

# Genomic Architecture of Hybrid Male Sterility in a Species Without Sex Chromosomes (*Tigriopus californicus*, Copepoda: Harpacticoida)

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## Abstract

Sterility among hybrids is one of the most prevalent forms of reproductive isolation delineating species boundaries and is expressed disproportionately in heterogametic XY males. While hybrid male sterility (HMS) due to the “large X effect” is a well-recognized mechanism of reproductive isolation, it is less clear how HMS manifests in species that lack heteromorphic sex chromosomes. We evaluated differences in allele frequencies at approximately 460,000 SNPs between fertile and sterile F<sub>2</sub> interpopulation male hybrids to characterize the genomic architecture of HMS in a species without sex chromosomes (*Tigriopus californicus*). We tested associations between HMS and mitochondrial-nuclear and/or nuclear-nuclear signatures of incompatibility. Genomic regions associated with HMS were concentrated on a single chromosome with the same primary 2-Mbp regions identified in one pair of reciprocal crosses. Gene Ontology analysis revealed that annotations associated with spermatogenesis were the most overrepresented within the implicated region, with nine protein-coding genes connected with this process found in the quantitative trait locus of chromosome 2. Our results indicate that a narrow genomic region was associated with the sterility of male hybrids in *T. californicus* and suggest that incompatibilities among select nuclear loci may replace the large X effect when sex chromosomes are absent.

**Key words:** hybrid male sterility, reproductive isolation, pool-seq, *Tigriopus californicus*.

## Significance

Reproductive incompatibility between divergent taxa can often be traced to hybrid male sterility (HMS). For many animals, hybrid male sterility is due to genetic incompatibilities associated with sex chromosomes. Yet not all species have sex chromosomes. In these species, sterility might be due to incompatibilities between mitochondrial and nuclear genes or strictly among nuclear genes themselves. We evaluated the causes of hybrid sterility in a copepod species without sex chromosomes and found that incompatibilities underlying sterility map to a single autosome where genes involved with spermatogenesis are enriched and are likely involved in male reproductive failure. Our results highlight how hybrid male sterility and the boundaries between species may be maintained through population-specific nuclear incompatibilities in taxa without sex chromosomes.

## Introduction

Inviability and reproductive failure in interspecific hybrids can often be attributed to mismatching genetic elements with differing evolutionary histories (Dobzhansky 1936;

Muller 1942; Orr 1995). Incompatibilities among sex and autosomal chromosomes make an especially strong contribution to hybrid reproductive failure and the formation of allopatric species (i.e., the “Large X Effect”; Presgraves

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2008; Phillips and Edmands 2012; Lima 2014). In many interspecific crosses, hybrid offspring that are members of the heterogametic sex are more susceptible to reproductive failure than their homogametic counterparts (Haldane 1922). “Haldane’s Rule” can often be attributed to recessive sex-linked factors that are hidden in homogametic individuals but exposed in heterogametic form (Turelli and Orr 1995; Presgraves and Orr 1998). Sterility is one such hybrid incompatibility that is disproportionately prevalent in males (or females in ZW systems), typically because of genetic elements that accumulate on sex chromosomes (Price and Bouvier 2002; Masly and Presgraves 2007; Presgraves and Meiklejohn 2021). In fruit flies, nematodes, mosquitos, and mice, HMS can be traced to incompatibilities between X-linked and autosomal loci (Tao et al. 2003; Slotman et al. 2004; Schwann et al. 2018; Bi et al. 2019). While the concept of the large X effect has a broad influence, it is not well understood how HMS manifests across animal species that lack sex-determining chromosomes or have a strong environmental contribution to sex determination (Willett 2008).

