflict theory, Haig 2000). While perhaps only approximately 100-200 autosomal genes showed strongly imprinted expression across tissues (Babak et al. 2015), disruption of genomic imprinting has emerged as an important cause of congenital disorders in humans (Hirasawa and Feil 2010; Lee and Bartolomei 2013) and as a potential driver of reproductive barriers between species (Crespi and Nosil 2013; Wolf and Brandvain 2014). Second, some mammals also show imprinted paternal X chromosome inactivation in extra-embryonic tissues (i.e., imprinted XCI; Heard and Disteche 2006), representing a striking deviation from random XCI found in most somatic cells of placental mammals (Lyon 1961; Dupont and Gribnau 2013). The X chromosome in general, and imprinted XCI in particular, has been shown to play important roles in placental development (Hemberger 2002; Mcgraw et al. 2013). Moreover, the mouse X chromosome appears enriched for genes preferentially expressed in the placenta (Khil et al. 2004).

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Resolution of genetic conflict may explain the origin of genomic imprinting in mammals (Haig 2000). However, theory also predicts that once imprinting is established adaptation among interacting genes can drive the evolutionary expansion of regulatory networks of similarly imprinted genes (Wolf and Hager 2006; Wolf and Brandvain 2014; Patten et al. 2016; O'Brien and Wolf 2017). Given the relative scarcity of autosomal imprinting overall (Babak et al. 2015), the X chromosome is expected to harbor the majority of genes showing imprinted (maternal) placental expression in species with imprinted XCI. Thus, co-evolutionary interactions between the maternal X chromosome and maternally expressed autosomal genes should be relatively common within placental regulatory pathways in imprinted XCI species. Despite these predictions, the overall importance of the X chromosome to the evolution of placental gene expression remains unclear. Many molecular genetic studies on the placenta have focused on established disease models or targeted genetic manipulations of imprinted autosomal genes (e.g., gene knockdowns or knockouts; Sanli and Feil 2015), revealing fundamental insights into the mechanisms and functional consequences of genomic imprinting in the placenta and other tissues. In parallel, metaanalyses of expression data have revealed that clusters of imprinted autosomal genes appear to fall within larger networks of co-expressed genes that include both imprinted and bi-allelically expressed loci (Varrault et al. 2006; Al Adhami et al. 2015). The extent and functional relevance of such regulatory networks remain unclear, but the emerging model of genome-wide networks of imprinted and nonimprinted genes represents a conceptual shift from the view of imprinting controlled primarily through local cis-regulatory effects (Patten et al. 2016).

Parent-of-origin effects for abnormal embryonic and placental growth are common in mammalian hybrids (Vrana 2007; Brekke and Good 2014), suggesting that hybrid systems may provide powerful models for understanding how the evolution of gene expression impacts placental development. Here, we focus on crosses between two closely related species of dwarf hamsters (Phodopus sungorus and Phodopus campbelli) that yield strong parent-of-origin growth effects in reciprocal F1 crosses (Brekke and Good 2014). Extensive placental overgrowth and disrupted embryonic development occur in hybrid crosses when the mother is P. sungorus (female P. sungorus × male P. campbelli; hereafter S × C with uppercase used to denote placental overgrowth), often resulting in hybrid and maternal death during birth. The reciprocal cross results in normal embryonic development (female P. campbelli \times male P. sungorus; hereafter c \times s with lowercase used to denote normal placental growth), although adult hybrid males are smaller (Brekke and Good 2014) and completely sterile (Safronova et al. 1999; Bikchurina et al. 2018). Intrinsic postzygotic reproductive isolation (i.e., hybrid inviability or sterility) generally tends to be asymmetric in reciprocal crosses between closely related species due to incompatible interactions at sex-linked or imprinted loci (Turelli and Moyle 2007). Although the genetic architecture of hybrid placental growth has not been determined in dwarf hamsters, massively overgrown F_1 hybrid $S \times C$ placenta do show extensive disruption of gene expression pathways that are highly enriched for embryonic growth and genomic imprinting (Brekke et al. 2016). Building on these previous studies, we combine quantitative genetic and transcriptomic analyses to test the hypothesis that the X chromosome plays a central role in the evolution of placental gene expression, embryonic development, and reproductive barriers between species.

Materials and methods Animals and experimental crosses

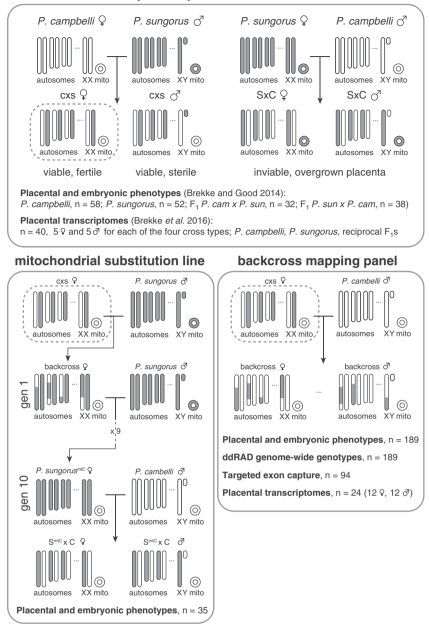
Experimental animals were drawn from established colonies of wild-derived *P. campbelli* and *P. sungorus* at the University of Montana (Brekke and Good 2014), which were originally established by Kathy Wynne-Edwards (Scribner and Wynne-Edwards 1994). Colonies were maintained as outbred, though overall inbreeding levels are high (Brekke *et al.* 2018). All breeding experiments were done in compliance with the University of Montana Institutional Animal Care and Use Committee regulations (animal use protocols 039-13JGDBS-090413 & 050-16JGDBS-082316).

We previously reported results from experimental crosses within and between P. campbelli and P. sungorus used to examine late-term placental and embryonic phenotypes (Brekke and Good 2014) and placental transcriptomes (n = 40 placental transcrip)tomes, five males and five females for each cross type; Brekke et al. 2016). Here we combined these results with new data from two additional genetic crossing experiments used to evaluate the role of the mitochondrial and nuclear genomes in the genetic basis of asymmetric hybrid placental and embryonic overgrowth (Figure 1). First, we generated mitochondrial substitution lines wherein c × s hybrid females were crossed to P. sungorus males for 10 additional backcross (BC) generations. This crossing scheme is predicted to recover hamsters that are ~99.9% P. sungorus across the nuclear genome but retain the mitochondria of P. campbelli (P. sungorus^{mtC}). Tenth-generation P. sungorus^{mtC} females were crossed to P. campbelli males to test for F1 overgrowth $(S^{mtC} \times C)$, thereby mimicking the overgrown $S \times C$ hybrid across the nuclear genome while substituting the P. sungorus mitochondria for P. campbelli mitochondria. Second, we crossed $F_1 c \times s$ hybrid females to P. campbelli males to generate a BC panel of lateterm embryos and placentas. In the context of genetic elements with sex-limited inheritance or expression, these BC hybrids have the same paternal contribution found in overgrown $S \times C F_1$ hybrids, while varying the species origin of maternally inherited alleles

All advanced crosses were conducted through $c \times s$ hybrid females as $S \times C F_1$ hybrids generally do not survive birth (Brekke and Good 2014) and $c \times s$ males are completely sterile (Safronova *et al.* 1999; Ishishita *et al.* 2015; Bikchurina *et al.* 2018). For both crossing experiments, females were sacrificed at late gestation and offspring placentas and embryos were collected, weighed, and snap-frozen on dry ice. Embryos were developmentally scored following (Brekke and Good 2014) to ensure that all offspring were in the final 4 days of gestation corresponding to Theiler's Stages 24–27 (Theiler 1972). Developmental abnormalities such as embryonic fluid accumulation (edema) and embryo reabsorption were noted, and embryo and placenta weights were assessed with stepwise model selection and adjusted for variation in edema, litter size and Theiler stage using simple linear models as implemented in JMP (v12).

