

1 Corresponding Author:
2 Dr. Megan L. Fritz
3 112 Derieux Place
4 1549 Thomas Hall
5 Department of Entomology
6 North Carolina State University
7 Raleigh, NC 27695
8 mlfritz@ncsu.edu
9 (919) 515-1651
10

11 **Application of a dense genetic map for assessment of genomic responses to selection and**
12 **inbreeding in *Heliothis virescens*.**

13

14 Megan L. Fritz^{1,2,§}, Sandra Paa¹, Jennifer Baltzegar², and Fred Gould^{1,2}

15 ¹Department of Entomology, North Carolina State University, Raleigh, NC, USA

16 ²Department of Biological Sciences, Program in Genetics, North Carolina State University, Raleigh,
17 NC, USA

18 [§]Corresponding Author

19 **Abstract**

20 Adaptation of pest species to laboratory conditions and selection for resistance to toxins in the
21 laboratory are expected to cause inbreeding and genetic bottlenecks that reduce genetic variation.
22 *Heliothis virescens*, a major cotton pest, has been colonized in the laboratory many times, and a few
23 laboratory colonies have been selected for Bt resistance. We developed 350 bp Double-Digest
24 Restriction-site Associated DNA-sequencing (ddRAD-seq) molecular markers to examine and compare
25 changes in genetic variation associated with laboratory adaptation, artificial selection, and inbreeding in
26 this non-model insect species. We found that allelic and nucleotide diversity declined dramatically in
27 laboratory-reared *H. virescens* as compared with field-collected populations. The declines were
28 primarily due to the loss of low frequency alleles present in field-collected *H. virescens*. A further,
29 albeit modest decline in genetic diversity was observed in a Bt-selected population. The greatest
30 decline was seen in *H. virescens* that were sib-mated for 10 generations, where more than 80% of loci
31 were fixed for a single allele. To determine which regions of the genome were resistant to fixation in
32 our sib-mated line, we generated a dense intraspecific linkage map containing 3 PCR-based, and 659
33 ddRAD-seq markers. Markers that retained polymorphism were observed in small clusters spread over
34 multiple linkage groups, but this clustering was not statistically significant. Here, we confirmed and
35 extended the general expectations for reduced genetic diversity in laboratory colonies, provided tools
36 for further genomic analyses, and produced highly homozygous genomic DNA for future whole
37 genome sequencing of *H. virescens*.

38 **Keywords:** *Heliothis virescens*, genetic variation, colonization, inbreeding, ddRAD-seq, linkage map

39 **Introduction**

40 Laboratory-reared insect colonies are important resources for many types of entomological
41 experiments. They are used to quantify physiological or behavioral differences between insect
42 populations or species (Dekker *et al.*, 2006; Dobzhansky & Spassky, 1954; Fritz *et al.*, 2015; Groot *et*
43 *al.*, 2005; Shaw *et al.*, 2000; Sokolowski, 1980; Tomaru *et al.*, 2000), identify the genetic architecture
44 of insect traits (Gahan *et al.*, 2010, Mackay *et al.*, 2012, Oppenheim *et al.*, 2012), develop insect
45 populations that express desirable traits (Collins, 1984; Goldman *et al.*, 1986; Gould *et al.*, 1995; Hoy,
46 1990; Pradeep *et al.*, 2005), and generate genetically modified species as a means of pest control (de
47 Valdez *et al.*, 2011). A major concern for researchers maintaining insect colonies is the degree to which
48 adaptation to the laboratory environment affects insect genotypic, and thereby phenotypic diversity
49 (Boller, 1972; Huettel, 1976).

50 The phenotypic consequences of adaptation to the laboratory depend upon the trait of interest,
51 and range from undetectable to severe (Baeshen *et al.*, 2014; Fox *et al.*, 2007; Gerloff *et al.*, 2003;
52 Raulston, 1975; Roush, 1986). Observed phenotypic changes can be attributed to inadvertent selection
53 for traits that are favorable in the laboratory environment (Roush, 1986), inbreeding depression (*i.e.*
54 reduction in fitness caused by matings between related individuals; reviewed in Charlesworth & Willis,
55 2009; Mackauer, 1976), or the interaction of the two. Indeed, the selection that occurs during colony
56 establishment creates conditions conducive to inbreeding (Roush, 1986). Families with higher fitness
57 under laboratory conditions contribute disproportionately to the reproductive pool, thereby increasing
58 the probability of matings between related individuals. Where selection is very strong, as in the
59 production of an insecticide resistant colony, measures must often be taken to minimize the effects of
60 inbreeding and thereby inbreeding depression (Gould *et al.*, 1995). Overall, the expectation is that the
61 selection and inbreeding that takes place during insect colonization results in an overall loss of genetic

62 diversity (Munstermann, 1994), and concomitant genome-wide increase in homozygosity (reviewed in
63 Etzel & Legner, 1999).

64 Previous studies that have examined genetic differences between field-collected, laboratory-
65 adapted, and inbred populations of non-model insects have primarily focused on Dipteran species and
66 were limited to small numbers of molecular markers (Mukhopadhyay *et al.*, 1997; Munstermann, 1994;
67 Norris *et al.*, 2001). Such small numbers of markers allow for estimation of the genome-wide average
68 change in genetic variability across populations, but cannot be used to examine fine-scale patterns of
69 genomic change. Examination of these patterns allows for identification of where and how genetic
70 variation, the raw material necessary for environmental adaptation, is maintained (Dobzhansky &
71 Spassky, 1954). The relatively recent development of high-throughput sequencing combined with
72 reduced-representation DNA library preparation techniques allows for the discovery of hundreds to
73 thousands of new molecular markers, even in species for which genomic data are absent (Davey *et al.*,
74 2012). Here we used Double-Digest Restriction-Site Associated DNA Sequencing (ddRAD-seq;
75 Peterson *et al.*, 2012), one type of reduced representation library preparation, for *de novo* construction
76 of molecular markers in the non-model species, *Heliothis virescens*.