Aside from sex chromosomes, incompatibilities between nuclear and mitochondrial genomes are at times linked to male sterility in plants and hybrid inviability across taxa (Chase 2007; Burton et al. 2013). In many angiosperms, sterility-causing mutations involve new open reading frames formed from recombining mitochondrial genomes, but these mutations are often counteracted by nuclear “restorer-of-fertility” genes (Chase 2007; Case and Willis 2008). Moreover, mito-nuclear interactions play an important role in the fitness breakdown of interpopulation or interspecific  $F_2$  hybrids in some animal and fungal taxa (Bolnick et al. 2008; Chou et al. 2010; Meiklejohn et al. 2013; Lima et al. 2019; Healy and Burton 2020). Hybrid breakdown due to mito-nuclear interactions in these systems are most often manifested via inviability, slow development, metabolic dysfunction, and smaller clutch sizes. Alternatively, incompatibilities strictly among nuclear autosomal loci contribute to hybrid pollen sterility in some hermaphroditic plants (Sweigart et al. 2006; Long et al. 2008; Feng et al. 2020), but the pervasiveness of this pattern in species with separate male and female sexes is uncertain. Assessing the genomic architecture of HMS in species with separate sexes but without sex chromosomes could provide valuable insight into the relative importance of nuclear and mito-nuclear interactions in the formation of reproductive isolation generally and HMS in particular.

We evaluated the genomic architecture of male sterility resulting from interpopulation hybridization in the tidepool copepod *Tigriopus californicus*. *Tigriopus californicus* has a polygenic sex-determining system and lacks heteromorphic sex chromosomes (Alexander et al. 2014; 2015). In some interpopulation crosses, male sterility in  $F_2$  hybrids is increased relative to its baseline frequency within populations (Willett 2008). In the absence of sex

chromosomes, asymmetries in the manifestation of HMS between reciprocal crosses might implicate mito-nuclear incompatibilities as an important source of sterility. Alternatively, HMS in *T. californicus* could be the result of mismatches among nuclear elements and these effects may be localized or dispersed throughout the genome.

