Population differentiation and structural variation in the *Manduca sexta* genome across the United States

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Abstract

Many species that are extensively studied in the laboratory are less well characterized in their natural habitat, and laboratory strains represent only a small fraction of the variation in a species’ genome. Here we investigate genomic variation in 3 natural North American populations of an agricultural pest and a model insect for many scientific disciplines, the tobacco hornworm (*Manduca sexta*). We show that hornworms from Arizona, Kansas, and North Carolina are genetically distinct, with Arizona being particularly differentiated from the other 2 populations using Illumina whole-genome resequencing. Peaks of differentiation exist across the genome, but here, we focus on the most striking regions. In particular, we identify 2 likely segregating inversions found in the Arizona population. One inversion on the Z chromosome may enhance adaptive evolution of the sex chromosome. The larger, 8 Mb inversion on chromosome 12 contains a pseudogene which may be involved in the exploitation of a novel hostplant in Arizona, but functional genetic assays will be required to support this hypothesis. Nevertheless, our results reveal undiscovered natural variation and provide useful genomic data for both pest management and evolutionary genetics of this insect species.

Keywords: tobacco hornworm; Carolina sphinx moth; inversion; sex chromosomes; pseudogene; resequencing; population structure; agricultural entomology

Introduction

One of the main objectives of evolutionary genetics is to unite the proximate molecular function of genes with the long-term evolutionary forces that govern their change (Nei 1987). The 2 halves of this pursuit necessitate different approaches, with well-controlled laboratory studies best suited to elucidating the biochemistry of gene function and surveys of natural variation required to understand the ecological and demographic context in which these genes exist. The most fruitful species to study in the lab tend to be those with small body sizes and short generation times. In the field, common species with large population sizes are easiest to sample. Consequently, many of the most popular models for evolutionary genetics come from insects, which fulfill all of the above criteria.

For a brief example, consider the well-studied flies in the genus *Drosophila*. Early pioneers of evolutionary genetics noted that *Drosophila* collected from different locations often had differing arrangements of genes along their chromosomes, in other words, inversion polymorphisms within and between populations (Sturtevant 1917; Dobzhansky and Sturtevant 1938). But without the context provided by population genomic methods, these authors could only catalog the various structural rearrangements they observed; the evolutionary significance of these traits was unknown. With the benefit of modern sequencing technologies, researchers have since shown that segregating inversions like these often exist in environmental clines and contribute to maintaining fitness across a variety of environmental conditions (Anderson et al. 2005; Kapun et al. 2016; Wellenreuther et al. 2017). The initial discovery could only come from controlled laboratory crosses, but the significance could only be established by studying individuals taken from natural populations.

In spite of the above successes and notable others (e.g. mice: Hoeksstra et al. 2004; and monkeyflowers: Lee et al. 2016), species for which both robust natural and controlled studies exist remain rare. Certainly, the establishment of a new laboratory model species is a massive undertaking and model status can only be achieved through widespread adoption in the research community, which takes years if not decades. Conversely, population genetic sampling is only becoming more tractable with advances in sequencing technology. From this perspective, it is more logical to identify well-studied species that lack population genetic data and bring that natural context to them.

The tobacco hornworm moth (*Manduca sexta*) is a well-established laboratory model (Reinecke et al., 1980), and likely the second most studied moth after the domestic silkmoth, *Bombyx mori* (The International Silkworm Genome Consortium 2008). The tobacco hornworm is used to study developmental biology...
parts of the genome for further studies of establishment patterns of regional genetic variation and differentiated continental United States. With these resequencing samples, we sent whole-genome resequencing of this model insect to investigate management strategies. In service of both of these goals, we present how is a pest of economically important crops, including tobacco the nightshade family, Solanaceae (Wagner 2005). Thus, under range. Second, as the name tobacco hornworm suggests, benefit from understanding natural genetic variation across its range. McNeil 1995) and the ability of pupae to survive subfreezing temperatures (Shapiro 2006). Surprisingly, there have been no genetic studies of population structuring of M. sexta to date.

This lack of research into the population genetics of this moth is puzzling on 2 fronts. First, in terms of basic research, much of the ecological knowledge about this moth, such as acclimation and adaptation to different environments and host plants (Mechaber and Hildebrand 2000; Contreras et al. 2013) could benefit from understanding natural genetic variation across its range. Second, as the name tobacco hornworm suggests, M. sexta is a pest of economically important crops, including tobacco (Nicotiana spp.), tomatoes, peppers, and generally other plants in the nightshade family. Solanaceae (Wagner 2005). Thus, understanding how M. sexta populations are structured will inform pest management strategies. In service of both of these goals, we present whole-genome resequencing of this model insect to investigate population structure across an east-west transect of the continental United States. With these resequencing samples, we establish patterns of regional genetic variation and differentiated parts of the genome for further studies of M. sexta.

Methods
Sample collection and sequencing
For an initial characterization of genetic variation in this broadly distributed species, samples were chosen, based on availability, to maximize the geographic range represented while still sampling local populations with enough individuals to detect large changes in allele frequency. Specifically, samples from 3 states in the United States were examined in this project: North Carolina (n = 12), Kansas (n = 4), and Arizona (n = 8), see Fig. 1. Collecting methods varied somewhat between these locales. Samples from North Carolina were already obtained and sequenced as part of Mongue et al. (2019). In brief, adults were collected over the course of a week from a mercury vapor light trap in a single locale in July of 2017. Kansas samples were obtained from a mixture of adult moths caught at light traps and larvae taken from tomato fields in different towns in eastern Kansas throughout the summer of 2018. For larval sampling, only one individual was sequenced per field to minimize the risk of sequencing siblings. Arizona samples were obtained from the University of Florida’s collections, where adult moths had been preserved in ethanol. At time of collection, they were collected from a mercury vapor trap at one location over several nights in July of 2012. A summary of sequenced individuals and their methods of collection can be found in Table 1.

For each sample, tissue was extracted from adult thoracic flight muscle to avoid eggs and sperm in the abdomen, and larval head capsules to avoid hostplant tissue in the gut. DNA was extracted using an Omniprep kit for genomic DNA (G-Biosciences, St Louis, MO, USA), following the manufacturer’s protocol. DNA was isolated via phase-separation with chloroform and precipitated with ethanol. DNA pellets were resuspended in buffer and sent for Illumina sequencing with Novogene Sequencing (Sacramento, CA, USA). Quality control and library preparation followed the company’s standard and sequencing generated 150 bp, paired-end reads sequenced to an average depth of 14–20× coverage across the genome (also summarized in Table 1). As part of the sequencing service, the company trimmed raw reads for both quality and adapter sequence content.