Genotyping

We genotyped 189 BC individuals (91 females and 98 males) and our original colony founders (14 P. campbelli and 11 P. sungorus) using double digest restriction site-associated DNA sequencing (ddRAD-seq; Peterson et al. 2012) with minor modifications. Genomic DNA was extracted from frozen embryos with a Machery-Nagel Nucleospin Tissue DNA extraction kit (catalog number 740952) following the manufacturer's protocol, except that 5 μ l RNase-A was added to the column and incubated for 15 minutes at room temperature. We then digested 1 μ g of



reciprocal species intercrosses

Figure 1. Summary of genetic crosses and experiments. Reciprocal species intercrosses between *P. sungorus* (gray chromosomes) and *P. campbelli* (white chromosomes) result in asymmetric placental overgrowth and inviability. Only females with a *P. campbelli* mother and *P. sungorus* father are both viable and fertile ($c \times s$, indicated with a dashed box). The first generation cross with normal sized placentas is indicated with lowercase letters ($c \times s$), and the reciprocal cross with overgrown placentas is indicated with uppercase letters ($S \times C$). A mitochondrial substitution line was created through 10 generations of backcrossing hybrid females to a *P. sungorus* male, resulting in females with *P. sungorus* nuclear genomes and *P. campbelli* michondria (S^{mtC}) that were then crossed to *P. campbelli* males. The BC mapping panel was created by a single generation of backcrossing a fertile hybrid female to a *P. campbelli* male, resulting in offspring showing a range of placenta phenotypes. Data types collected for each experiment are reported in the corresponding panels.

genomic DNA using the high-fidelity restriction enzyme *SbfI-HF* (New England BioLabs, catalog number R3642L), followed by *MspI* (New England BioLabs, catalog number R0106L) both with the CutSmart buffer (New England BioLabs). Libraries were prepared with a dual barcoding scheme incorporating both Illumina indexes and inline barcodes to uniquely identify each sample (Peterson *et al.* 2012). Size selection of adapter-ligated fragments (200–500 bp) was done with Agencourt AMPure XP beads. Both the size selection and PCR amplification were done prior to sample pooling to assure more even representation across samples.

Combined sample pools were sequenced on 50% of an Illumina HiSeq 2500 lane in rapid-run mode and then on 50% of a lane of Illumina Hiseq 2500 in normal mode at the University of Oregon Genomics and Cell Characterization Core Facility. All samples were sequenced in both lanes and combined for subsequent analyses. We also independently determined the sex of embryos using a PCR assay of the Y-linked Sry gene as described previously (Brekke *et al.* 2016).

RAD libraries were cleaned and demultiplexed with Stacks (v1.20; process_radtags parameters -e sbfl -renz_2 mspl -r -c -q; Catchen et al. 2013). An initial list of unique RADtags from both read pairs was generated using ustacks (-H -r -d) using data from one female founder from each species. RADtag reference libraries were then generated using cstacks (-n 4). Reads from all the colony founders were aligned to the RADtag reference library with bwa-mem (v0.7.9a; Li and Durbin 2009) and single-nucleotide variants (SNVs) were called with the GATK HaplotypeCaller (v3.1-1, -stand_call_conf 30; Van Der Auwera et al. 2013). All SNVs that were polymorphic within a species in our colony founders were filtered out using GATK selectVariants (v3.1-1). BC individuals were genotyped at the ascertained SNVs sites using GATK UnifiedGenotyper (v3.1-1; -stand_call_conf 30; Van Der Auwera et al. 2013).

We also used an exon capture experiment to anchor placentalexpressed genes onto the *Phodopus* genetic map (described below). We designed custom hybridization probes to target 9,756 fixed SNVs between *P. campbelli* and *P. sungorus* ascertained from species-specific transcriptomes (Genbank BioProject PRJNA306772 and DDBJ/EMBL/GenBank Accessions GEVA00000000 and GEVB00000000; Brekke *et al.* 2016). Exon boundaries were annotated for each transcript with a local BLAT search (Kent 2002) against the golden hamster (*Mesocricetus auratus*) draft reference genome (The Broad Institute Genome Assembly & Analysis Group, MesAur1.0). For each gene, we selected 1-2 SNVs located furthest from inferred exon boundaries and included probes matching both alternative bases to avoid species bias. Capture baits were manufactured by MYcroarray.

We selected 94 individuals (44 males and 50 females) from the BC panel and prepared Illumina sequencing libraries following the Meyer-Kircher protocol (Meyer and Kircher 2010). Ten cycles were used during the indexing PCR step. The indexed libraries were then combined into four pools and target enriched following the MyBaits-1 Custom Target Capture protocol and using mouse C_oT-1 DNA supplied by the manufacturer as a blocking agent. The four captured pools were then reamplified with 20 PCR cycles, quantified with a Kappa qPCR quantification kit (catalog number KK4824), and pooled for Illumina sequencing. Enriched libraries were initially sequenced at the University of Montana Genomics Core on an Illumina MiSeq 75-bp paired-end sequencing, and followed by one lane of Illumina HiSeq 2500 100-bp single-end sequencing at the University of Oregon Genomics and Cell Characterization Core Facility. Raw sequence reads were adapter trimmed with Cutadapt (v1.6; -O 5 and -e 0.1; Martin 2011) and quality-filtered with Trimmomatic (v0.3.2; LEADING : 5, SLIDINGWINDOW : 4:15, MINLEN: 36, and HEADCROP : 13; Bolger et al. 2014). Filtered reads were then aligned to published transcriptome assemblies (Brekke et al. 2016) and the targeted SNVs were genotyped with GATK HaplotypeCaller (v3.1-1; stand_call_conf 30.0 -stand_emit_conf 30) and filtered (selectvariants -restrictAllelesTo BIALLELIC -select "QD > 10.0") so that only high-quality genotypes were used for estimating the location of each gene.

Quantitative genetic analysis

We constructed a genetic map using RADtag SNVs identified between the *P. campbelli* and *P. sungorus* colony founders and the program R/qtl (v1.45; Broman and Sen 2009). Putative X-linked RADtags were identified as markers that were heterozygous or homozygous for *P. campbelli* genotypes in females and always homozygous *P. campbelli* or *P. sungorus* in males. To estimate the map, we removed two BC individuals with low-sequencing coverage, identified putative X-linked markers based on Hardy– Weinberg expectations, and dropped all autosomal markers that were genotyped in fewer than 177 individuals (95%). We formed linkage groups and ordered the markers on each linkage group with the ripple(), compareorder(), and switch.order() functions until each linkage group was a short as possible. Then we sequentially dropped each marker to see if the likelihood of the map improved. Once all poor quality markers were removed, we repeated the ripple(), compareorder(), and switch.order() functions until the likelihood was maximized. The linkage groups in the final map were ordered by descending length in centiMorgans (cM).

We then used R/qtl to test for single-quantitative trait loci (QTL) associated with the variation in BC embryo and placental weight using the extended Haley-Knott method with imputation (Haley and Knott 1992; Feenstra et al. 2006). Next, we incorporated sex as a covariate and re-estimated QTL for both phenotypes. For all of single-QTL scans, we used 10,000 permutations to estimate genome-wide significance thresholds for autosomes and 337,364 permutations for the X chromosome. Finally, we used the QTL identified in the first two analyses as additive cofactors and re-scanned for additional QTL that were contingent on the presence of the earlier identified QTL (Broman and Sen 2009). We used 1000 permutations for autosome-autosome interactions, 1687 permutations for autosome-X interactions, and 113,815 permutations for X-X interactions to establish significance thresholds. QTL intervals were established based on 95% Bayesian confidence intervals, and the proportion of phenotypic variance explained by QTL was estimated as $1 - 10^{-\frac{2}{n}LOD}$ (Broman and Sen 2009).

Expressed genes were then integrated onto the genetic map by comparing captured SNV genotypes to RADtag genotypes. Following Brekke et al. (2019), we counted shared genotypes between each RADtag and each gene across all individuals, placing genes at the location of the RADtag with which they shared the most genotypes. In the event of a tie between multiple RADtags, the gene was placed at the proximal map location and only genes sharing at least 90% of genotypes with at least one RADtag were placed on the map. Given the low number of recombination events and the high number of genes, these RADtag-anchored positions represent coarse genetic locations for each gene. Instances where multiple genes were associated with a single RADtag were treated as a single unordered linkage block. Once integrated, likely genotyping errors in the capture data were identified using calc.errorlod() and the highest errors were extracted with top.errorlod() and a very strict cutoff of 1 such that even moderately questionable genotypes were identified. The genotypes of these sites were removed and then filled in with the fill.geno() function using the imputation method, which imputes the missing genotypes from the surrounding sites for each individual. These corrected genotypes were used for evaluating imprinting status of select candidate genes (see below).

