Interaction between the X chromosome and an autosome regulates size sexual dimorphism in Portuguese Water Dogs

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Size sexual dimorphism occurs in almost all mammals. In Portuguese Water Dogs, much of the difference in skeletal size between females and males is due to the interaction between a Quantitative Trait Locus (QTL) on the X-chromosome and a QTL linked to Insulin-like Growth Factor 1 (IGF-I) on the CFA 15 autosome. In females, the haplotype of CFA 15 resulting in small size is dominant. In males, the haplotype for large size is dominant. Females, homozygous at the CHM marker on the X chromosome and homozygous for the large size CFA 15 haplotype are, on average, as large males. However, all females that are heterozygous at the CHM marker are small, regardless of their CFA 15 genotype. This interaction suggests a genetic mechanism that in turn leads to a scenario for the evolution of size sexual dimorphism consistent with a proposal of Lande that sexual dimorphism can evolve because females secondarily become smaller than males as a consequence of natural selection for optimal size. Our results also can explain Rensch's Rule, which states that size is often positively correlated with the level of size sexual dimorphism.

Since Darwin (1875), sexual selection for dimorphism in size has been a subject of intense interest to biologists. A difference in size between the sexes (sexual dimorphism) is observed in almost all mammals (Abouheif and Fairbairn 1997), including dogs (Canis familiaris). The mechanisms for maintaining sexual dimorphism in a population are not completely understood, but may involve interaction between the sex chromosomes and autosomes. For example, the Sry locus on the Y chromosome plays an important role in sex determination and dimorphism (McLaren 1990). The X chromosome has been shown to harbor an excess of genes related to sex and reproduction, mental functions, and skeletal muscle (Bortoluzzi et al. 1998; Saifi and Chandra 1999; Wang et al. 2001; Zechner et al. 2001; Qiu et al. 2002; Vallender and Lahn 2004). Several autosomal loci are expressed differentially in males and females with concomitant effects on sexual dimorphism (Udy et al. 1997; Vaughn et al. 1999; McMahon et al. 2003; Salih et al. 2005).

We are analyzing the genetic basis for skeletal morphology in Portuguese Water Dogs (Chase et al. 1999, 2002, 2004). Portuguese Water Dogs (PW dogs) are a recent breed of dog, whose descent from a few founders is documented by accurate and complete pedigree records (Molinari 1993). They are descended from two major founding kennels that disagreed on an appropriate size standard, resulting in considerable size variation in the founding population. The population structure of the breed is favorable for detailed genetic analysis of quantitative phenotypes (Chase et al. 1999) and variation in size and shape is still encountered within the current population. This is, in part, due to a fairly lax size standard for the breed, although a different average standard is maintained for males and females. Moreover, any

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pure-bred PW dog can compete in "performance events" (obedience, agility, waterwork, etc.) regardless of size.

Insulin-like Growth Factor 1 (IGF-1) regulates postnatal skeletal growth in mammals (Baker et al. 1993). Here we present evidence that an autosomal genetic marker, linked to *IGF1*, is associated with size sexual dimorphism in PW dogs. This locus interacts with another locus on the X chromosome to regulate differences in size between the female and male populations.

Results

Female PW dogs are, on average, 85% the size of males. Based on 42 metrics from the pelvis, fore- and hind-limbs, we have estimated size as the first Principal Component (PC1) of skeletal variation (Chase et al. 2002). (PC1 is significantly correlated with body mass (r = 0.7) and represents an averaging over each dog's normalized skeletal metrics. It is less influenced by environmental factors such as diet and exercise.) This trait is highly heritable ($h^2 = 0.45$) and we have associated PC1 variation in PW dogs with quantitative trait loci (QTLs) on four autosomes and the X chromosome (approximate locations on the dog genome are presented in Fig. 3 of Carrier et al. [2005]).

Figure 1 presents size distributions of PC1 for male and female PW dogs. Sexual dimorphism is evident as a shift in population means, but the ranges of size overlap extensively and several females are within the size range of the largest 10% of the male population.

A QTL regulating size variation was associated ($P \le 0.028$) with tetra nucleotide repeat marker *FH2017* on autosome 15 (CFA 15; see Carrier et al. 2005). Two major *FH2017* alleles (*A* and *B*) account for >90% of *FH2017* genotypes. On the canine genetic map (Guyon et al. 2003), this marker is linked to canine *IGF-1* (Chase et al. 2002). The variation in PC1 associated with genotypes of *FH2017* is shown in Figure 2. The data are consistent with an additive mode of inheritance in which genotypic means