Starting points for genomic analyses
Recent work on the M. sexta genome assembly has created both the opportunity and need for some comparative work on the multiple available assemblies. Manduca sexta has had an Illumina-only (i.e. fragmented) genome assembly for several years (Kanost et al. 2016); much more recently a chromosome-level assembly has been published (Gershman et al. 2021). Naturally, the chromosome-level assembly’s more contiguous scaffolds facilitate identification of large-scale variation; however, the previous assembly features high quality gene models informed by dozens of RNA-sequencing experiments and manually updated over the last 5 years (Kanost et al. 2016). To take advantage of the strengths of both assemblies, the same analysis pipeline was carried out on both assemblies to leverage the wealth of available data. Generally speaking, the newer assembly was employed for genome-wide investigation of differentiation and the older assembly was used to validate patterns observed in the former and investigate specific gene(set)s of interest. For both assemblies, the same processed short-read data were aligned to the reference using the very-sensitive-local alignment setting in Bowtie2 version 2.2.9 (Langmead and Salzberg 2012). Alignments
Table 1. Sample collection and alignment information.

| Sample | State | Location | Collecting method | Sex | Overall alignment rate (bowtie2 vsl) | Coverage depth (x) |
|--------|-------|----------|-------------------|-----|-------------------------------------|-------------------|
| A36    | AZ    | Madera Canyon | Blacklight trapped adult | M   | 94.08% | 18 |
| A70    | AZ    | Madera Canyon | Blacklight trapped adult | M*  | 94.13% | 18 |
| A71    | AZ    | Madera Canyon | Blacklight trapped adult | F*  | 93.60% | 17.8 |
| A76    | AZ    | Madera Canyon | Blacklight trapped adult | F   | 93.61% | 13.7 |
| A78    | AZ    | Madera Canyon | Blacklight trapped adult | M   | 94.00% | 14.7 |
| A82    | AZ    | Madera Canyon | Blacklight trapped adult | M   | 94.05% | 14.5 |
| A84    | AZ    | Madera Canyon | Blacklight trapped adult | M   | 94.18% | 14.2 |
| A85    | AZ    | Madera Canyon | Blacklight trapped adult | M   | 94.20% | 14 |
| WK2    | KS    | Williamstown | Larva from hostplant | F   | 93.65% | 14.5 |
| KC1    | KS    | Kansas City | Adult from hostplant | F   | 93.77% | 15.9 |
| LK5    | KS    | Lawrence    | Larva from hostplant | F   | 94.04% | 16.2 |
| GK3    | KS    | Garnett     | Larva from hostplant | M   | 94.55% | 18 |
| S32    | NC    | Rocky Mount | Blacklight trapped adult | M   | 94.12% | 18.13 |
| S33    | NC    | Rocky Mount | Blacklight trapped adult | M   | 93.90% | 19.82 |
| S34    | NC    | Rocky Mount | Blacklight trapped adult | M   | 93.84% | 19.8 |
| S35    | NC    | Rocky Mount | Blacklight trapped adult | M   | 94.02% | 16.93 |
| S36    | NC    | Rocky Mount | Blacklight trapped adult | M   | 94.04% | 19.13 |
| S37    | NC    | Rocky Mount | Blacklight trapped adult | M   | 94.06% | 20.08 |
| S38    | NC    | Rocky Mount | Blacklight trapped adult | M   | 94.08% | 22.44 |
| S39    | NC    | Rocky Mount | Blacklight trapped adult | M   | 94.01% | 26.16 |
| S40    | NC    | Rocky Mount | Blacklight trapped adult | M   | 93.79% | 19.72 |
| S42    | NC    | Rocky Mount | Blacklight trapped adult | M   | 94.12% | 25.67 |
| S44    | NC    | Rocky Mount | Blacklight trapped adult | M   | 94.07% | 22.10 |
| S45    | NC    | Rocky Mount | Blacklight trapped adult | M   | 94.16% | 20.49 |
| Q6     | NC    | Rocky Mount | Blacklight trapped adult | M   | 85.44% | 11.2 |

Sample names follow sample labeling in SRA. States are abbreviated as AZ (Arizona), KS (Kansas), and NC (North Carolina). Basic genomic information on overall alignment rates to the published M. sexta genome (Kanost et al. 2016) and depth of sequencing coverage are given as well. All sequencing data are available with the following accession: SRP144217, PRJNA639154 on NCBI. Individuals with an * next to their sex did not have sex information recorded during sampling; however, we were still able to assign sex based on coverage ratios between the Z chromosome and autosome after sequencing. Q6 is the M. quinquemaculata sample used as an outgroup to polarize allele frequencies.

were coordinate sorted and optical duplicates were removed with Picardtools v. 2.8 (Wysoker et al. 2013). Finally, alignments around insertions and deletions were adjusted with GATK v. 3.7 (McKenna et al. 2010). This end result generated a set of curated alignment files (bams) that were the starting point for all downstream analyses.

For some of these analyses, it was necessary to analyze the genotypes of single nucleotide variants. To generate sets of high-confidence variants, the curated alignments were taken through GATK v.3.7’s best practices pipeline for SNP calling (McKenna et al. 2010), including hard quality filters (quality by depth > 2.0 and Fisher Strand-bias < 60 and mapping quality > 40). The end result was a variant call format (vcf) file containing SNPs for each population. This filtering resulted in a total of 16,674,625 SNPs resulting was a variant call format (vcf) file containing SNPs for each species. These putative divergences were combined with the polymorphism data from M. sexta to create a new "ancestral" reference (fasta) for use in polarizing allele frequencies, using the following parsimony logic.

In a majority of cases, polymorphic sites did not overlap with divergent sites as well. In other words, these sites that varied in M. sexta but not in M. quinquemaculata showed the reference allele in the outgroup; thus, the reference allele was taken to be ancestral and left as was. For each site that varied in both M. sexta and M. quinquemaculata, allele identities were compared between species. If the alternate allele in M. quinquemaculata was shared with the alternate allele in M. sexta the alternate allele was inferred to be ancestral, and the reference derived. Likewise, if the outgroup was heterogeneous at the site, but shared only one allele with the focal species, that allele was inferred to be ancestral (e.g. a tri-allelic site with reference: T, M. sexta alternate allele: G, M. quinquemaculata alternate alleles: G, C). Considering both of these scenarios, a total of 1,327,986 sites were updated in the reference. In a minority of cases (e.g. a tri-allelic site with reference: T, M. sexta alternate allele: C, M. quinquemaculata alternate allele: G), there was no most-parsimonious solution. These sites were updated to "N" at the site in the new ancestral reference to mask them from analysis (n=275,410 sites). The code to generate the updated reference was written in R and can be found in the git repo listed in Data accessibility.