Gene expression analyses

We chose 24 placentas from our BC mapping panel for genomewide expression analysis using RNA-seq (Wang *et al.* 2009), including six males and six females with large placentas (mean = 0.232 ± 0.010 g) and six males and six females with normal sized placentas (mean = 0.140 ± 0.008 g). RNA was extracted from whole frozen placenta with an Omega Bio-tek E.Z.N.A. Total RNA Kit I (catalog number R6834) including a DNase digestion following the manufacturer's protocol. All RNA samples were checked for quality and concentration on the bioanalyzer and all samples used had RNA integrity numbers >8.0. RNA-seq libraries were constructed from 2 µg of input RNA with the Agilent Sure-Select Strand-Specific RNA-seq Kit (catalog number G9691B) following the manufacturer's recommendations. Libraries were amplified with 14 cycles of PCR, and pooled based on a Kappa qPCR Quantification Kit (catalog number KK4824). The pooled libraries were sequenced with two lanes of Illumina HiSeq2500 100-bp single-end sequencing.

RNA-seq data were processed as described previously (Brekke et al. 2016). Briefly, Illumina adapters were trimmed off reads with Cutadapt (v1.6; -O 5 -e 0.1; Martin 2011) and quality trimmed with Trimmomatic (v0.3.2; SE -phred 33 LEADING: 5 SLIDINGWINDOW: 4:15 HEADCROP: 13; Bolger et al. 2014). While an initial draft of the P. sungorus genome has been generated using Illumina shotgun sequencing, current annotation and assembly quality remains insufficient for reference-guided transcriptome analyses (Bao et al. 2019). Therefore, reads were aligned with bowtie2 (v2.2.3; Langmead and Salzberg 2012) to a published de novo placental transcriptome assembly (Genbank BioProject PRJNA306772 and DDBJ/EMBL/GenBank Accessions GEVA00000000 and GEVB00000000; Brekke et al. 2016), and filtered for potentially chimeric transcripts using draft genomic resources by excluding 1422 "genes" with exons that multiply mapped to different contigs. To evaluate expression level, we created a table of counts at the gene level using featureCounts (v1.4.2; Liao et al. 2014), which counted fragments (-p) and discarded reads with too long an insert (-P) or are chimeric (-C) or have a mapping quality (-Q) below 20. This table of counts was normalized with the trimmed mean of m-values (TMM) method (Robinson and Oshlack 2010).

We used the WGCNA package (version 1.68; Langfelder and Horvath 2008) to infer weighted gene co-expression networks using expression data from previously published parental and F_1 genotypes (Brekke et al. 2016) and the newly generated BC. This network approach uses adjacency correlation matrices to identify hierarchical clusters of co-expressed genes (Zhang and Horvath 2005), enabling the reduction of complex clusters into representative expression profiles (i.e., module eigengenes) defined as the first component of a principle component analysis. A scale-free topology index was determined by soft thresholding (Supplementary Figure S1), which was then used to automatically detect signed, Pearson correlated modules via dynamic cutting. The signed module method splits otherwise correlated genes with increases in expression into separate modules from those with decreases in expression, which allowed us to evaluate upregulated gene sets independently from downregulated gene sets. We merged similar modules using a threshold of 0.25, which combines modules with a correlation of 0.75 or greater into a single module. For each module, we tested for correlations between the module eigengene and placental and embryonic weights. For modules showing significant correlations after correction for multiple testing, we then retested associations using a more stringent ANOVA model that controlled for developmental stage and sex.

Each module was assessed with a binomial exact test (R/stats package 3.6.1) for enrichment of candidate imprinted genes previously identified based on patterns of allele-specific expression in the placenta (Brekke *et al.* 2016) and for X-linked genes. Network connectivity was determined through pairwise correlation between genes, with *P*-values corrected with the *q*-value package (version 2.18.; Storey *et al.* 2019), using a false discovery rate threshold of 0.05. We then counted the number of additional genes significantly correlated with each gene within the module. Hub genes were defined as the top 5% most connected genes in each module. Overlap between F₁ and BC modules was determined by comparing gene lists to get counts of shared genes. We also evaluated module conservation by comparing how strongly each gene was correlated to each module across data sets. To connect module conservation to

phenotypes, we compared the concordance between each gene and placenta weight across F₁ and BC datasets using a bivariate correlation as implemented in JMP (v12).

Genotypes at each gene placed on the genetic map were inferred by evaluating genotypes at flanking RAD markers. If a gene was placed in the same linkage block (cM location) as a RADtag, the marker genotype was used for the gene. Likewise, if the gene was placed between two markers with the same genotype, the concordant marker genotypes were used for the gene. If flanking RADtag genotypes were discordant, the genotype at the gene was treated as unknown. We then tested for expression interactions (EIs) between the X chromosome and diploid autosomal genotypes within the BC as follows. Autosomal genotypes were imputed for the 23 BC individuals with transcriptome data. EI scores were then calculated for all mapped autosomal genes by comparing the difference in fold change in expression of that gene for the two possible autosomal genotypes (i.e., homozygous for the P. campbelli allele or heterozygous) dependent on the genotype of the maternal X chromosome (P. campbelli or P. sungorus). To generate EI scores for each gene, normalized gene expression count tables generated during WGCNA were used to calculate log2-fold changes between alternative maternal X chromosome genotypes and the two genotypes at each autosomal locus. We excluded unmapped autosomal genes and genes with imputed genotypes for fewer than three individuals for each of the four X-by-autosome genotypic classes. We calculated the absolute value of EI scores, where a value of zero indicates no difference between the two autosomal genotypes when inheriting different maternal X chromosomes and a value of one indicates a onefold difference in expression between the two autosomal genotypes (i.e., a candidate X-autosome EI). We also considered a polarized (i.e., nonabsolute value) version of the statistic where positive values reflected greater change when maternal X and autosomal alleles genotypes were discordant.

EI was calculated as follows:

$$\begin{split} EI &= abs \left(log2 \left(\frac{mean \; expression_{X_{cam}Gene_{CC}}}{mean \; expression_{X_{cam}Gene_{CC}}} \right) \\ &- log2 \left(\frac{mean \; expression_{X_{cam}Gene_{SC}}}{mean \; expression_{X_{sun}Gene_{SC}}} \right) \right) \end{split}$$

$$\end{split} polarized EI &= log2 \left(\frac{mean \; expression_{X_{cam}Gene_{CC}}}{mean \; expression_{X_{sun}Gene_{CC}}} \right) \\ &- log2 \left(\frac{mean \; expression_{X_{cam}Gene_{CC}}}{mean \; expression_{X_{sun}Gene_{SC}}} \right) \end{split}$$

where

x_{sun} = samples with a P. sungorus maternal X chromosome;

 $x_{cam} =$ samples with a P. campbelli maternal X chromosome;

Gene_{CC} = samples that are homozygous for the *P. campbelli* allele at the gene for which the EI score is calculated;

 $\mbox{Gene}_{\rm SC} = \mbox{samples}$ that are heterozygous at the gene for which the EI score is calculated.

Results

Mitochondrial interactions were not a major factor in extreme parent-of-origin hybrid overgrowth

If hybrid placental overgrowth is caused by deleterious interactions between the maternal P. sungorus mitochondrial genome and the P. campbelli nuclear genome, then crossing females from the P. sungorus^{mtC} mitochondrial substitution lines to P. campbelli males should recover normal development. Alternatively, if heterospecific mitochondria have no effect on hybrid growth, then S^{mtC} ×C hybrids should be similar in size to overgrown S×C hybrids. In support of the second hypothesis, $S^{mtC} \times C$ placentas were found to be extremely large ($F_{4,213} = 106$, P < 0.001, ANOVA), but not statistically different from $S \times C$ hybrids based on a Tukey's honest significant difference (HSD) test (Supplementary Figure S2). We note that low levels of residual autosomal heterozygosity could theoretically mask mitochondrial incompatibilities in the 10th generation mitochondrial substitution P. sungorus^{mtC} lines. However, such residual variation cannot also explain why the substitution lines did not recover normal placental development in the F_1 hybrid crosses. On balance, these results strongly indicate that heterospecific mitochondrial interactions are not a major factor in extreme parent-of-origin hybrid overgrowth in dwarf hamsters.