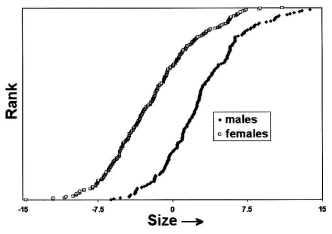


Figure 1. Size sexual dimorphism in the PW Dog. Forty two skeletal metrics were corrected to a mean of 0 and a standard deviation of 1. A matrix composed of the corrected skeletal metrics of 463 dogs was used to calculate the Principal Components of skeletal variation. Size is represented by values of PC1 calculated using the Pearson correlation option of XLSTAT (http://www.xlstat.com) for 42 skeletal metrics (males ♦; females □).

are ranked: BB(small) < AB < AA(large). This interpretation changes, however, when the population is further subdivided according to sex (Fig. 3). In males, haplotype *A* (large-size phenotype) is dominant (AA = AB > BB), whereas Haplotype *B* (small size phenotype) is dominant in females (BB = AB < AA). This difference provides a genetic basis that explains part of the sexual size dimorphism observed between the male and female populations.

We deconstructed PC1 by comparing the effects of PC1 QTLs on individual bone metrics. With one exception, no significant correlation could be established between the individual QTLs. However, variation associated with the CFA 15 locus (FH2017) was significantly correlated with trait variation explained by the X chromosome PC1 QTL. This suggested the existence of an interaction between CFA 15 and X chromosome PC1 OTLs. This was confirmed when we searched for an interaction between loci on the X chromosome and the FH2017 autosomal locus: A QTL on the X chromosome, associated with the SSR marker CHM, interacts significantly ($P \le 0.001$) with the FH2017 QTL. CHM is located in a region of the X chromosome that has been shown in humans to escape X inactivation (Carrel and Willard 1993, 2005). Similar to FH2017, CHM has only two haplotypes, α (51%) and β (49%) identified by two alleles of an SSR marker in an intron of this gene. In females homozygous at CHM (e.g. $\alpha\alpha$ or $\beta\beta$), the size of FH2017 AA genotypes approach values found in FH2017 AA males (top, Fig. 4). In contrast, CHM heterozygous females (e.g., $\alpha\beta$) are small and FH2017 genotypes show no segregation for size (Fig. 4, bottom). Hemizygous males, α and β , each do not differ significantly from the pattern of FH2017 size segregation observed for males in Figure 3. This interactive effect, confined to the CHM region of the X, reduces the average size of female PW dogs, but maintains a small fraction of very large females in the population (alternatively, the interaction increases the size of a small portion of female dogs).

The effects of dominance reversal and the interaction between *CHM* and *FH2017* are significant for each of the individual traits contributing to variation in PC1, but we lack sufficient statistical power to determine whether some traits are affected more than others. The overall effect of dominance reversal and the interaction between *CHM* and *FH2017* is responsible for \sim 50% of the size sexual dimorphism observed in the breed.

Finally, we have analyzed parental PW dog phenotypes and genotypes, but failed to find evidence for imprinting. There was no observed effect of *FH2017* on size in the female *CHM*- $\alpha\beta$ population regardless of which *CHM* allele was inherited from the sire. Similarly, there is no significant parent of origin effect for *FH2017* alleles (e.g., *FH2017-AB* females who received the *A* allele from the sire are no different from those that received the *A* allele from the dam).

Discussion

These data present two sex-related phenomena demanding explanation (as shown in Fig. 5), the reversal of dominance at the *FH2017* QTL, and the heterozygote-specific interaction between the *CHM* locus and the *FH2017* QTL. The first phenomenon requires the existence of another sex-specific factor. For example, the *FH2017* QTL might contain two distinct genes associated with two haplotypes; *FH2017* haplotype *A* acts in both males and females to up-regulate size, while *FH2017* haplotype *B* not only does not up-regulate size, but contains another gene that suppresses this up-regulator. It is this suppressor that is itself suppressed by an unknown factor in males, perhaps derived from the Y chromosome.

To explain the second phenomenon, we propose that activation of the haplotype *A* up-regulator requires interaction with a *CHM*-associated multimeric product, either $\alpha\alpha$ or $\beta\beta$. The failure of such activation by *CHM* heterozygotes would occur if a mixed multimeric product ($\alpha\beta$) were inactive (spoiler effect) (Kadouri et al. 1978). Males, being hemizygous, are never heterozygotes and cannot produce the proposed mixed multimer. The difference between heterozygotes and homozygotes at the *CHM* locus is consistent with the observation that this region of the X chromosome escapes inactivation (both copies of the X are active) (Carrel and Willard 1993, 2005), a phenomenon that has been observed at other X-linked loci (Brown and Greally 2003).