Updating genomic resources

In addition to preparation of resequenced samples and variants, the newer reference genome required ancillary metadata, namely: the assignment of scaffolds to chromosomes to separate chromosome-specific effects (e.g. differing dynamics on the Z sex chromosome). Curiously, the newer M. sexta reference contains no chromosomal linkage information for its scaffolds. The older assembly does have linkage information however, from previous ortholog counting analyses comparing M. sexta to B. mori to identify and separate sex-linked scaffolds; this effort generated a
chromosome identity for each scaffold, but did not contain information about the ordering of scaffolds within a chromosome (Mongue and Walters 2018). To anchor the newer assembly, it was aligned to the older one using SatsumaSynteny in Satsuma v.3.1 (Grabherr et al. 2010). This tool generated a list of alignments between the older, anchored assembly and the newer, more contiguous but unanchored assembly. Chromosomes were assigned to the new assembly’s scaffolds by tabulating the number of fragmented (older) scaffolds aligning to each chromosome-length scaffold and taking the highest scoring chromosome. This effort generated unambiguous assignment of the largest scaffolds in the newer assembly (which should be mostly complete chromosomes) as well as putative assignments for the other, unplaced scaffolds. These placements were mostly unambiguous based on syntenic alignments alone, but in the case of 2 ambiguities, additional information from BAC-FISH probes between M. sexta and B. mori were used to resolve chromosomal assignment (Yasukochi et al. 2009). As this information may be of future use beyond the scope of this research, both this assignment and the alignment summary file used to generate it can be found in the Supplementary Files.

Population structure
With linkage established, alignments were filtered to separate out the Z chromosome, as unequal sampling of sexes between populations (e.g. North Carolina’s sample is male only) could create spurious patterns. The autosomal genome was used to infer population structure while accounting for uncertainty in genotyping using the software NGSAdmix v. 3.2 (Skotte et al. 2013) as well as a principle-component-based approach with PCAngsd v.0.98 (Meinier and Albrechtsen 2018). For the admixture analysis, the best-supported number of populations (K) was determined by assessment of ten bootstrapped replicates each for possible K values from 2 to 6, evaluated using the logic from CLUMPACK’s web-hosted application (Kopelman et al. 2015). After a first round of analyses revealed a large (~8 Mb) block of differentiation on chromosome 12 segregating in population, structuring analyses were repeated with this chromosome removed as well.

Characterization of genomic differences
After establishing basic population relationships, population differences were examined in detail. A baseline expectation of differentiation was set by estimating the genome-wide pairwise $F_{ST}$ (excluding the Z chromosome) analyzing the curated alignments with the population genetic software ANGSD v. 0.916 (Korneliussen et al. 2014). In more detail, we input into ANGSD the list of curated bam files and the parsimony-informed ancestral reference sequence and generated site allele frequency likelihoods ( -dosaf 1) separately for each population using the SAMtools method of genotype likelihoods ( -gl 1). Next, a 2-dimenational site frequency spectrum was generated for each pair of populations for a total of 3 comparisons using the realSFS command. With the relevant site frequency spectrum as a prior, we used the -fstout command to generate genome-wide $F_{ST}$ on using both assemblies as references. Additionally, we estimated $F_{ST}$ separately for silent (4-fold degenerate) and nonsilent (zero-fold degenerate) sites to assess whether differentiation skewed toward silent sites or those more likely to be under selection. Identification of zero-fold and 4-fold degenerate sites was completed in previous analyses with custom R scripts (in R v 3.3, R Core Team 2017; Mongue et al. 2019) but was only carried out on the Kanost et al. assembly due to the higher quality gene models.

With a baseline established, candidates for further investigation were identified via a preliminary scan for $F_{ST}$ outlier regions in windows of 10 kb with a 5-kb step size to identify regions for further study (using the fst command again with -win 10,000 -step 5,000). All pairwise comparisons were performed: North Carolina (NC)—Arizona (AZ), Kansas (KS)—Arizona, and NC—KS. For these analyses, published chromosome assignments of scaffolds were employed for the older assembly (Kanost et al. 2016; Mongue and Walters 2018; Mongue et al. 2022) and assignments generated for this manuscript were used for the new assembly.

For the older assembly, these assignments did not contain information on ordering of scaffolds within a chromosome. Thus, we focused primarily on the new assembly for identification of differentiated regions, but we performed analogous $F_{ST}$ analyses with alignments to the old assembly to assess differentiation in and around coding regions.

These analyses revealed multiple large $F_{ST}$ peaks, as would be expected of segregating inversions. To further investigate these regions and characterize which population(s) carried the ancestral or derived orientation in the putatively inverted regions, genetic diversity was examined as follows. Single-nucleotide polymorphism (SNP) frequencies were examined in each population, under the logic that if loci within an inversion cannot freely recombine with the noninverted orientation, they should share allele frequencies across longer distances than those outside of the inversion. For these analyses, the polarized allele frequencies described above were used. SNP variant files (vcfs) were generated for each population and alternate allele frequencies were extracted and examined. Beyond allele frequency, the allele identity was also examined. The VCFtools utility “extract-FORMAT-info GT” (Danecek et al. 2011) was used to examine sample genotypes directly and determine which individuals carried nonreference alleles across genetic features of interest. Finally, to further evaluate the interpretation of these regions as inversions, a pair of individuals ostensibly carrying the inversions was investigated with Delly2 v 0.8.7 (Rausch et al. 2012) to identify structural variants and potential inversion breakpoints. Because Delly makes inferences based on read-pair alignment positions, the individuals with the deepest coverage of those carrying the potential inversions were chosen (A36 and A70) to compare to a pair ostensibly lacking the inversion (S39 and S42).

Finally, in the interest of establishing baseline metrics for these populations, we calculated a number of population genetic summary statistics. In particular, we calculated measures of variability: pN (nonsynonymous changes per nonsynonymous site), pS (synonymous changes per synonymous site), pN/pS the scaled ratio of nonsynonymous variation, $\pi_0$ (variation at zero-fold degenerate sites), $\pi_4$ (variation at 4-fold degenerate sites), Tajima’s $D_0$, and Tajima’s $D_4$ (estimators of deviation from neutral evolution at zero- and 4-fold degenerate sites, respectively). These metrics were all calculated using alignments to the Kanost et al. assembly for reasons of gene model quality described below. Metrics related to pN/pS relied on SNP calls from the vcf files but estimates of $\pi$ and Tajima’s D were performed directly from alignments using angd’s “saf2theta” command. In addition, we calculated $\rho^2$ linkage disequilibrium for nonoverlapping 50-bp windows across the more contiguous Gershman assembly using the vcftools utility “–geno-r2”. These results are reported in Table 3 but we refrain from formal statistical comparisons between populations as the uneven sampling effort between them makes estimates of variation difficult to compare directly.
Genetic characterization of differentiated regions

For each genomic region-of-interest, annotated genes in the region were examined based on the original assembly; this version’s annotations have a high degree of manual curation not present in the newer assembly (Kanost et al. 2016). Manduca sexta has a wealth of available RNA sequence data from both sexes and different life-stages, aggregated as a set of RNAseq collected by Cao and Jiang (2017) and quantified with this older reference. From this dataset, the specificity of gene expression was calculated for each gene in the genome for life stage, tissue, and sex specificity (following methods in Mongue et al. 2022) to inform potential function of genes of interest. More precisely, the specificity metric (SPM) was used to assess the proportion of a given gene’s expression found in a specific class. This metric ranges from zero to one, inclusive, and denotes the proportion of a given gene’s expression unique to larva, pupa, or adult of tissues collected in all 3 stages (head, midgut, and fat bodies).