The quantitative genetic basis of extreme parentof-origin hybrid overgrowth

We generated a BC panel of 189 late-term embryos and placentas sampled from 38 litters. Using 1215 ddRAD SNVs, we constructed a 1231.7 cM genetic map that comprised 14 major linkage groups (Supplementary Figure S3 and Supplementary Table S1). The karvotype of P. sungorus comprised 14 chromosomes (2N = 28;Romanenko et al. 2007), suggesting that each of our major linkage groups correspond to an individual chromosome. The X chromosome was inferred to have the shortest overall genetic distance (35.5 cM or 2.9% of the genetic map), while it appears mediumsized in ranked karyotype analyses (i.e., middle 30%; Gamperl et al. 1977; Romanenko et al. 2007) and comprises ~10% of the haploid female karyotype (Haaf et al. 1987). The Phodopus X chromosome is metacentric with an Xp arm that has been described as heterochromatic (Gamperl et al. 1977) and nonrecombinant in females of both species and in $c \times s F_1$ hybrids (Bikchurina et al. 2018). Our inferred genetic map was consistent with strong repression of recombination on one end of the X chromosome in c × s females, albeit not complete repression as suggested in recent study that quantified signals of mismatch repair through immunolocalization of MLH1 (Bikchurina et al. 2018).

Using this genetic map, we then tested for QTL associated with variation in BC embryo and placenta weights (Figure 2). We observed a single QTL of large effect for placental weight on the X chromosome, with 52.3% of the observed variation in BC placental weight determined by the genotype of the maternally inherited X chromosome (Figure 2C). We estimated a QTL peak at 31.1 cM with a 95% Bayesian confidence interval between 29.6 and 32.6 cM. This X-linked QTL localized near the proximal boundary of where we also observed repressed recombination (Supplementary Figure S4), although the entire X chromosome exceeded a permutation-based significance threshold (P = 0.01). Male and female embryos inheriting a maternal P. sungorus X chromosome genotype at this QTL showed an ~60% increase in average placenta weight ($F_{1,179} = 178.4$, P < 0.0001, ANOVA; Figure 2C inset). No additional placental QTL was uncovered when using sex, developmental stage, and litter size as covariates, nor when using the X-linked QTL as a cofactor.

No QTL for embryo weight was recovered in our experiment (P = 0.05 permutation-based significance threshold; Supplementary Figure S5), despite considerable variation in BC embryo weights (Figure 2A) and significant overgrowth of S × C embryos when compared to normally developing cross-types (Brekke and Good 2014).

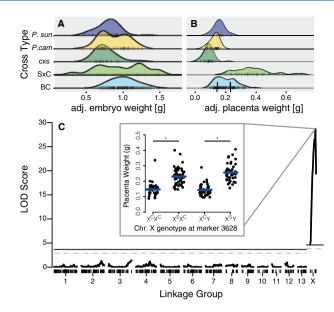


Figure 2. Phenotype distributions and QTL for placental size. Late term embryo size adjusted for Theiler stage and edema (A) and placenta size adjusted for Theiler stage (B) for *P. sungorus*, *P. campbelli*, reciprocal F₁s and 189 BC conceptuses. (C) Genetic mapping revealed a single QTL for placenta weight on the X chromosome. The *P. sungorus* X chromosome increases placenta weight by ~60% (inset, $F_{1,179} = 178.4$, $P \ll 0.0001$, stars indicate significant differences assigned by a Tukey's HSD test). Placenta weights were grouped by the genotype at marker 3,628. Genotypes are denoted with the maternal allele first followed by the paternal allele. Significance thresholds are denoted by solid (P = 0.01) and dashed (P = 0.05) horizontal lines.

Severe embryonic swelling or edema is common in $S \times C$ hybrids and appears to drive overall differences in embryo weights between overgrown S \times C hybrids and either species or c \times s hybrids (stepwise best model: embryo weight \sim developmental stage + edema; adjusted $r^2 = 0.40$, $F_{2,182} = 61.6$, $P \ll 0.0001$; adjusted embryo weight \sim cross-type $F_{3,182} = 0.74$, P = 0.5312, ANOVA; Supplementary Figure S6). However, BC placenta and embryo weights remained moderately correlated after adjusting for developmental stage and edema (adjusted $r^2 = 0.159$, $F_{1,184} = 36.0$, $P \ll 0.0001$, ANOVA; Supplementary Figure S7A), with males showing a stronger correlation than females (males: adjusted $r^2 = 0.257$, $F_{1,95} = 33.8$, $P \ll 0.0001$; females: adjusted $r^2 = 0.065$, $F_{1,88} = 7.15$, P = 0.0090, ANOVA). When we expanded our analysis of embryonic weights among genotypes to include the BC, we also detected a small but significant increase in embryo weight in the overgrown crosses after controlling for age and edema (adjusted $r^2 = 0.176$, $F_{4,376} = 21.1$, $P \ll 0.0001$, ANOVA; Supplementary Figure S7B). These apparent differences in embryonic growth were likely too subtle to detect in our QTL mapping experiment.

Networks of placental gene expression in firstgeneration reciprocal hybrids

We previously demonstrated extensive parent-of-origin dependent disruption of hybrid gene expression, with hundreds of genes significantly up- or downregulated in overgrown S \times C hybrid placenta relative to both species (i.e., transgressive expression; Brekke *et al.* 2016). In this previous study, examination of allele-specific expression revealed 88 candidate imprinted genes in the placenta overall, with 79 genes showing strong bias toward maternal expression (i.e., paternally imprinted). Notably, 68% of candidate imprinted genes (60 genes) showed transgressive expression in overgrown hybrids, suggesting a link between misexpression of autosomal genes with biased parent-of-origin expression and placental overgrowth. Imprinted XCI was not disrupted in F_1 hybrids. In contrast to a predominantly autosomal regulatory phenotype in F_1 hybrid placenta, our BC experiments indicated that $S \times C F_1$ hybrid placental overgrowth was likely caused by genetic incompatibilities exposed on the maternally inherited P. sungorus X chromosome (Figure 2C).

Motivated by these parallel observations, we sought to identify groups of co-expressed genes associated with overgrowth phenotypes exposed in both of our first- and second-generation hybrid models of placental overgrowth. We first used our published lateterm placental expression data from the parental species and reciprocal F₁ hybrids to construct weighted gene co-expression networks, removing one female S \times C sample during filtering (n = 39placental transcriptomes, 5 males and 5 females per cross-type; Supplementary Figure S8). We placed 11,392 genes (including 70 candidate imprinted genes) into 29 signed clusters, or "modules," of nonoverlapping gene sets. For each module, expression values were summarized with an "eigengene," or the principal component capturing the largest proportion of the variance in gene expression. We then assessed each module for mode of eigengene inheritance and association with placental phenotypes (Supplementary Table S2 and Supplementary Figure S9). Two key gene networks emerged from this analysis. One module was comprised of 565 genes that tended to be highly expressed in S × C hybrid placenta relative to all other genotypes. The eigengene value for this module was positively correlated with placental weights (Figure 3A), and included only one candidate imprinted gene (Figure 3B).

The other module comprised 1,160 genes that tended to show a lower eigengene value in S × C hybrid placenta (Figure 3C). Expression of this downregulated set was negatively correlated with placental weights (Figure 3C), and included nearly half (44%) of the downregulated transgressive genes identified by Brekke *et al.* (2016). Eigengene values for this module exhibited a stronger parent-of-origin mode of inheritance than the upregulated set (Figure 3C), and was positively correlated with candidate imprinted gene expression (Figure 3D). The downregulated module included over 50% of the candidate placental imprinted genes overall (36 of 70 genes, $P \ll 0.0001$; Supplementary Table S2). These findings mirror results from pairwise contrasts (Brekke *et al.* 2016) where overgrown S × C placentas showed an overall reduction in the expression levels of several putatively imprinted genes (Figure 4).