The hypothesis in Figure 5 suggests a two-step scenario for

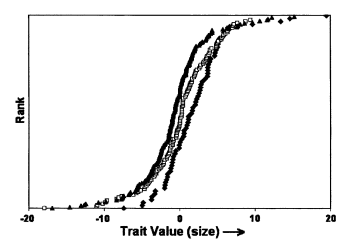


Figure 2. PC1 variation subdivided according to *FH2017* genotypes $(AA \Leftrightarrow; AB \sqsupset; BB \blacktriangle)$. Dogs are ranked on the *y*-axis according to pedigree-corrected values of PC1 on the *x*-axis (Chase et al. 2002, 2004). Pedigree effects were removed using BLUP (Lynch and Walsh 1998; Chase et al. 2004).

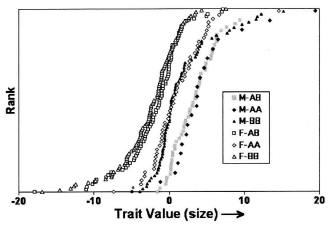


Figure 3. Size variation (values of PC1) subdivided according to sex and *FH2017* genotypes. The *FH2017* genotype data in Figure 2 were further subdivided into the male and female subpopulations.

the evolution of sexual dimorphism. (1) Male competition results in selection for larger males. This sexual selection acts on many body-size genes on the X and autosomal chromosomes. For example, a dominant haplotype A might arise at the FH2017 locus, driving both males and females to become larger. (2) Natural selection then acts to favor a smaller, optimal body size in females. Modifications such as the reversal of dominance and the interaction with the X chromosome create smaller females and sexual dimorphism. Interestingly, this system maintains sexual size dimorphism only when the alleles involved segregate nearly evenly in the population, as we indeed observe. Our hypothetical evolutionary scenario thus supports Lande's proposal (Lande 1980) that sexual dimorphism evolves when females secondarily return to their natural selection optimum after having undergone correlated evolution due to sexual selection acting on males.

The interaction between the X-chromosome and CFA 15 also maintains size polymorphism in females. Long-term changes in food availability and the strength of competition could favor maintenance of a body-size polymorphism in long-lived animals. Thus, large individuals would be most effective in fighting with neighboring social groups and capturing large prey, while smaller individuals can use their higher agility to capture small prey (Peters 1983). Such factors are known to be important for ants (Holldobler and Wilson 1990) and could operate in other social systems including wolves (*Canis lupus*) from which *Canis familiaris* is descended (Vila et al. 1997).

If such a genetic system works across species, those species with a major gene (such as that linked to *FH2017*) of large effect will simultaneously show larger average size and a larger degree of sexual size dimorphism caused by repression of that effect in females. Our data are thus consistent with Rensch's Rule that the level of sexual size dimorphism is often positively correlated with body size (Rensch 1950, 1960; Abouheif and Fairbairn 1997) and female body size in particular. Species in which this correlation holds include primates (Abouheif and Fairbairn 1997; Leutenegger 1978), other mammals (Jarman 1983, 1989; Reiss 1986; Abouheif and Fairbairn 1997), birds (Webster 1992; Abouheif and Fairbairn 1997).

Many hypotheses have been proposed to explain this rule (for review, see Webster 1992; Abouheif and Fairbairn 1997). The only hypotheses that can be generalized across taxa attributes the pattern to sexual selection on males (Fairbairn and Preziosi 1994). In one scenario, sexual selection favoring large males produces an increase in male size and a smaller, correlated increase in female size because of the high genetic correlation between the sexes (Leutenegger 1978; Lande 1980; Fairbairn and Preziosi 1994; Aboufeif and Fairbairn 1997).

As we have argued, our data are more consistent with Lande's prediction (1980) that sexual dimorphism evolves because females secondarily become smaller than males as a result of natural selection for optimal size. Reduction of female size relative to that of males through an inhibition of genes that enhance growth, such as that associated with *FH2017* on CFA 15, may commonly be related to the evolution of sexual size dimorphism. Such a pattern of inverse dominance, in which small is dominant in females, whereas large is dominant in males, should be relatively easily tested in other species where controlled mating is feasible and hybrid progeny can be examined.

Our data indicate an interactive X chromosome role in the regulation of size sexual dimorphism. We have not identified the specific genes involved in this interaction. While some candidates are suggested by proximity to QTLs (e.g., *IGF-1*), the actual identification of the phenotype-associated variants will be a challenging process. In order to truly establish cause and effect, a large number of dogs representing a range of well-characterized phenotypes must be sequenced in regions of interest and rigorous statistical methods applied to the resulting data. Given that we are not searching for mutations causing a profound disease, rather a phenotype associated with some variability, it is likely