Stage specificity was defined as gene expression unique to larva, pupa, or adult of tissues collected in all 3 stages (head, midgut, and fat bodies). Tissue specificity within a stage was based on tissue types available for that stage (see Table 2). And, naturally, sex specificity was based on amount of expression unique to males or females. For instance, a gene with an adult SPM of 0.90 shows 90% of its expression in adults as opposed to pupae or larvae. The same gene has separate SPM values that denote how specific its expression is to males or females and which tissues of those individuals. This method has the benefit of generating a sense of gene function independent of functional annotation based on sequence homology with existing annotations.

All genes across the genome were annotated with SPM values; however, for large outlier regions spanning numerous genes, it was not feasible to examine individual genes in depth. Instead, using the above annotations, these regions were examined for biased composition compared to nondifferentiated regions. In other words, enrichment or depletion of functional classes within putative inversions was tested. For small regions containing one or a few genes, each could be manually examined. In these cases, SNPs in and around these genes were examined for evidence of coding-sequence variation between populations. SNPs were annotated with the tool SNPeff v. 4.2, which uses gene models to predict the coding sequence changes introduced by polymorphisms (e.g. loss of a canonical stop codon, Cingolani et al. 2012).

Results

Chromosomal assignment of the new M. sexta assembly

The older, Illumina-only assembly of M. sexta already possesses chromosomal assignment based on syntenic alignments to B. mori, a well-studied moth in a sister family to M. sexta. It has previously been shown that these 2 species share not only a conserved karyotype (n = 28) but also extensive conservation of sequence along these chromosomes (Yasukochi et al. 2009). Initially, we compared synteny between the 2 assemblies of M. sexta to maximize similarity and thus alignment information. From these alignments, 25 of the 28 largest scaffolds in the new assembly matched cleanly and overwhelmingly to one chromosome. The remaining 3 were more complicated and resolved as follows.

HiC_scaffold_31 showed synteny to both chromosome 2 and chromosome 26, in a way that suggested a potential fusion or misassembly, i.e. the first ~9 Mb of the scaffold was syntenic to chromosome 26 and the remaining ~6 Mb to chromosome 2. Closer inspection of published BAC-FISH mapping reveals that there has been structural turnover between species and both B. mori chr2 and chr26 are orthologous to a fused M. sexta chr2 (Yasukochi et al. 2009); thus this scaffold was correctly assembled and was manually annotated as chromosome 2.

Second, both HiC_scaffold_2 and HiC_scaffold_25 showed homology to B. mori chromosome 11 almost exclusively. Again, the BAC-FISH mapping revealed the reason for the anomaly. Chromosome 11 in B. mori is split in M. sexta into the larger chromosome 11 and smaller chromosome 26 (Yasukochi et al. 2009). Based on the relative sizes of these scaffolds, we annotated the larger (~10 Mb) HiC_scaffold_2 as chromosome 11 and the smaller HiC_scaffold_25 (~5 Mb) scaffold to chromosome 26.

With these manual annotations, each of the largest 28 scaffolds in the new assembly is now anchored to one of the 28 chromosomes in M. sexta. Smaller, more fragmented scaffolds were also assigned to chromosome based on syntenic alignment counts, but without manual curation. These assignments are presented in a Supplementary File.

Table 3. Population genetic parameters across the autosomes of the populations surveyed here.

| Population genetic summary statistics for each population surveyed |
|---------------------------------------------------------------|
| | North Carolina | Kansas | Arizona |
| pN | 0.0068 | 0.0044 | 0.0063 |
| pN/pS | 0.2850 | 0.2852 | 0.2851 |
| r0 | 0.0027 | 0.0029 | 0.0030 |
| r4 | 0.0218 | 0.0204 | 0.0210 |
| Tajima’s D0 | -0.0795 | -0.0367 | -0.0673 |
| Tajima’s D4 | -0.0259 | -0.0148 | -0.0259 |
| ρ² | 0.3538 | 0.5990 | 0.4053 |

The Z chromosome was omitted due to uneven sampling of sexes between populations. Median values are given for polymorphism (pN and pS, to avoid skew from outliers), while means are reported for r and Tajima’s D (as in every case, the median value is centered on zero). For all statistics involving silent and replacement sites (i.e. everything but linkage), the older Kanost et al. assembly was used to take advantage of higher confidence gene models. Mean linkage disequilibrium (ρ²) reported for 50 bp windows using the Gershman et al. assembly to take advantage of more robustly supported linkage groups. Standard deviations appear in parentheses.

Table 2. Tissue-specific RNA sequencing available for each life-stage of M. sexta.

| Antenna | Head | Muscle | Midgut | Malphigian Fat Gonad tubes |
|---------|------|--------|--------|---------------------------|
| Larva: | | | | |
| Adult: | M | F | M | F |
| Pupa: | | | | |
| M | F | M | F |

This dataset forms the basis for tissue specificity labeling annotation of genes in this study. Grey cells indicate presence, empty cells denote absence of data. If sex-specific data are available, it is noted with an M (for male) and F (female) in the relevant cell. For larvae, few tissues were sampled at multiple instars, so no further developmental division was made.

Table 4. Genetic characterization of differentiated regions.

| Genetic characterization | Chromosome | Population genetic parameters across the autosomes of the populations surveyed here. |
|--------------------------|------------|----------------------------------------------------------------------------------|
| | North Carolina | Kansas | Arizona |
| pN | 0.0068 | 0.0044 | 0.0063 |
| pN/pS | 0.2850 | 0.2852 | 0.2851 |
| r0 | 0.0027 | 0.0029 | 0.0030 |
| r4 | 0.0218 | 0.0204 | 0.0210 |
| Tajima’s D0 | -0.0795 | -0.0367 | -0.0673 |
| Tajima’s D4 | -0.0259 | -0.0148 | -0.0259 |
| ρ² | 0.3538 | 0.5990 | 0.4053 |

The Z chromosome was omitted due to uneven sampling of sexes between populations. Median values are given for polymorphism (pN and pS, to avoid skew from outliers), while means are reported for r and Tajima’s D (as in every case, the median value is centered on zero). For all statistics involving silent and replacement sites (i.e. everything but linkage), the older Kanost et al. assembly was used to take advantage of higher confidence gene models. Mean linkage disequilibrium (ρ²) reported for 50 bp windows using the Gershman et al. assembly to take advantage of more robustly supported linkage groups. Standard deviations appear in parentheses.