Networks of placental gene expression in secondgeneration backcrossed hybrids

To test for links between our BC QTL mapping experiment and emergent patterns of placental expression in F_1 hybrid models, we next analyzed 24 transcriptomes from BC placentas (12 large and 12 normal sized placentas; 11,396 genes placed in the network). One large female placenta was removed during outlier filtering (Supplementary Figure S10). WGCNA analysis of the BC transcriptomes revealed seven modules that were correlated with placenta weights. No clusters were significantly associated with embryo weights after controlling for developmental stage and sex (Supplementary Table S3). The recombinant genotypes within this BC sample allow us to more clearly differentiate

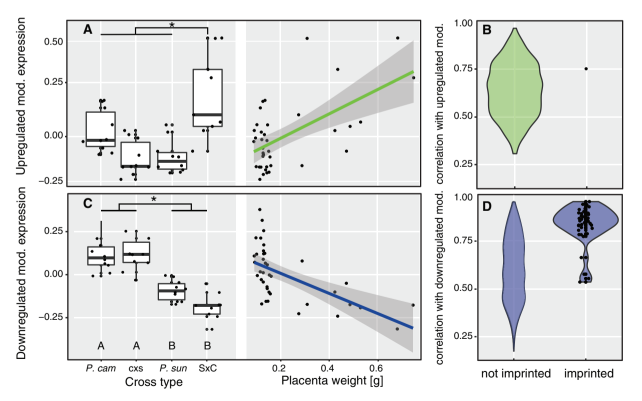


Figure 3. F_1 placenta size-associated gene expression modules. Eigengene gene expression values summarize the group of genes that are upregulated (A and B) and downregulated (C and D) in overgrown S × C placentas, shown as differing by cross-type (transgressive expression; stars indicate significance groups at P < 0.05 based on a Tukey's HSD test) and placental size (association with hybrid incompatibility phenotype controlling for Theiler stage and sex, P < 0.001, ANOVA, see Supplementary Table S2). The upregulated F_1 module was not enriched for imprinted genes (C; total genes, N = 565; imprinted genes, N = 1), whereas the downregulated F_1 module was enriched for highly connected candidate imprinted genes (Brekke *et al.* 2016) showing maternally biased expression (D; total genes, N = 1160; imprinted genes, N = 36). Data points rendered only for imprinted genes in (B and D).

disrupted expression in overgrown hybrid placenta versus species differences (P. sungorus versus P. campbelli) or interspecific hybridization per se (hybrids versus parental species). Consistent with this, only about one-third of the core genes included in the upand downregulated F_1 modules were captured in the seven placenta-associated BC modules (563 of 1,725 genes). However, most of these overlapping genes (>90%, 513 of 563) were derived from the downregulated $F_1 S \times C$ module (Supplementary Table S3). The BC module that was most strongly associated with placental weights (432 genes) tended to show lower summary expression in overgrown BC placenta (Figure 5A). This downregulated BC module was also highly enriched for autosomal imprinting (33

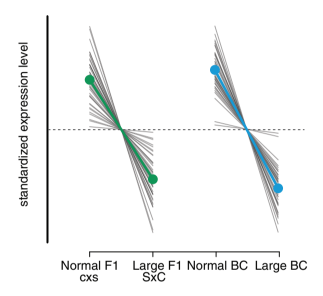


Figure 4. Reduction in candidate imprinted gene expression in overgrown F_1 and BC placentas. Each gray line represents the relative change in expression of a candidate imprinted gene between normal sized and large hybrid placentas. Genes were standardized by mean expression per gene (dotted line), and centered to display direction rather than magnitude of gene expression change. Both F_1 and BC experiments demonstrate reduced gene expression of genes showing parent-of-origin bias accompanying placental overgrowth (grand mean indicated by colored line).

candidate imprinted genes, binomial exact test $P \ll 0.0001$) and for genes in the downregulated $F_1 S \times C$ module (206 genes, binomial exact test $P \ll 0.0001$; Figure 4; Supplementary Table S3 and Supplementary Figure S11). In concordance with the overlap between gene lists, genes that were positively correlated with the F_1 downregulated module eigengene were also positively correlated with the BC downregulated module eigengene (Supplementary Figure S12), indicating module conservation across first- and second-generation hybrids.

Connectivity within scale-free expression networks is commonly defined by the extent to which the expression level of a given gene is correlated with the expression of other genes. We found that candidate imprinted genes were much more highly connected than nonimprinted genes within the downregulated, placenta-associated BC module (Figure 5B). Co-expression modules are usually characterized by a few highly connected "hub" genes (Ghazalpour et al. 2006; Mack et al. 2019). We found that the top 5% most connected (hub) genes in the downregulated module were involved in 60% (300) of the top 500 pairwise interactions (Figure 5C, hub interactions indicated by thicker lines and larger circles). Candidate imprinted genes were highly overrepresented as hub genes in this network-one-third of hub genes (8 of 21 genes, binomial exact test $P \ll 0.0001$) were candidate imprinted loci with maternally biased expression, including the top five most highly connected genes (Plxdc2, ProcR, Scara5, CD68, and Wnt4). Indeed, nearly half (238) of the top 500 pairwise correlations (Figure 5C, blue lines) involved at least one candidate imprinted gene (binomial exact test $P \ll 0.0001$). These highly connected, downregulated genes represented many core biological functions of the placenta, ranging from broadly expressed genes involved in growth and development to those with specialized placental function (Table 1).

Our transcriptome analyses revealed a central link between the X chromosome and the disruption of autosomal regulatory pathways in the placenta. To integrate our expression and placental phenotypes more directly, we next tested for QTL that explained expression variation in the overall module eigengene. Despite a small sample size (n = 23), the X chromosome was a significant predictor of the downregulated BC module expression after permutation (Supplementary Figure S13A). In principle, this signal could represent a predominant contribution of X-linked

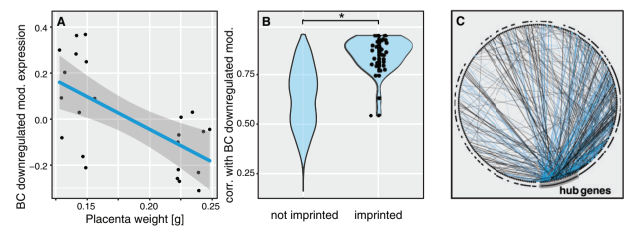


Figure 5. BC downregulated module is associated with placenta size and candidate imprinted genes. (A) Summary expression of the BC downregulated module was significantly associated with placenta size controlling for Theiler stage and sex (P < 0.001, ANOVA, see Supplementary Table S3). (B) This module was enriched for highly connected, candidate imprinted genes ($\mu_{imprinted}=0.8489 \pm 0.015$; $\mu_{non\ imprinted}=0.6370 \pm 0.008$; Welch's two-sided t-test, $t_{49.315} = 12.255$, P < 0.0001; total genes, N = 432; imprinted genes, N = 33). Data points rendered only for imprinted genes. (C) The top 500 pairwise connections within this network largely involved the top 5% most connected genes, with interactions involving hub genes denoted by a thicker line and a solid black circumference notation at the interaction partner. Interactions involving a candidate imprinted gene are indicated in blue.

Gene name	No. of interactions in top 500	Module membership	Imprinting status	Function (UniProtKB)	Clotting/ angiogenesis	Immune	Growth and development	References
Plxdc2	32	0.9259	Candidate	Cell surface signaling	Maybe		Yes	Cheng et al. 2014
Procr	28	0.9451	Candidate	Cell surface signaling, blood coagulation	Yes	Yes	_	Sood et al. 2006; Bouwens et al. 2013
Scara5	23	0.9218	Candidate	Iron transport, blood co- agulation	Yes	—	Yes	Li et al. 2009
Cd68	20	0.9133	Candidate	Immune function	—	Yes	—	Chistiakov et al. 2017
Wnt4	20	0.9451	Candidate	Developmental signal- ing	_	_	Yes	Sonderegger et al. 2010; Knöefler and Pollheimer 2013
Ppard	19	0.9181	_	Transcription factor; steroid hormone re- ceptor	_	_	Yes	Schmidt et al. 1992
Adm	18	0.9515	_	Immune function	Yes	Yes	_	Li et al. 2013
Вос	18	0.9217	_	Cell–cell adhesion, tis- sue differentiation	_	—	Yes	Zakaria et al. 2019
Igsf11	17	0.8696	—	Cell–cell adhesion, cell growth, neurogenesis	_	—	Yes	Jang et al. 2016
Erv3	17	0.9463	Candidate	Tissue identity and immunomodulation	—	Yes	Yes	Mangeney et al. 2007
Tbc1d2b	16	0.8934	_	Protein modification and transport	_	—	Yes	Manshouri et al. 2019
Olfml3	15	0.8884	Candidate	Developmental signal- ing	Yes	_	—	Miljkovic- Licina et al. 2012
Lmcd1	13	0.9115	—	Transcription factor	—	—	Yes	Ferreira et al. 2019
Fn3krp	12	0.8801	_	Protein modification/ glycation	Maybe	Maybe	Yes	Karabag et al. 2007
Numbl	11	0.879	_	Cell–cell adhesion, neu- rogenesis	_	—	Yes	Wilson et al. 2007
Ipmk	11	0.8049	—	Lipid metabolism	—	—	Yes	Malabanan and Blind 2016
F13a1	10	0.888	_	Metal binding, blood co- agulation	Yes	—	—	Muszbek et al. 2011
Pdpn	10	0.8987	—	Cell–cell adhesion, de- velopmental signaling	—	Yes	—	Astarita et al. 2012
Larp6	10	0.824	—	Collagen biosynthesis, fibrosis	—	_	Yes	Stefanovic et al. 2019
Gypc	10	0.8474	—	Cell membrane stability, red blood cells	Yes	—	—	Wilder et al. 2009
Fut4	10	0.8666	—	Cell–cell adhesion, im- mune function	Maybe	Yes	—	Wang et al. 2013