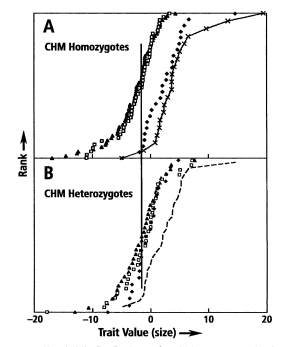


Figure 4. Size (PC1) distributions of *FH2017* genotypes in *CHM* females: (*A*) homozygous ($\alpha\alpha$ and $\beta\beta$); (*B*) heterozygous ($\alpha\beta$). The data represent genotypic subpopulations of the data in Figure 2. As in Figure 2, dogs are ranked by pedigree-corrected PC1 values on the *x*-axis. *FH2017* genotypes are shown ($AA \Leftrightarrow$; $AB\Box$; $BB \blacktriangle$). For comparison, male *FH2017 AA* genotypes (-x-) also are presented. The 50% rankings of *BB* genotypes (\blacktriangle) are almost the same (vertical line between *A* and *B*). For purposes of comparison, we have added a dashed line to *B*, reproducing the distribution of *AA* females from *A*.

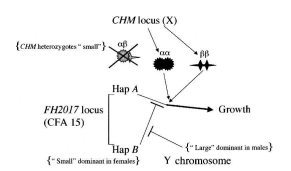


Figure 5. An interactive network between the X (*CHM*), Y, and CFA 15 chromosomes suggested by the data in Figures 1–4.

that DNA variants involved in gene expression levels, exonskipping message stability, or protein folding will prove to be important. Such variants, indeed, are likely to be the syntax associated with genes like *IGF-1* and others that account for the continuum of phenotypic variation observed in dogs. However, once the causative variants are truly identified, it will be possible to search for this mechanism in other canids, such as the wolf, as well as other mammals including humans, to test whether variants and/or molecular mechanisms are preserved by selection (Garrigan and Hedrick 2003).

Methods

We have increased the number of dogs in the data set previously used for analysis of the full skeleton and focused on metrics of the hip and limb (Fig. 3B,D,E in Chase et al. 2002). We have obtained phenotypes (skeletal metrics from radiographs) and DNA genotypes (alleles of simple sequence repeat (SSR) markers) from 463 PW dogs. Radiographs for skeletal metrics and blood for DNA were obtained from owners of individual dogs. These radiographs, together with the measurements taken from them, have been described in a previous publication (Chase et al. 2002).

Methods for identifying QTLs and for estimating the genotypic means of markers near a QTL in an unstructured population have been published (Lynch and Walsh 1998; Chase et al. 2002, 2004). Briefly, QTLs are identified using an allele-sharing method. This method compares pedigree-corrected allele-sharing values at a marker with phenotypic similarity to establish an association between marker and phenotype. Monte Carlo simulations of a random trait with heritability equal to the target phenotype are used to estimate the NULL distribution of the marker-phenotype association. This NULL distribution is used to estimate the significance of the association between the marker and the target phenotype. Marker P-values are adjusted for the ca. 700 markers tested. Marker genotypic means are estimated using a mixed model in which the marker genotypes are treated as fixed effects and the additive genetic background and error deviations are treated as random effects. Infrequent genotypes are bulked for this analysis.

Interaction between *FH2017* and *CHM* was evaluated using females only. The variation explained by *FH2017* in the *CHM* $\alpha\alpha$, $\alpha\beta$, or $\beta\beta$ subpopulations is $R^2_{\alpha\alpha}$, $R^2_{\alpha\beta}$, and $R^2_{\beta\beta}$, respectively. The Variance (VR) among these R^2 values was used as a test statistic for the interaction between the two loci. We used permutation tests to establish the significance of this interaction. For each permutation test, the genotypes for *CHM* were permuted with respect to the individuals and VR recalculated. Each permutation test created a random assortment of *CHM* genotypes, but maintained the relationship for the *FH2017* genotypes, allowing us to

fit the animal for *FH2017* in each sub group. We used 5000 permutation tests to establish the null distribution for the VR statistic.

We tested for imprinting using parent of origin tests. Heterozygotes at each marker were subdivided according to the parental origin of a specific allele (A or α) and tested for significant differences in QTL effects. We tested for differences between the *CHM*- $\alpha\beta$ (α from sire) and *CHM*- $\alpha\beta$ (α from dam) as well as the *FH2017-AB* (A from sire) and *FH2017-AB* (A from dam).

We calculated the amount of size sexual dimorphism explained by these loci and the interaction between them as $1-R_a^2/R_b^2$; where R_b^2 is the amount of variation explained by sex in the raw data and R_a^2 is the amount of variation explained by sex in the residuals after accounting for the two loci and the interaction. Treating males as homozygous at the X locus size was modeled as: size = $\mu + A_i + A_{md} + A_{fd} + X_{het} + e$ where μ is the mean, A_i is the allele count for *FH2017-A*, A_{md} is a male-specific dominance factor (1 for male *FH2017-AB* and 0 for all others), A_{fd} is a factor for heterozygotes at the *CHM* locus.

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