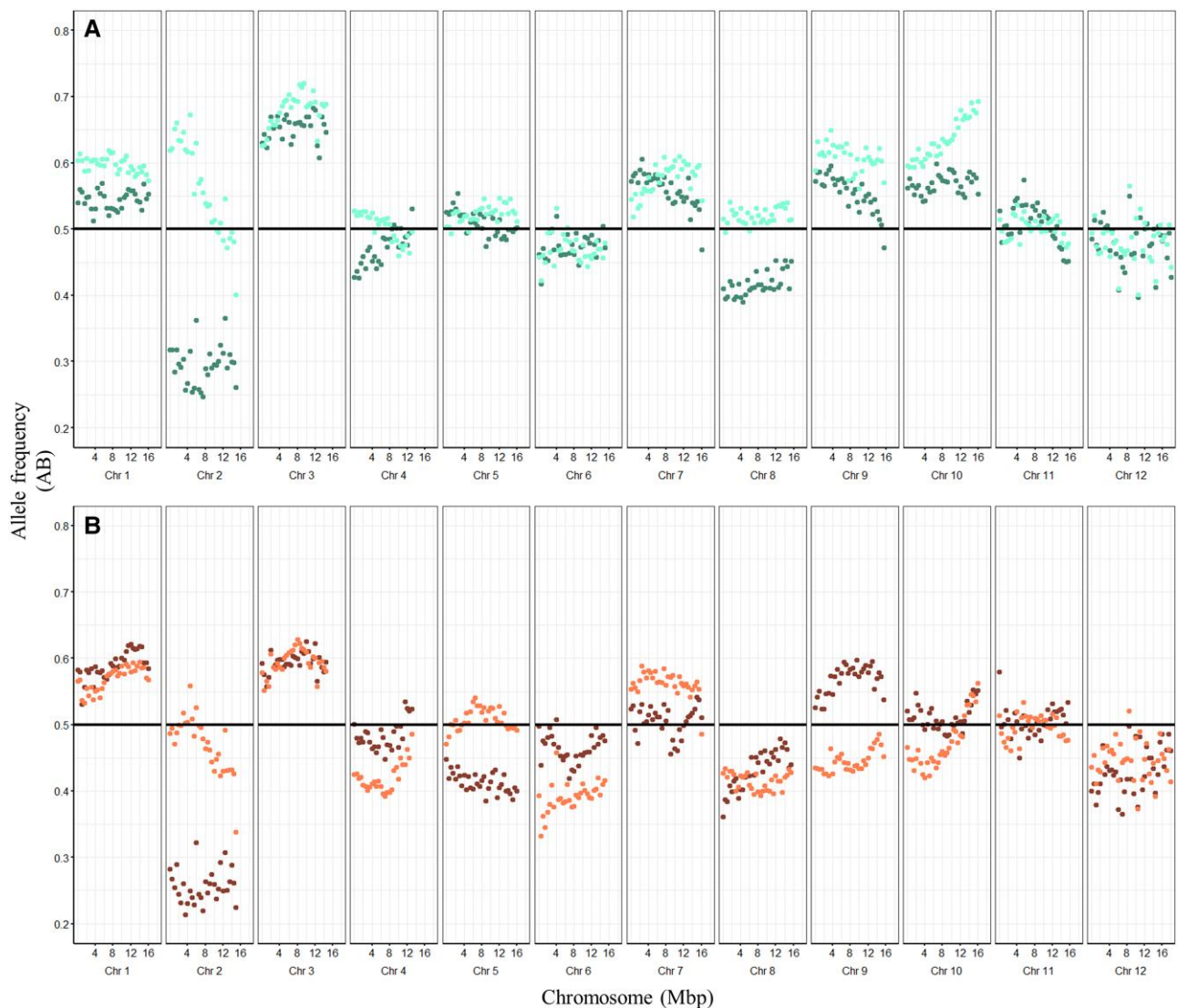
## Results

In reciprocal crosses between copepods from Abalone Cove (AB) and San Diego (SD) CA, USA, we scored fertility and sterility across a total of ~2,000  $F_2$  interpopulation male hybrids, and then performed whole-genome sequencing of four pools (i.e., a pool of fertile males and a pool of sterile males for each cross direction). Allele frequencies in  $AB\text{♀}\times SD\text{♂}$  and  $SD\text{♀}\times AB\text{♂}$  crosses were evaluated at 462,330 and 458,526 population-diagnostic SNPs, respectively (supplementary table S1, Supplementary Material online). The average depth of coverage per locus ranged from  $68.824 \pm 0.100$  (mean  $\pm$  95% C.I.) to  $85.975 \pm 0.159$  across the four hybrid pools. The allele frequencies of each pool are plotted in figure 1.

Regions of the genome associated with HMS were identified by quantifying differences in allele frequencies among fertile and sterile males within each of the reciprocal crosses using a bulk segregant/quantitative trait locus (QTL) approach.  $G'$  analyses detected a single 15-Mbp segment on chromosome 2 that was associated significantly with HMS (fig. 2). Within the region detected in  $G'$  analyses, we identified three 2-Mbp QTL peaks where differences in allele frequencies were greatest: *QTL-1* (0.8–2.4 Mbp), *QTL-2* (3.4–5.7 Mbp), and *QTL-3* (6.2–7.6 Mbp) (fig. 2, supplementary fig. S1, Supplementary Material online). The segment corresponding to *QTL-2* expressed the largest difference in allele frequencies between fertile and sterile male hybrids relative to the rest of the genome, and it overlapped fully between the reciprocal crosses.

A primary goal of this experiment was to assess the potential role of mito-nuclear incompatibilities as a source of HMS. Such incompatibilities would be inferred if sterile males showed patterns of mismatched nuclear allele frequency relative to their respective mitochondrial backgrounds. In other words, sterile males from the  $AB\text{♀}\times SD\text{♂}$  cross would have elevated SD nuclear allele frequencies, while sterile males from the  $SD\text{♀}\times AB\text{♂}$  cross would show elevated AB nuclear allele frequencies. Within the detected QTLs, however, sterile males from both cross directions showed elevated frequencies of SD alleles, clearly rejecting the mito-nuclear incompatibility hypothesis for HMS in hybrids between these populations of *T. californicus* (fig. 1).