Table 1 Hub genes in the downregulated, placenta-associated BC module

The top 5% most connected genes in the downregulated module, with connection defined as the number of times the gene was included in the top 500 strongest pairwise correlations in gene expression between genes in the module. Gene imprinting status (Brekke *et al.* 2016) and functions are indicated.

genes to the eigengene summary of expression within this parent-of-origin module, or a genome-wide *trans*-regulatory signal dependent on the species origin of the X chromosome. Consistent with the latter hypothesis, only ~6% of genes in downregulated BC module were X-linked (25 of 432 genes, binomial exact test P = 0.006, not significant after correction for multiple testing). These genes were significantly under-represented in the correlation network with only 7 of the top 500 pairwise correlations including an X-linked gene (56 pairs expected; binomial exact test $P \ll 0.0001$). It is also possible that the strong X-linked signal could be a correlated side effect of overall placental phenotypes on expression levels caused by X-linked hybrid incompatibilities. However, we found no significant QTL for the six other expression modules correlated with placental weight (Supplementary Figure S13, B and C), thus this signal seems unlikely to be a consequence of a spurious phenotypic correlation.

Integration of the placental transcriptome onto the *Phodopus* genetic map

These results strongly support the hypothesis that X-linked incompatibilities interacting with a hybrid autosomal background are the primary determinant of disrupted autosomal expression observed in both F_1 and BC hybrids, but the architecture of underlying X-autosome incompatibilities remain unresolved. The current genome assembly for dwarf hamsters is highly fragmented and has not been arranged into an ordered physical map

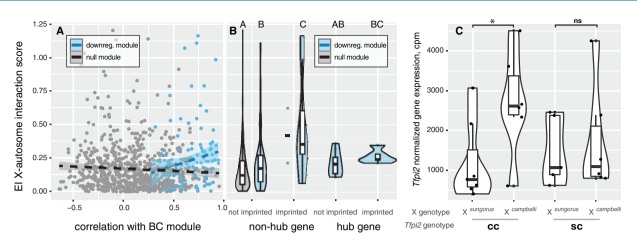


Figure 6. Expression interactions exposed in BC hybrids. (A) Greater disruption in gene expression with X chromosome–autosome mismatch (EI score) was weakly associated with genes more connected in the downregulated BC network module ($F_{1,122} = 4.063$, P = 0.046, ANOVA). Shown are comparisons between the gray module (gray; genes passing filter, N = 733) and the downregulated BC module (blue; genes passing filter, N = 124). The gray module includes all genes that were not placed in any module in the network analysis, and was used to generate a null expectation for the distribution of the EI score. (B) Increased EI score was explained by candidate imprinted genes, rather than highly connected hub genes ($F_{1,122} = 8.75$, P = 0.0037, ANOVA; downregulated BC module not imprinted nonhub genes, N = 102; imprinted nonhub genes, N = 9; not imprinted hub genes, N = 8; imprinted hub genes, N = 5). (C) EI expression pattern is illustrated with top gene Tfpi, where mismatch between maternal X genotype and maternal Tfpi genotype results in an average reduction in gene expression in addition to inheritance of a maternal P. surgorus X chromosome. Gene expression is reported as counts per million (CPM) transcripts. Letters and star indicate significant differences at P < 0.01, assigned by a Tukey's HSD test.

(Bao et al. 2019), limiting our ability to fully integrate our transcriptome and quantitative genetic analyses. To begin to overcome these limitations, we used a custom exon capture experiment to anchor 3,616 placenta-expressed genes onto the *Phodopus* genetic map, including 159 X-linked and 34 autosomal imprinted genes (Supplementary Table S13, Supplementary Figure S14). An additional 212 X-linked genes were identified based on patterns of inheritance and orthology with mouse, but were not ordered on the genetic map (371 X-linked genes total). We placed approximately one-third of the BC downregulated network genes (162 of 432 genes) on to the genetic map, including 17 of the 21 hub genes (Table 1). Genes in this anchored network were distributed across 12 of the 13 autosomes and the X chromosome (Supplementary Figure S15).

A scan for genetic interactions that influence gene expression

If there was a hybrid interaction between the X chromosome and a specific diploid genotype at an autosomal locus that influenced the expression of that gene, then individuals that inherited a maternal P. sungorus X should show a larger change in gene expression for one of the autosomal genotypes than for the other. To test for such interactions, we calculated EI scores by comparing the fold change difference in expression between the two possible autosomal genotypes of all mapped genes (i.e., homozygous for the P. campbelli allele or heterozygous) dependent on the genotype of the maternal X chromosome (P. campbelli or P. sungorus). Mapped genes from the BC downregulated module (n = 124)showed higher EI scores on average when compared to genes not placed in any WGCNA module ($\mu_{downregulated}{=}0.229\pm0.008,$ $\mu_{\rm null}{=}0.165\pm0.006,\;F_{1,870}$ = 14.13, P=0.0002, ANOVA). Next, we used a linear model to assess the relationship between gene connectivity and EI score in the downregulated module. Hub genes disproportionately overlap with male sterility loci in hybrid mice (Morgan et al. 2020), suggesting that incompatibilities may be more common among highly connected genes. However, we found support for only a slight increase in EI score for genes that were the most connected to the module (EI \sim module correlation, adjusted $r^2 = 0.0243$, $F_{1,122} = 4.063$, P = 0.046; Figure 6A). Interestingly, much of the signal for increased EI scores appeared to be driven by candidate imprinted genes rather than hub gene status per se (EI ~ candidate imprinting status, adjusted $r^2 =$ 0.059, $F_{1,122} = 8.75$, P = 0.0037; EI ~ hub, $F_{1,122} = 0.005$, P = 0.942; Figure 6B and Supplementary Figure S16), again underscoring the central role of genes with maternally biased expression within the downregulated module. One such candidate imprinted (nonhub) gene was *Tfpi2* (Figure 6C). Individuals with an interspecific mismatch between the maternally inherited X chromosome (*P. sungorus*) and maternal allele at *Tfpi2* (*P. campbelli*) showed more than a twofold (2.26×) decrease in expression compared to individuals with matching maternal genotypes.

We next polarized EI scores to evaluate specific X-autosome combinations with the downregulated module for candidate hybrid incompatibilities. Within the BC architecture, hybrid incompatibilities involving autosomal recessive or imprinted genes could manifest when the maternally inherited P. sungorus X chromosome was combined with maternally inherited P. campbelli autosomal alleles (all paternal alleles were P. campbelli). Consistent with this prediction, we observed positive polarized EI scores for imprinted genes within the downregulated module demonstrating that these EIs were driven largely by mismatches between the maternal autosomal and X chromosome genotypes (P = 0.043, Tukey's HSD test, nonhub genes, imprinted vs. null module; Supplementary Figure S17A). Furthermore, we found that maternally mismatched imprinted genes in the downregulated module showed larger fold changes when compared to nonimprinted genes regardless of their status as a hub gene (P=0.045, hub imprinted vs. nonimprinted; P < 0.0001, nonhub imprinted vs. nonimprinted, Tukey's HSD test; Supplementary Figure S17B). Collectively, these patterns suggest that imprinted autosomal genes with biased maternal expression within the downregulated module are more likely to be involved in incompatible hybrid interactions with the maternal X chromosome.