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Structure
We found 3 populations via the heuristic of Evanno et al. (2005); namely, the most probable K is the one the biggest change in likelihood from the previous K value. Per this metric, each of the 3 states’ moths form distinct populations, although there was some apparent admixture between Arizona and Kansas, including an Arizona individual confidently grouping with the Kansas samples (Fig. 2a).

Taking a complimentary approach, we assessed principal components of SNP variation across our samples and observed the same clustering pattern as found in the structure analysis. Each state’s samples do not cluster with other states, but Arizona samples showing a more dispersed pattern than the others, including some individuals more similar to Kansas individuals than other Arizona individuals (Fig. 2b). When excluding chromosome 12, samples sort by collecting location more cleanly, both in admixture analyses and principal component analyses (Fig. 2c and d), suggesting the signature of apparent admixture was driven by mainly the inversion presence or absence. However, even without chromosome 12, one Arizona sample shows some inferred admixture with Kansas, suggesting there may be smaller variable genomic regions that still generate more similarity between the 2 populations.

Differentiation across the genome
To start, we provide population genetic summary statistics for each state in Table 3. Due to the unequal sampling effort between populations, we do not formally test for statistical differences between populations. And indeed, absolute values for many measures of variation (e.g. pS or n) are lowest in Kansas, the population with the fewest sampled individuals. Still, the relative amount of nonsynonymous variation scaled by synonymous variation (pN/pS) is remarkably consistent across populations, suggesting the strength of selection relative to drift is comparable across states. Moreover, Tajima’s D is more negative at zero-fold sites (i.e. those exposed to selection) than 4-fold (putatively neutral sites) in each population, which further implies a role for directional selection in each population. Finally, apparent linkage disequilibrium is inversely correlated with sample size, as expected if artificially tight associations are observed between common alleles due to shallower sampling failing to capture rare alleles.

A simple genome-wide average of $F_{ST}$ set a baseline level of differentiation of 0.04–0.09, depending on the pairwise comparison between states (Table 4). A sliding window scan across the genome revealed a few regions with markedly higher $F_{ST}$ (>0.5, Fig. 3). From this analysis, several regions of interest became apparent. Most notably, a large proportion of chromosome 12 shows high levels of differentiation between Arizona and the other 2 populations. The Z chromosome showed the strongest peak of elevated differentiation, and chromosome 19 showed another above our threshold level. We compared these results on the newer assembly to a parallel $F_{ST}$ scan on the older assembly (Supplementary Fig. 1). As expected, this turned up many differentially scaffolds assigned to chromosome 12, the Z, and 19, but also 2, 6, and 13. In the following sections, we describe each of these regions in detail and resolve apparent discrepancies.

An inferred segregating inversion on chromosome 12
We observed a large $F_{ST}$ peak across a substantial portion of chromosome 12 in the more contiguous assembly (Fig. 4). This peak corresponds with the Delly2 annotation of an inversion located on HiC_scaffold_12 from 5,414,046 to 13,233,866 bp, a total of 7.82 Mb. Samples from North Carolina annotated with Delly lacked this called inversion, as expected given that the reference
individual was sourced from lab stock which was originally collected in North Carolina.

In the old assembly, this peak correspond to 13 scaffolds, listed in Supplementary Table 2 and plotted in Supplementary Fig. 2, summing to 7.6 Mb of sequence, approximately 56% of the total (assigned) length of the chromosome. The peaks contain 321 annotated genes. Note the discrepancy of ~200 kb of differentiated sequence.

The putative inversion’s frequency of 0.6875 corresponds to 11–16 sampled chromosomes with the inversion in the Arizona population. Per investigation of sample genotypes, this variation is distributed as 4 individuals homozygous for the nonreference orientation (A36, A70, A78, A82), 3 heterozygous for it (A71, A76, A84), and one homozygous for the ancestral orientation (A85). Note that the heterozygous individuals correspond to those inferred to be admixed with Kansas and the one homozygous non-inverted individual was grouped with Kansas completely in structure analyses. As discussed above, removing chromosome 12 from structuring analyses leads to less similarity between AZ and KS samples, suggesting the inversion presence/absence is large portion of the signal of the Arizona population. A scan of linkage disequilibrium along the chromosome (Supplementary Fig 3) showed elevated linkage within the bounds of the hypothesized inversion in each of the 3 populations, suggesting that this region resists gene flow between populations.

Table 4. Genome-wide point estimates (excluding the Z chromosome) for differentiation between each pairwise combination of the 3 populations studied here.

| F stk | NC | KS |
|------|----|----|
| 0 x  |    |    |
| 4 x  |    |    |
| KS   | 0.064, 0.054 | 0.041, 0.044 |
| AZ   | 0.090, 0.081 |    |

In each comparison differentiation at sites exposed to selection (i.e. zero-fold degenerate sites, 0 x) and putatively neutral sites (4-fold degenerate sites, 4 x) were estimated separately in angsd using the gene models from the Kanost et al. assembly.

The large size of this inferred inversion (roughly 8 Mb and 321 genes contained within) makes identifying a causal adaptive locus challenging. We used the RNA sequencing metadataset to classify genes as specific to one sex (male or female), life stage (larva, pupa, or adult), and tissue (antennae, head, fatbody, mid-gut, Malpighian tube, muscle, ovaries, or testes) if a given gene showed 70% or more of its expression exclusively in that category. By this metric, the putative inversion on chromosome 12 does not differ in sex-biased composition ($X^2_2 = 1.31, P = 0.518$), lifestage-biased composition ($X^2_6 = 2.06, P = 0.560$), or tissue-biased composition ($X^2_7 = 7.45, P = 0.384$) compared to the rest of the genome. Thus, even if there is a set of locally adaptive alleles in this region, it is not one enriched for any of the annotations available to us. Nevertheless, we identified one gene of interest in this region thanks to a misassembly.

Both the ostensible chromosome 2 and 13 peaks from the more fragmented assembly belong to the putatively inverted region of chromosome 12 in the newer assembly. The former is rather unremarkable, some 50 kb of a scaffold containing 2 annotated genes (Msex2.02572 and Msex2.02573) with highly differentiated SNPs localized to areas outside the coding region of either. These variants may be truly neutral or impact the regulation of the nearby genes, but without a better understanding of regulatory elements (e.g. promoter sequences) in this species, it is impossible to say. The peak previously assigned to chromosome 13, deserves more consideration however.

A potential pseudogene within the chromosome 12 inversion

Despite overall congruence of results from the old and new assembly, one difference in the F stk scans proved particularly interesting. The older assembly indicated a peak of differentiation on a scaffold anchored to chromosome 13; however, none of the more contiguous chromosome 13 is particularly differentiated using the new assembly as a reference. To reconcile this contradictory pair of results, we investigated the peak in the old assembly first, then placed it in new assembly.