We identified 121, 279, and 173 protein-coding genes within *QTL-1*, *QTL-2*, and *QTL-3*, respectively (supplementary table S2, Supplementary Material online). Across the protein-coding genes within the three QTL



**FIG. 1**—Allele frequencies of fertile and sterile *T. californicus* male  $F_2$  hybrids produced from (A)  $AB♀ \times SD♂$  and (B)  $SD♀ \times AB♂$  crosses. Allele frequencies are based on the AB allele perspective. Light-shaded symbols are allele frequencies of fertile males; dark-shaded symbols are allele frequencies of sterile males. Data points represent mean allele frequency across SNPs in nonoverlapping 500-kb sliding windows. Total assembly lengths vary across the 12 *T. californicus* chromosomes (13–18 Mbp).

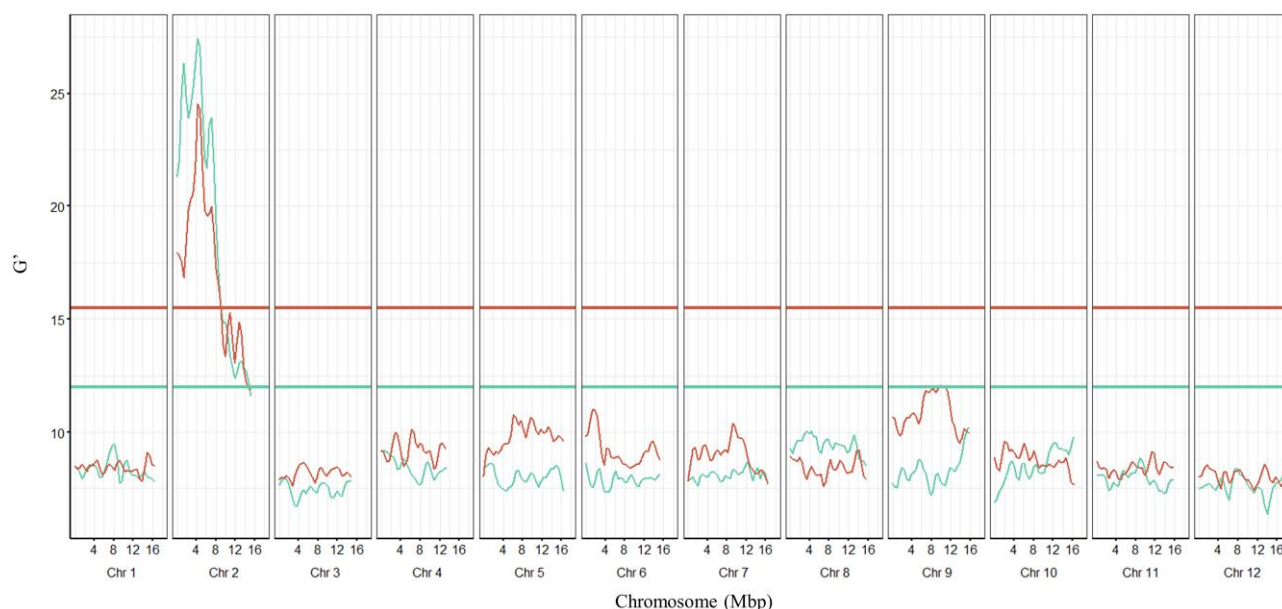
regions, a total of 55 Gene Ontology (GO) terms were over-represented relative to their frequencies in the whole genome (table 1; supplementary table S3, Supplementary Material online). Interestingly, the most overrepresented term was “spermatogenesis”, with nine genes in *QTL-2*, which is  $\sim 4.1 \times$  more common than expected by chance when considering that 107 total genes are annotated with this GO term.

## Discussion

We found a 2-Mbp genomic region closely associated with the sterility of *T. californicus*  $F_2$  interpopulation male hybrids. The boundaries of the region were remarkably

consistent in reciprocal crosses and, unlike other forms of hybrid breakdown in *T. californicus*, displayed patterns indicative of nuclear rather than mito-nuclear incompatibility. Our results suggest mismatches among select nuclear loci were responsible for the sterility of male hybrids and point to the role of autosomal incompatibilities in forming HMS when sex chromosomes are not an available target of intrinsic reproductive isolation.