Finally, we tested for parent-of-origin bias in expression for candidate imprinted genes in the BC. We found 31 genes previously identified as showing either maternal or paternal bias in expression in F1 placentas (Brekke *et al.* 2016) with expression data for at least seven heterozygous individuals in the BC. Of these, 22 genes show significant parent-of-origin bias in expression in the direction shown previously (Supplementary Figure S18, one-sample t-test, Bonferroni corrected P < 0.05). Although limited to a subset of BC individuals, these results suggest that imprinting status was at least partially maintained at many genes despite large changes in overall expression levels. We also confirmed a strong bias in expression of the maternal allele on the X chromosome (adjusted $r^2 = 0.17$, $F_{1,3560} = 930.3$, $P \ll 0.0001$; Supplementary Figure S19), indicating that paternal imprinted XCI is likely intact in BC females as has been previously reported for F_1 females (Brekke *et al.* 2016).

Discussion

By combining quantitative genetic mapping of placental overgrowth with transcriptomic data, we uncovered genome-wide networks of gene expression that were disrupted as a consequence of incompatible genetic interactions with the X chromosome. These data indicate that genetic interactions between the X chromosome and networks of imprinted and nonimprinted autosomal genes are critical for proper placental development in dwarf hamsters. Below we compare our results with findings from other systems to examine the contribution of the X chromosome to the accumulation of reproductive barriers between species and the evolution of placental gene expression.

The evolution of hybrid placental overgrowth

A positive parent-of-origin correlation between hybrid placental and embryonic growth effects has been observed in some mammal hybrids (e.g., Vrana et al. 2000; Brown et al. 2012), providing a possible mechanistic link between the disruption of early development and extreme adult parent-of-origin growth effects observed in many mammal hybrids (Brekke and Good 2014). Our quantitative results underscore that the relationship between placental and embryonic growth may be weak early in development, especially given the difficulty of differentiating the effects of edema and other developmental defects from overall embryonic growth (Supplementary Figures S6 and S7). Consistent with this, parent-of-origin placental growth effects described in some mouse (Zechner et al. 1996; Kurz et al. 1999) and equine hybrids (Allen et al. 1993) also were not strongly correlated with hybrid embryonic growth.

Regardless of the unclear functional relationship between gross placental and embryonic growth phenotypes, our results show that the species origin of the maternal X chromosome is the major genetic factor responsible for placental overgrowth in Phodopus hybrids. Reproductive barriers between species often evolve through negative epistatic interactions between two or more loci, generally referred to Dobzhansky-Muller incompatibilities (DMIs; Dobzhansky 1937; Muller 1942; Orr and Turelli 2001, see also Bateson 1909). DMIs disproportionately involve X-linked loci, a phenomenon known as the large X-effect (Coyne and Orr 1989b). Hybrid incompatibilities presumably occur between the maternally inherited P. sungorus X chromosome and P. campbelli autosomal loci (e.g., dominant, recessive, or imprinted incompatibilities). However, paternal loci were invariant in our BC and thus paternal contributions to putative X-autosome mismatches could not be mapped in this experiment. Large-X effects for disrupted placental development are perhaps not surprising given the central role that the X chromosome tends to play in the evolution of reproductive isolation (Coyne and Orr 1989b; Turelli and Orr 2000; Masly and Presgraves 2007; Turelli and Moyle 2007) and the parent-of-origin dependent nature of placental hybrid inviability in dwarf hamsters (Figure 2). However, we had previously associated placental overgrowth with widespread disruption of autosomal gene expression (Brekke *et al.* 2016); a regulatory phenotype that could manifest entirely from hybrid incompatible interactions between imprinted autosomal genes. Our mapping results rule out this possibility.

Large effect X-linked QTL also underlie placental overgrowth in hybrid deer mice (Peromyscus maniculatus × P. polionotus; Vrana et al. 2000) and house mice (Mus spretus × M. musculus; Zechner et al. 1996; Hemberger et al. 1999). In all three rodent systems, imprinted XCI occurs in the placenta and incompatibilities on the maternally inherited X chromosome emerge as a central genetic determinant of placental overgrowth in hybrids. However, broader connections between these developmental phenotypes and the disruption of placental gene expression have remained unclear. Studies in deer mice have linked the X chromosome with disrupted imprinted placental pathways, including a putative interaction with the autosomal maternally imprinted gene Peg3 (Loschiavo et al. 2007). X-linked incompatibilities are also the primary cause of hybrid placental overgrowth growth in some hybrid crosses of house mice (M. spretus \times M. musculus; Zechner et al. 1996; Hemberger et al. 1999), but no direct link between disrupted expression of candidate imprinted genes and placental overgrowth has been established (Shi et al. 2004; Zechner et al. 2004). However, a recent genome-wide study on the same Mus hybrid system showed transgressive autosomal expression in undergrown hybrid placentas (reciprocal F1 crosses were not performed), including disruption of the imprinted Kcnq1 cluster (Arévalo et al. 2021). In these experiments, males showed more severe placental undergrowth and disrupted gene expression, which the authors proposed may involve interactions with the imprinted X chromosome (Arévalo et al. 2021). Artificial insemination experiments between more divergent Mus species pairs (M. musculus × M. caroli) resulted in massively abnormal placenta showing local demethylation and overexpression of an X-linked retroelement (Brown et al. 2012). Similarly, loss of genomic imprinting has been correlated with overgrowth in fetuses derived from assisted reproduction between divergent cattle breeds (Chen et al. 2015, 2016). Most of these works have focused on F₁ crosses or candidate gene approaches, limiting further insights into the role that X-autosome interactions play in the broader disruption regulatory pathways in hybrid placenta. Building on these previous studies, we show that X-linked hybrid incompatibilities underlie the disruption of placental growth, with widespread effects on the misexpression of imprinted autosomal pathways.

Gene expression plays a central role in organismal development and morphological evolution (King and Wilson 1975; Carroll 2008; Sears et al. 2015), but overall importance of regulatory incompatibilities to species formation has remained unclear (Butlin et al. 2012; Guerrero et al. 2016). In mammals, progress has been made in linking disruption of specific epigenetic regulatory mechanisms on the X chromosome (e.g., meiotic sex chromosome inactivation) to the evolution of hybrid male sterility during the relatively early stages of speciation (Bhattacharyya et al. 2013; Campbell et al. 2013; Davis et al. 2015; Larson et al. 2017). However, it has remained unclear if other developmental pathways are also predisposed to disruption in animal hybrids (Coyne and Orr 2004; Brekke and Good 2014). We suggest that placental development may represent a second developmental hotspot for the evolution of postzygotic reproductive isolation through the widespread disruption of gene expression.

There are several interesting parallels between the evolution hybrid male sterility and abnormal hybrid placental development in mammals. The X chromosome appears to play a central role in the genetic basis of both hybrid male sterility (Storchová et al. 2004; Good et al. 2008; Davis et al. 2015) and placental overgrowth (Figure 2; Zechner et al. 1996; Hemberger et al. 1999; Vrana et al. 2000). Both forms of postzygotic isolation manifest during developmental stages where the X chromosome is subject to chromosome-wide epigenetic silencing. Whereas meiotic sex chromosome inactivation may often be disrupted in the case of hybrid male sterility (Lifschytz and Lindsley 1972; Bhattacharyya et al. 2013; Larson et al. 2017), imprinted XCI appears to be maintained in hybrid hamster placentas (Supplementary Figure S19; Brekke et al. 2016). Although the specific mechanisms underlying abnormal placental development are yet to be determined, tissues with imprinted XCI may select for the evolution of coadapted epistatic networks of gene expression that are subsequently prone to disruption in hybrids (see below). Finally, aspects of placental development and spermatogenesis are both thought to evolve rapidly in response to genomic conflict (Haig 2000; Larson et al. 2018), which may contribute to the rapid evolution of hybrid incompatibilities (Crespi and Nosil 2013). It is wellestablished that hybrid male sterility tends to evolve rapidly in animals (Coyne and Orr 1989a; Wu et al. 1996), while hybrid growth effects have been detected across a broad range of evolutionary divergence in mammals (Brekke and Good 2014). All three systems where X-linked placental incompatibilities have been found (Phodopus, Mus, Peromyscus) involve crosses between relatively closely related species pairs (Brekke and Good 2014), suggesting that placental hybrid inviability can evolve rapidly and contribute to the early stages of reproductive isolation.