![Population differentiation across the genome](image.png)

**Fig. 3.** Pairwise differentiation ($F_{ST}$) across the *M. sexta* genome. North Carolina (NC) versus Kansas (KS, black squares), Kansas versus Arizona (AZ, blue diamonds), and North Carolina versus Arizona (red points). Plotting is ordered by chromosomal linkage. For ease of visualization, the genome is split into 2 rows. We examined regions enriched for differentiation greater than 0.5 (dashed line). By this metric, one peak on the Z, and one peak each on chromosomes 12 and 19. Additionally, note that the highest peaks almost always come from KS versus AZ and NC versus AZ, indicating that Arizona’s population is more distinct from the others than NC and KS are from each other. Chromosomes are numbered based on syntenic assignment of HiC scaffolds from the Gershman assembly. Unplace scaffolds are all those beyond the 28 main chromosomal scaffolds.
This peak comes from one scaffold (scaffold00381) starting around 200 kb and continuing for the remaining 100 kb (Fig. 5a). This region contains a single gene, Msex2.10493, and it displays a striking pattern of differentiation between states. Within population variation is low in Kansas and North Carolina, with only 1 and 3 coding-sequence polymorphisms observed, respectively; in contrast, 61 SNPs were called in the gene body from the Arizona samples (summarized in Fig. 5b). Moreover, many of these SNPs are at high frequency (Fig. 5c), including the predicted loss of a stop codon.

The massive difference in called variants theoretically could arise from population-specific differences in mapping rates in this region, either through poor mapping in populations with low SNP counts or (far less likely) a pile-up of reads in Arizona owing to a collapsed segregating paralog. To rule out these artefactual explanations, we examined per-base coverage around this gene and found no differences between populations (Fig. 5d; note that slightly deeper coverage in NC is consistent with genome-wide differences in sequencing effort between the populations). Thus, Arizona M. sexta appear to truly differ from the other populations with regard to the molecular evolution of this gene, which begs the question of the gene's functional role.

The final question to consider was why such a striking pattern of difference is absent in the newer assembly. To answer this, we examined the syntenic alignment of scaffold00381 to the more contiguous assembly. A discrepancy immediately arose: roughly 2/3rds of the scaffold aligned to HiC_scaffold30 (which we have anchored as chromosome 13), but the remaining 1/3rd, including the coding region for Msex2.10493, aligns to chromosome 12 (Supplementary Fig. 4). This pattern is suggestive of a misassembly in the older reference genome. Furthermore, the roughly 100 kb length of this apparently chromosome 12 sequence accounts for half of the 200 kb discrepancy in the estimated chromosome 12 differentiation peak between the 2 assemblies.

More direct evidence suggests that this potential pseudogene lies within the chromosome 12 inversion. First, the coding sequence of the old assembly's Msex2.10493 is homologous to the new assembly's XM_037442453.1, which lies in the inverted region (roughly 9.257–9.261 Mb into chromosome 12). Additionally, the frequency of the stop-loss is 0.6875 (¼ 11/16) and matches that of the most common SNP frequency within the inversion. Moreover, the genotypes (homozygous present, heterozygous, homozygous absent) of the inversion perfectly match the stop-loss variant's genotype. In conclusion, this potential pseudogene sits in the middle of the inferred inversion on chromosome 12, but it was only identified thanks to a serendipitous misassembly in the older version of the M. sexta reference.

The Z holds another putative segregating inversion

The Z chromosome’s outliers are attributable to roughly 1 Mb in the middle of the chromosome-level scaffold (HiC_scaffold_19). The scale of elevated differentiation suggests another inversion, and our structural variant scan identified an inversion that precisely overlaps this region (from 13,706,400 to 14,710,588 bp, for a total length of 1,004,188 bp). Further evidence for a lack of recombination in this region can be seen in the nonreference allele.

This region contains a single gene, Msex2.10493, and it displays a striking pattern of differentiation between states. Within population variation is low in Kansas and North Carolina, with only 1 and 3 coding-sequence polymorphisms observed, respectively; in contrast, 61 SNPs were called in the gene body from the Arizona samples (summarized in Fig. 5b). Moreover, many of these SNPs are at high frequency (Fig. 5c), including the predicted loss of a stop codon.
frequencies which vary across the scaffold in both NC and AZ but show a tract of SNPs with a common frequency of 0.93 (¼ 13/14) within the putative inverted region in Arizona (Fig. 6).

This peak of differentiation is also identifiable in the older assembly as a roughly 1 Mb region of scaffold00022, suggesting we are capturing essentially all of the inversion on one scaffold, even in the more fragmented assembly (Supplementary Fig. 5).

Chromosomes 6 and 19 show differentiation but no clearly differentiated genes

Chromosome 6 shows a single peak of divergence, though this region is only weakly differentiated, barely meeting our cut-off threshold for exploration (\(F_{ST} > 0.5\) between any 2 populations). This region contains 3 full annotated genes (Msex2.12148, Msex2.12149, and Msex2.12150), but no SNPs within coding regions.

Two outlier scaffolds in the older assembly localize to chromosome 19; both represent differentiation between Arizona and the other 2 states. In the newer, more contiguous assembly, these peaks remain distinct from each other (Supplementary Fig. 6), suggesting they are not parts of a common feature. One is localized to scaffold00118, to approximately the last 400 kb of an ~850-kb scaffold. This region contains 11 annotated genes (Msex2.05775-Msex2.05786), but none of these shows skews in coding region SNPs between populations as extreme as seen on the potential pseudogene mentioned above. However, 2 genes do show more modest deviation between states. Msex2.05777 is variable in both North Carolina (\(pN = 6, pS = 10\)) and Kansas (\(pN = 7, pS = 12\)) but monomorphic (\(pN = 0, pS = 0\)) in Arizona. Conversely, Msex2.05784 is nonvariable in the North Carolina and Kansas states.

Fig. 5. Extreme population differentiation of a gene within chromosome 12’s putative inversion. a) Finer-scale plotting of \(F_{ST}\) of the differentiated Kanost et al. scaffold, previously assigned to chromosome 13 shows that differentiation occurs beyond 200 kb and, as with the inversions above, appears that Arizona is markedly different from North Carolina and Kansas. Boxes at the bottom of the plot show syntenic alignments with the Gershman et al. assemble and reveal that the differentiated region belongs to chromosome 12 rather than 12. A single annotated gene, Msex2.10493, lies in this region (gray box). b) Annotation of SNPs in the coding region reveals a stark difference in variation between populations. Arizona moths hold more than 10 times the nonsynonymous polymorphisms than those in the other 2 states. c) Frequencies of the SNPs in Arizona. Gray boxes represent exons and the dashed vertical line marks a variant that disrupts the annotated stop codon. d) Depth of coverage of the gene in question does not vary between populations in a way that suggests differences in alignment between populations can explain the differences in called SNPs.
samples but contains 3 nonsynonymous variants and one synonymous variant in Arizona.