Within the identified QTL, sterile  $F_2$  male hybrids showed elevated frequencies of SD alleles in both cross directions (fig. 1), indicating some incompatibility between these SD alleles and nuclear components of the AB genome. In characterizing the frequency of HMS in *T. californicus*, Willett (2008) found that crosses between AB females and SD



**FIG. 2**—Differences in allele frequencies between fertile and sterile *T. californicus* male  $F_2$  hybrids quantified with the  $G'$  statistic. Plot shows smoothed  $G'$  values estimated across 1-Mbp windows for  $AB♀ \times SD♂$  (teal) and  $SD♀ \times AB♂$  (red). Horizontal lines represent statistical thresholds of a genome-wide false discovery rate of 0.01. Total assembly lengths vary across the 12 *T. californicus* chromosomes (13–18 Mbp).

**Table 1**

GO biological process terms overrepresented in three QTLs identified on Chromosome 2. Allele frequency differences between fertile and sterile  $F_2$  male hybrids were greatest in the region associated with *QTL-2* in both cross directions. Listed are only the five most significant terms in each QTL with at least ten genomic annotations included (see [supplementary Tables, Supplementary Material](#) online for complete list)

GO term (region on chromosome 2)	GO term ID	Genomic annotations	Expected regional	Regional annotations	P-value
<i>QTL-1</i> (0.8–2.4 Mbp)					
Cellular response to manganese ion	0071287	10	0.06	2	0.002
DNA ligation	0006266	10	0.06	2	0.002
Maintenance of protein localization in the endoplasmic reticulum	0035437	15	0.09	2	0.004
Nucleotide-excision repair, DNA duplex unwinding	0000717	15	0.09	2	0.004
Regulation of endoplasmic reticulum unfolded protein response	1900101	16	0.1	2	0.004
<i>QTL-2</i> (3.4–5.7 Mbp)					
Spermatogenesis	0007283	107	2.15	9	<0.001
Protein ubiquitination	0016567	411	8.27	21	0.001
Ubiquitin-dependent protein catabolic process	0006511	310	6.24	14	0.004
Phosphatidylinositol phosphorylation	0046854	17	0.34	3	0.004
Phosphatidylinositol biosynthetic process	0006661	56	1.13	5	0.005
<i>QTL-3</i> (6.2–7.6 Mbp)					
Negative regulation of actin filament depolymerization	0030835	21	0.26	3	0.002
Regulation of hydrolase activity	0051336	525	6.54	15	0.002
Substrate adhesion-dependent cell spreading	0034446	57	0.71	6	0.002
Histone monoubiquitination	0010390	25	0.31	3	0.004
Regulation of intracellular steroid hormone receptor signaling pathway	0033143	58	0.72	4	0.006

males had a higher occurrence of  $F_2$  male sterility by 11–21% compared with parental populations. Furthermore, sterility was infrequent in  $F_1$  hybrids, and crosses between

either AB or SD and other populations did not lead to  $F_2$  HMS (Willett 2008). These phenotypic patterns suggest AB and SD genetic elements are particularly prone to

forming male sterility when recombined in  $F_2$  hybrids. Given that within the QTL the frequency of AB alleles in sterile males averaged  $<0.3$  (as opposed to the 0.5 expectation for all alleles in the  $F_2$  generation), we can infer a very strong effect at one or more genes in this region; indeed, if all SD homozygotes are sterile and heterozygotes are unaffected, we would expect an AB frequency of 0.33. Taken together with the GO enrichment analysis (table 1), our results suggest that deleterious interactions among SD and AB nuclear elements on chromosome 2 disrupt spermatogenesis and culminate in HMS.