Through analysis of these parallel systems of hybrid sterility and inviability, a trend is emerging where sex chromosome evolution and genetic conflict within regulatory systems appears to fuel divergence within these key developmental processes (Crespi and Nosil 2013; Larson *et al.* 2018), ultimately leading to the formation of reproductive barriers between species. Striking parallels also exist in plants, where hybrid seed inviability also evolves rapidly (Garner *et al.* 2016; Coughlan and Matute 2020) and has been linked to the intensity of parental conflict (Coughlan *et al.* 2020) and the disruption of imprinted gene expression in the extra-embryonic endosperm (Wolff *et al.* 2015).

The evolution of placental gene expression networks

In both rodents and humans, fetal-derived trophoblast cells shape the vasculature at the maternal-fetal interface, allowing for nutrient transport and immune modulation (Gris et al. 2019). Notably, nine of the most connected genes in the BC downregulated module had functions related to coagulation and/or angiogenesis. Endothelial protein C receptor (ProcR) is an important anti-coagulant receptor in the trophoblast coagulation cascade (Bouwens et al. 2013). Allelic variants that result in under-expression of ProcR are associated with fetal loss in humans (Cochery-Nouvellon et al. 2009), and there is some evidence that maternal and fetal ProcR genotypes can interact to either prevent or induce placenta-mediated adverse pregnancy outcomes (Sood et al. 2006). In a healthy rodent placenta, the coagulation initiating tissue factor is counterbalanced by anti-coagulation proteins produced in differentiated syncytiotrophoblast tissue (Sood et al. 2006). Development fails without the early expression of the coagulation cascade in the placenta (Isermann *et al.* 2003). However, low levels of anticoagulants later in development are associated with preeclampsia and pregnancy loss (Ebina *et al.* 2015).

Several of the hub genes (Table 1) are known to contribute to differentiation of placental layers. For example, Wnt signaling is broadly important in placentation and embryonic development, and Wnt4 specifically may be involved in signaling between the fetal and maternal placental layers (Sonderegger et al. 2010; Knöefler and Pollheimer 2013). Another such specialized hub gene, Erv3, is part of a family of genes co-opted from endogenous retroviruses and are involved in immunomodulation, fusion, and differentiation of trophoblasts (Mangeney et al. 2007), and are increasingly recognized for their role in regulating placental gene expression (Pavlicev et al. 2015; Chuong 2018). Similarly, the Plexin domain containing 2 gene (Plxdc2) encodes an endothelial cell-surface transmembrane receptor (Cheng et al. 2014) that is often co-expressed with Wnt signaling genes (Miller et al. 2007). Other candidate hub genes play roles in cell-cell adhesion and differentiation (Wilson et al. 2007; Jang et al. 2016; Zakaria et al. 2019), immune function (Astarita et al. 2012; Li et al. 2013; Wang et al. 2013; Chistiakov et al. 2017), nutrient metabolism and delivery (Schmidt et al. 1992; Li et al. 2009; Malabanan and Blind 2016), and transcriptional regulation (Ferreira et al. 2019).

Overall, our expression data suggest a strong connection between placental overgrowth, the maternally expressed (P. sungorus) X chromosome, and the imprinted expression of autosomal genes. Our candidate imprinted gene set included several genes known to be maternally (e.g., Igf2, Mest, and Peg3) or paternally imprinted (e.g., Axl, H19, Tfpi2, and Wt1) in mice, as well as several novel candidates including most of the hub genes (Table 1). Confirmation that these candidates reflect the evolution of novel parent-of-origin epigenetic silencing in Phodopus (e.g., through DNA methylation or other mechanisms) awaits detailed functional validation beyond the scope of the current study. Others have argued that contamination of maternal blood or tissue may often bias patterns of allele-specific expression in the post-embryonic placenta (Wang et al. 2011, but see Finn et al. 2014). We previously found little evidence for extensive maternal contamination of dissected placental tissue in Phodopus (i.e., genome-wide paternal: maternal allele ratios were \sim 1:1 in overgrown S \times C placentas), but it is possible that maternally biased expression of some of these candidates reflects a large maternal contribution to overall placental expression levels. Indeed, hub genes such as Wnt4 are thought to be directly involved in signaling between the fetal and maternal placental layers (Sonderegger et al. 2010; Knöefler and Pollheimer 2013). Regardless of the underlying regulatory mechanisms-epigenomic imprinting or fetal-maternal transcript sharing—our results suggest that X-linked and autosomal genes with maternally biased expression play a central role in the evolution of placental development and the disruption of placental pathways in hybrids.

The existence of placental networks of maternally biased gene expression is consistent with some predictions of the co-adaptation theory of gene expression, whereby maternal expression at one gene can select for maternally biased expression at other positively interacting genes (Wolf and Hager 2006; Wolf 2013; Wolf and Brandvain 2014; O'Brien and Wolf 2017). Such a co-evolutionary process should result in the broader integration of imprinted gene networks, the evolution of separate co-expressed networks of maternally and paternally expressed genes, and the exposure of epistatic DMIs in second- (or later) generation hybrids with recombinant genotypes (Wolf and Brandvain 2014; Patten et al. 2016). We uncovered evidence for many such interactions in our preliminary screen for genetic interactions that influence gene expression levels in BC placenta (Figure 6; Supplementary Figures S16 and S17). For example, the maternally expressed *Tfpi2* gene showed a greater than twofold decrease in expression when a maternally inherited *P. campbelli* allele was combined with a maternally inherited *P. sungorus* X chromosome (Figure 6C). *Tfpi2* is imprinted in the placenta where its expression may limit trophoblast invasion (Jin et al. 2001) and also downregulated in several types of cancer (i.e., a potential tumor suppressor; Konduri et al. 2001; Ribarska et al. 2010). More data are needed to determine if this and other genes showing significant EIs contribute directly to abnormal placental growth phenotypes in hybrids.

More generally, we propose that the X chromosome is likely to play a central role in the evolution of maternally biased placental networks in many eutherian mammals. The strength of this prediction is dependent on patterns of X inactivation in the placenta and other extra-embryonic tissues of females as males only have a maternally derived X. The paternal X chromosome appears to be silenced in the placenta and other extra-embryonic tissues in at least four genera of rodents (i.e., Mus, Takagi and Sasaki 1975; Rattus, Wake et al. 1976; Phodopus, Brekke et al. 2016; Peromyscus, Vrana et al. 2000), resulting in predominantly maternal expression of X-linked genes in the placenta of males and females (Dupont and Gribnau 2013; Lee and Bartolomei 2013). Imprinted XCI is expected to contribute the vast majority of maternally expressed genes in the rodent placenta, which as a consequence should favor the evolution of maternal expression at interacting autosomal genes. In contrast, XCI appears to be random in extraembryonic tissues of humans (Moreira De Mello et al. 2010), cattle (Chen et al. 2016), pigs (Zou et al. 2019), horses (Wang et al. 2012), and possibly rabbits (Okamoto et al. 2011). The more frequent occurrence of random XCI in this limited sample suggests that imprinted XCI may have evolved more recently in rodent extraembryonic tissues (Okamoto et al. 2011), and therefore may not apply broadly across the radiation of placental mammals. However, it is worth noting that rodents comprise over 40% of all placental mammal species (Burgin et al. 2018). The current sample of XCI in extraembryonic tissues is also biased toward a just few major lineages (i.e., ungulates, primates, rodents, and lagomorphs) and thus insufficient for accurate reconstruction of the ancestral state of extra-embryonic XCI in placental mammals. Moreover, male hemizygosity in species with random XCI may still favor the evolution of maternal expression at interacting autosomal genes under some conditions (Wolf and Brandvain 2014). In addition, the physiological integration of maternal blood supply and trophoblast-generated fetal vasculature is a particularly compelling biological context that could favor the evolution of coordinated maternal-fetal gene expression networks, regardless of the pattern of XCI. Given our data and these general theoretical predictions, the broader relevance of X-autosomal gene expression networks to placental evolution and development warrants further consideration.

Data Availability

All raw sequencing reads are archived at NCBI under BioProject PRJNA306772. Accession numbers for individual libraries are provided in Supplementary Table S5. Supplementary material is available at figshare: https://doi.org/10.25386/genetics.14171075.

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Conflicts of interest

The authors declare no competing financial interests.

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