The second peak, on scaffold00419, seems to span the length of this 200 kb scaffold, but this may reflect the limits of resolution of our FST window size. In any case, the entire scaffold contains 11 annotated genes, many of which have no expression data and no called SNPs in any population. As such, we do not place much emphasis on the differentiation around these dubiously annotated regions.

Discussion

We conducted whole-genome resequencing to investigate the natural variation of M. sexta populations across the United States. In brief, we found that individuals from 3 populations across an east-west transect are genetically differentiated. This result may not seem surprising, but studies of other Lepidoptera have at times found evidence for continent-wide panmixis (specifically for monarch butterflies, Lyons et al. 2012; Zhan et al. 2014). And while monarchs may be an extreme case of lepidopteran long-distance travel, sphinx moths, including M. sexta, are renowned for their flight and dispersal ability, both in terms of small-scale speed and dexterity (Stevenson et al. 1995) and long-distance dispersal (Janzen 1984; Haber and Frankie 1989). Indeed, M. sexta has been posited as migratory in the past (Ferguson 1991). Our results, however, suggest that gene flow is less common than expected of a species with routine migration. In fact, the genome-wide mean FST for the least differentiated pair (0.04, Kansas—Arizona) is on par with the differentiation between migratory North American monarchs and the Hawaiian population that is thought to have split roughly 200 years ago (Lyons et al. 2012).

More striking than this level of differentiation is the nature of the differences that contribute to it. Namely, large (>1 Mb) putative inversions, are key contributors to the variation between populations. Sustained tracts of differentiation are most apparent on 2 chromosomes, the Z chromosome and chromosome 12. In both cases, the Arizona population is distinct from Kansas and North Carolina. We note that the smaller sample size of Kansas M. sexta decreases the precision of allele frequency estimation (Lou et al. 2021), and may obscure some of the more subtle population differences for the time being. For the present study, we discuss in detail the genomic and genetic nature of the large differences, with the caveat that functional genetics will be required to truly confirm the genomic structure and downstream fitness consequences of this variation.

Minor differentiated regions of the genome

The inversions discussed below are the most apparent sources of regional differentiation, but chromosomes 6 and 19 also showed elevations in differentiation at smaller scales. For the most part, differentiated regions fell outside of coding regions or near dubiously annotated genes (i.e. those not supported by expression data). Lacking further information for hypothesized importance, they are not discussed in detail here for brevity’s sake.

The largest signal for population differentiation: a likely inversion on chromosome 12

A long tract of chromosome 12 showed high levels of differentiation in our FST scan. This signal belonged to a single block of sequence, spanning 8 Mb (roughly half the length of the chromosome), containing over 300 annotated genes, and found at a frequency of almost 70% in the Arizona population.
Individuals that are heterozygous for this putative inversion or lacking it entirely cluster more closely with the Kansas population than other Arizona samples which are homozygous for it. When removing chromosome 12 from analyses, all Arizona samples cluster more closely with each other than with Kansas however, suggesting that the 2 populations are differentiated across the genome, but that the sheer number of variants associated with the putative inversion swamps other signals. These 2 observations combine to suggest that the inversion has been segregating in the population for some time or that selection on some variant(s) in the inversion has driven frequency changes at all linked loci.

In general, and especially in large populations, inversion frequencies are thought to imply something about their selective effects, with only beneficial variants reaching high frequency (Lande 1984, Kirkpatrick and Barton 2006). In other species in which naturally occurring inversions have been studied, this expectation has been borne out. Low frequency inversions (e.g. 8%) are associated with fitness tradeoffs (Lee et al. 2016) and more common inversions (~50%) may have little to no fitness costs (Knief et al. 2016), at least locally. Their absence elsewhere may suggest a lack of gene flow or that there are fitness costs in other environmental backgrounds. Indeed, linkage disequilibrium is higher within the bounds of the inverted region even in populations in which we did not detect the variant, suggesting a dampening of gene flow between populations here. From this perspective, the common chromosome 12 variant seems likely to be at least selectively neutral, if not beneficial, in Arizona. However, the large number of encompassed genes present a challenge for identifying selectively relevant loci from purely computational methods. Still, one gene of interest arose from our efforts to cross-check results with the older M. sexta reference (Kanost et al. 2016).

A likely candidate for local adaptation in the inversion: an apparent pseudogene

An FST scan of the older assembly with the same resequencing data revealed numerous outlier scaffolds anchored to chromosome 12 as expected. However, an additional similar peak was detected on a scaffold assigned to chromosome 13. Alternate allele frequencies on this peak matched those most commonly observed on a scaffold assigned to chromosome 13 (in the old annotation; XM_037442453.1 in the new annotation). In this regard, the single gene is one of many potential candidates for ecologically significant differentiation in the inversion; its misassembly is the main cause of our additional scrutiny and it may hold no special relevance for the Arizona population. Still, the patterns of variation are striking here. This gene carries almost no variation in North Carolina or Kansas but has a wealth of high frequency variants (61 SNPs in our sample) in the Arizona population, one of which is a predicted stop loss in the coding region. A large-effect SNP like this could easily break the protein function via translation of otherwise noncoding DNA. While inconclusive from molecular data alone, a relaxation in selective pressure at this gene would allow for pseudogenization and subsequent accumulation of numerous variants. And indeed, the high frequency of this stop-loss suggests that it cannot have an appreciable fitness cost, or it would not have persisted in the population. Naturally, this raises the question of what function this gene has that it could be lost without strong selective consequences. Or, alternatively, could a loss of function be adaptive under certain circumstances?

Based on our expression characterization, the Msex2.10493 gene is almost exclusively expressed in larval antennae, which are known to play a role in olfaction and food choice in M. sexta (de Boer and Hanson 1987). Olfaction genes are among the most commonly observed to pseudogenize in other species, ostensibly because the fitness consequences of their loss are rarely lethal (Nei et al. 2008; Prieto-Godino et al. 2016). Based on these observations, it is possible that Msex2.10493 is merely a broken olfaction gene that got caught in the massive inversion found in Arizona by chance. Intriguingly however, while most populations of M. sexta feed almost exclusively on plants in the family Solanaceae, larval Arizona hornworms have been recorded eating the unrelated devil’s claw (Martyniaceae: Proboscidea parviflora, Mechaber and Hildebrand 2000). It is tempting to make the connection between the apparent loss of function of a larval sensory gene and the exploitation of a new hostplant, as research on the silkmoth (B. mori) has shown that knockout of a single chemoreceptor is sufficient to turn naturally specialist larvae into generalists (Zhang et al. 2019); however, it will take further study to establish whether or not a similar situation is occurring in M. sexta.