The genetic basis of HMS differs from other forms of  $F_2$  hybrid breakdown and inviability in *T. californicus* (Burton and Barreto 2012; Lima et al. 2019; Healy and Burton 2020; Han and Barreto 2021). While mito-nuclear incompatibilities are disproportionately responsible for reductions in developmental rate, survivorship, and salinity tolerance in *T. californicus*  $F_2$  hybrids (Burton 1987, 1990a, 1990b; Edmands 1999), no mito-nuclear signature was associated with HMS. A similar pattern has been described in *Caenorhabditis* nematodes, where incompatibilities between mtDNA and nuclear genes contribute to hybrid inviability, but male sterility is the result of a single X-autosome incompatibility (Bundus et al. 2018). Because the AB and SD populations do not show evidence of HMS when crossed to other populations (Willett 2008), the genomic architecture of HMS appears to be more complex in *T. californicus* and involves population-specific interacting elements. This pattern is more akin to polygenic X-autosome interactions described in *Drosophila* clades (Presgraves and Meiklejohn 2021), and epistatic incompatibilities among autosomal loci in some plant taxa that culminate in HMS (Sweigart et al. 2006; Long et al. 2008; Feng et al. 2020). Overall, our results suggest that in some instances without sex chromosomes, nuclear incompatibilities on a single chromosome may make a disproportionate contribution to forming HMS and delineating species boundaries.

## Methods

### Interpopulation Crosses and Identification of Fertile and Sterile Male Hybrids

*Tigriopus californicus* were collected from intertidal rocky pools in Abalone Cove (AB; 33°44' N, 118°22' W) and San Diego (SD; 32°44' N, 117°15' W), CA. Copepods were maintained in mass cultures in 400 mL of 35 ppt seawater and fed ground commercial fish food and natural algal growth. Cultures were kept in incubators at 20 °C with a 12-h light/dark cycle. Males and females used in interpopulation crosses were sampled from different mass cultures so that replicate crosses reflected as much of the genetic variation present in parent populations as possible.

Forty-eight reciprocal crosses between AB and SD were established in 24-well culture plates, with a single male–female pair in each well. The resulting  $F_1$  interpopulation hybrids were separated as juveniles and raised to maturity in isolation.  $F_1 \times F_1$  crosses were conducted with a single pair per well, with an effort to maximize the mixing of offspring from different parental population crosses (i.e., no sibling crosses were performed).

The fertility of  $F_2$  interpopulation male hybrids was assessed by pairing them with a single female in two sequential trials. Females used to assess male fertility were from the population that matched the hybrid male's mitochondrial background. Approximately 1,000  $F_2$  male hybrids were evaluated for both cross directions. We sorted two classes of  $F_2$  male hybrids: a) Fertile males, which successfully produced offspring in both sequential trials, and b) Sterile males, which failed to produce offspring in both trials. Males that produced offspring with only one of the two females were excluded from the study.

### DNA Isolation and Sequencing

We collected fertile and sterile  $F_2$  hybrid males from both of the reciprocal crosses resulting in four pools: Fertile  $F_2$  males from  $AB \text{♀} \times SD \text{♂}$  crosses ( $n = 100$ ), Sterile  $F_2$  males from  $AB \text{♀} \times SD \text{♂}$  crosses ( $n = 88$ ), Fertile  $F_2$  males from  $SD \text{♀} \times AB \text{♂}$  crosses ( $n = 100$ ), and Sterile  $F_2$  males from  $SD \text{♀} \times AB \text{♂}$  crosses ( $n = 76$ ). DNA was isolated using a phenol:chloroform protocol (Sambrook and Russell 2006). Each pool was sequenced as 100-bp paired end libraries on the Illumina HiSeq 2000. Adapters were trimmed with Trimmomatic (Bolger et al. 2014) and only reads of at least 50-bp were kept after trimming.

### SNP Database Between Parental Populations

*Tigriopus californicus* has a reference genome that was developed de novo for the SD population where greater than 90% of the genome is anchored to 12 chromosomes (Barreto et al. 2018). Lima et al. (2019) generated an AB consensus genome by mapping AB reads from Lima and Willett (2018) to the most recent SD reference (GenBank accession: GCA\_007210735.1). Furthermore, Lima and Willett (2018) and Lima et al. (2019) identified population-diagnostic SNPs for SD and AB. These original analyses were conducted with a depth of coverage of  $\sim 40 \times$  and with reads sequenced from large pools of individuals from SD and AB ( $\sim 3,000$  and  $\sim 300$ , respectively) (GenBank accessions SD: SRX469409; AB: SRX2746703). Briefly, the SNP list was developed by a) performing reciprocal mapping of reads of a population to the reference sequence of the other, b) considering only those positions where all mapped reads showed an alternative nucleotide to the reference (i.e., fixed differences), and c) comparing the reciprocal mappings and keeping only SNPs that were fixed

differences in both mappings (Lima and Willett 2018). Here we used the SD and AB assemblies and the population-diagnostic SNPs developed by the previous authors.

In the current study, reads from the four hybrid pools were mapped reciprocally to both the SD and AB reference using Burrows-Wheeler Aligner (BWA) with default parameters (Li and Durbin 2009), and only reads with a mapping quality (MAPQ) score >20 were kept. PoPoolation2 (Kofler et al. 2011) was used to find positions across the genome where all reads had an alternative nucleotide to that of the reference, considering only biallelic positions with total coverage  $\geq 80$  and  $\leq 400$ . Read counts for every variable position were determined using PoPoolation2 (Kofler et al. 2011). Allele frequencies were calculated as the AB allele frequency in reciprocal crosses.

### Identification of Regions Associated With HMS

To assess the genomic architecture underlying HMS in *T. californicus*, we evaluated the allele frequency distributions of fertile and sterile  $F_2$  hybrid males with a two-step approach. First, we identified regions of the genome associated with HMS by quantifying differences in allele frequencies among fertile and sterile males within each of the reciprocal crosses (fertile vs. sterile  $AB\text{♀}\times SD\text{♂}$ ; fertile vs. sterile  $SD\text{♀}\times AB\text{♂}$ ) using a QTL approach in R 4.1.2 with the package “QTLseqr” (Mansfeld and Grumet 2018). We estimated the  $G'$  statistic described by Magwene et al. (2011) for analyzing bulk segregant data from next-generation sequencing under the default settings of the package and 1 Mbp nonoverlapping sliding windows. Varying window size (250 kb–1 Mbp) did not qualitatively change our results. Second, we distinguished the effects of nuclear and mitochondrial incompatibilities on HMS within these regions by evaluating allele frequency patterns among reciprocal crosses. We expected that if HMS was caused by incompatibilities between nuclear factors, allele frequencies in sterile males would be shifted in the same direction in both crosses. On the other hand, incompatibilities between the nuclear and mitochondrial genomes would be expected to skew allele frequencies in sterile males toward the population that mismatched the mitochondria (see Lima et al. 2019; Pereira et al. 2021).

### Enrichment Analysis

We used GO annotations (terms) to search for functional categories that were overrepresented in genomic regions associated with HMS. We focused on three 2-Mbp QTLs where differences in allele frequencies between fertile and sterile hybrid males were greatest and compiled gene identifications within these regions with an integrative genomics viewer (Thorvaldsdottir et al. 2013). We used a Fisher's exact test in the R package “topGO” (Alexa and Rahnenfuhrer 2021) to identify GO terms that were

overrepresented compared with the rest of the genome and report only those with  $P \leq 0.01$  and at least 10 genomic annotations.

### Supplementary Material

Supplementary data are available at *Genome Biology and Evolution* online (<http://www.gbe.oxfordjournals.org/>).

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### Author Contributions

T.G.L. and R.S.B. conceived the study. T.G.L. performed laboratory and bioinformatic processing. K.C.O. and F.S.B. conducted statistical analyses. K.C.O. wrote the manuscript. All authors contributed to interpreting results and editing the manuscript.

### Data Availability

The data supporting this article are available in NCBI's Sequence Read Archive (SRA) under accession numbers SRR24623753-SRR24623756.

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