Still, the placement of this apparent pseudogene within a massive block of elevated differentiation between populations and its potential relationship with hostplant ecology is suggestive of a locally adapting region. As genomic scans of natural variation become more common, so too does the observation that many species harbor regions of large-scale differentiation that distinguish individuals adapted to different environments (Fuller et al. 2019; Todesco et al. 2020; Mérot et al. 2021). In such instances, suppressed recombination, typically through inversions, keep locally beneficial alleles from being decoupled from each other, even in the face of gene flow from other populations. The frequencies of these large variants typically follow an environmental (or more directly a selective) cline (Kapun et al. 2016; Wellenreuther et al. 2017). We do not have the geographic resolution to detect any potential cline in the frequency of this inversion and pseudogene, but presence and absence data are at least consistent with a role in expanded hostplant usage.

The devil’s claw plant’s native range is limited to the desert southwest of North America (particularly: northern Mexico and California, Arizona, New Mexico, Utah, and Nevada in the United States; see Berry et al. 1981). As expected based on this range, we did not observe any M. sexta carrying the putative inversion in North Carolina or Kansas. With only 4 sampled individuals from Kansas, we cannot exclude the possibility that the Arizona inversion is present there as well, albeit at a much lower frequency, but to follow the logic of the devil’s claw hypothesis, there would be no local benefit for individuals carrying the pseudogene outside of the southwest. Indeed, in the absence of a novel hostplant with the correct set of complex phytochemistry, becoming less choosy could be maladaptive. Zhang et al. (2019) found that although chemoreceptor-knockout silkworms readily ate many food sources, they failed to finish development and pupation on all but the traditional hostplant. To reiterate, these inferences are highly speculative, but they create easily testable hypotheses. Laboratory studies of Arizonan M. sexta with the putative inversion and pseudogene will be needed to unpack the functional consequences of this striking variation.
A Z inversion in AZ: a role for sex-bias driving rearrangement

The single highest peak of differentiation in our genome-wide scan comes from the Z chromosome. This region, with roughly 1 Mb of extended differentiation and shared allele frequencies in Arizona also fits the pattern of a segregating inversion between populations. Moreover, the nonreference orientation is nearly fixed in the Southwest population, being homozygous (or present in hemizygous females) in all but one individual. As discussed above, an inversion is unlikely to reach such high frequency without some selective advantage.

Here, even without specific gene functional annotations, we have a reasonable hypothesis for what could drive this structural variant to near fixation. We found that the inversion contains a significant excess of genes expressed in both sexes (i.e. unbiased genes), compared to the rest of the Z chromosome, which is male biased (Mongue et al. 2020). As has been recently shown, unbiased genes are an important source of positive selection on the Z in M. sexta (Mongue et al. 2022). These genes are expressed in a haploid state in females (which have only one Z chromosome); thus, even recessive beneficial mutations are exposed to positive selection (Rice 1984; Charlesworth et al. 1987). The capture of positively selected variants of such genes in a nonrecombining region would ensure their co-transmission and seems a reasonable explanation for this inversion’s prevalence.

Moreover, these sex chromosome dynamics intersect with the population genetics of local adaptation in synergistic ways. Modeling suggests that the exposure of recessive alleles to selection should facilitate local adaptation and make the sex chromosomes hotspots for this phenomenon (Lasne et al. 2017). For similar reasons, sex-linked inversions are thought to sweep to fixation more easily than autosomal inversions (Connallon et al. 2018). As such, it is unsurprising to find the Z involved in short-term differentiation as well as long-term adaptation (Mongue et al. 2022) in this species.

Conclusions

We report that M. sexta from Arizona, Kansas, and North Carolina are genetically differentiated. The confirmation of regional differentiation is a valuable addition to research in both laboratory and field studies, as well as to development of management plans for this crop pest between regions. That said, we have yet to establish the minimum geographic scale at which M. sexta populations are structured (for instance, how much gene flow exists between adjacent states or even within a single state). This fine-scale sampling will be particularly important for agricultural centers on the applied side, and additional sampling will also provide more insight into the large-scale variation we report here.

At least 2 large inversions and a potential pseudogene differentiate Arizona from the Kansas and North Carolina populations. All of these variants are high frequency (>0.5) within Arizona but absent in the other populations. However, the nature of our sample collection limits the inferences we can make at present. For instance, the high frequency of the variants suggests they are not deleterious, but with only a single time-point of sampling in Arizona, it is hard to say whether or not these variants have reached an equilibrium frequency. Additionally, other studies have shown inversion polymorphisms to follow geographic or environmental clines (Anderson et al. 2005; Wellenreuther et al. 2017). In such cases, the inversions often hold locally adaptive alleles that are maintained in spatial polymorphism by differences in environmental factors, but for M. sexta, with each sampled state separated from the others by at least 1,500 km, it is impossible to say if these segregating variants also follow a cline in the region or to which climactic variables they may relate. Additional sampling of the Southwest will be required to assess how widespread the inversions are.

Likewise, controlled laboratory studies will be required to assess the fitness consequences of the segregating stop-loss in a larval antennal gene within the chromosome 12 inversion. In particular, the coincidence of a novel larval hostplant in Arizona and a large-effect mutation in a gene expressed in a larval chemosensory gene are suggestive but far from conclusive. The relationship between the 2 will require hostplant choice experiments for larvae with and without the stop-loss mutation.

Finally, it is worth noting that these dramatic differences between populations could not be observed in standard lab strains, which are all derived from the North Carolina population to our knowledge. On the other hand, without extensive lab studies and RNA-sequencing, we would not have the information to form functional hypotheses for these variants found in wild populations. Beyond the immediate utility of these data, the results we present here serve as a reminder of the efficacy of integrating laboratory and field research within the same study system.

Data availability

All newly generated sequencing data are available with the following accessions: SRP144217, PRJNA639154 on NCBI. Accessions for RNAseq used in functional inference can be found in Cao and Jiang (2017). Custom scripts for analysis can be found at github.com/amongue/Manduca_Demography.git. New assembly chromosome assignments can be found in the Supplementary Files. Supplemental material is available at G3 online.

Acknowledgments

The authors wish to acknowledge Wesley Mason and Michael Hulet and the rest of the Information and Telecommunication Technology Center (ITTC) staff at the University of Kansas for their support with our high-performance computing. The authors also thank the University of Kansas’s Entomology Endowment for additional funding support in sequencing. Thanks to Paul Hime for guidance on population structure analysis. Thank you to Clyde Sorenson, Judie Bronstein, and Coggy Davidowitz for initial and continued support in investigating population genomics of M. sexta. Kelly Dexter and Rebecca Messcher helped provide the samples from University of Florida’s McGuire Center collection. Thanks also to the members of Ashworth 2.02 for continued patience in the discussion of the material of this manuscript.

Funding

This project was partially funded by National Science Foundation Doctoral Dissertation Improvement Grant to AJM (Division of Environmental Biology #1701931) and DEB #1557007 and Integrative Organismal Systems #1920895 to AYK.

Conflicts of interest

The authors declare no conflicts of interests.
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Communicating editor: A. Whitehead