77 The tobacco budworm, *H. virescens*, is an historically important pest of cotton throughout much
78 of the Southeastern United States (Blanco, 2012). This non-model Lepidopteran species has been
79 colonized a number of times for investigations of mating and host-selection behaviors (Sheck & Gould,
80 1995; Sheck & Gould, 1996; Sheck *et al.*, 2006), as well as detecting the underlying genetic basis for
81 insecticide resistance (Gahan *et al.*, 2001; Gahan *et al.*, 2010; Taylor *et al.*, 1993). We used our newly
82 developed ddRAD-seq markers to examine and compare the effects of colonization, selection, and sib-
83 mating on *H. virescens* genome-wide measures of genetic diversity. To examine fine-scale patterns of
84 change in genetic diversity, we also used our ddRAD-seq markers to generate a dense intraspecific

85 genetic map for *H. virescens*. This map consists of 659 high quality 350-bp markers which will serve
86 as an important genomic resource to the entomological community.

87 Overall, our research aims to:

- 88 1. Quantify overall patterns of change in genomic diversity across field-collected, laboratory-
89 reared (non-selected), Bt-selected, and sib-mated *H. virescens*.
- 90 2. Determine whether the observed degree of inbreeding in our sib-mated *H. virescens* calculated
91 from our ddRAD-seq genotypic data matched theoretical expectations (Falconer & Mackay
92 1996).
- 93 3. Use ddRAD-enabled linkage mapping to determine if specific genomic regions were resistant to
94 fixation, even under intense inbreeding, by comparing genotypic data from long-term colony
95 and sib-mated lines.

96

97 **Results**

98 We sequenced 204 *H. virescens* individuals from a total of 6 populations that were used in a
99 population-level analysis of genomic change associated with laboratory colonization, artificial selection
100 and inbreeding. These populations were comprised of 2 field-collected (LA, TX populations collected
101 in 2012), 2 laboratory-reared (BENZ, YDK), and 1 Bt-selected population (YHD2), as well as
102 specimens from a single inbred family following 10 generations of full-sibling mating (see Table 1 for
103 information on population history, sample sizes, and read counts). Three of these populations, YDK,
104 YHD2, and the inbred line were founded from a collection in Yadkin County, NC, in 1988 (Gould *et*
105 *al.*, 1995), but were thereafter subjected to different rearing conditions, allowing us to make
106 comparisons of population-genomic change within the same genetic background. In addition, 99
107 individuals (1 BENZ parent, 1 BENZ-YHD2 hybrid parent and 97 progeny) were sequenced for

108 linkage analysis. This produced a total of 105,487,499 Illumina MiSeq reads (38,221,995 and
109 67,265,504 for linkage- and population-level analyses, respectively) that passed quality filters (data
110 available upon request).

111 *Genomic diversity among H. virescens populations*

112 On average, 338,892 sequencing reads (*s.d.* = 113,397) were produced per individual, and the
113 variation in read count was spread uniformly across populations (Supplementary Figure 1). Ninety-
114 four percent of individuals had read counts between 90,000 and 688,000, and these were fed into the
115 Stacks pipeline (Catchen *et al.*, 2011; 2013) for *de novo* locus construction (Supplementary Figure 1).
116 Loci constructed by Stacks had an average of 6× depth of coverage per individual. In total, 4,281
117 polymorphic 350-bp ddRAD-seq markers (hereafter loci) were detected in at least one individual per
118 population across all populations. Two well-documented challenges commonly encountered when
119 working with moderate coverage reduced representation library data like ours are: 1) uneven
120 distribution of missing data across sets of loci (Davey *et al.*, 2012; Xu *et al.*, 2014), and 2) under-
121 sampling of heterozygotes (Li *et al.*, 2009; Nielsen *et al.*, 2011). Both reduce confidence in final
122 genotypes called by genotyping-by-sequencing SNP calling algorithms, including the algorithm used in
123 Stacks. To overcome these challenges, we examined several subsets of these 4,281 polymorphic loci
124 for our downstream population genomic analyses. These subsets contained between 125-1231 loci, and
125 were chosen based upon the overall proportion missing genotype calls in the subset. The smallest
126 subset consisted of loci for which over 75% of individuals per population had genotypic data present
127 and were therefore likely sequenced to greatest depth of coverage. Each larger subset allowed
128 additional loci at the expense of coverage (*i.e.* more missing genotypic data were allowed;
129 Supplementary Table 1). By using multiple datasets, we were able to examine whether the presence of
130 missing genotype calls influenced overall estimates of genomic diversity across populations.

131 We first examined all subsets of loci and determined the mean and maximum numbers of alleles
132 per locus (Supplementary Table 1). In our case, alleles were not analogous to SNPs, but rather the
133 accumulation of SNPs per 350 bp locus per individual. For the total sequenced population ($n = 192$
134 total *H. virescens*), the mean numbers of unique alleles detected per locus ranged from 29 to 34
135 depending upon the number of loci included in the analysis. As more loci were included, the average
136 number of unique alleles detected per locus decreased. However, the maximum number of unique
137 alleles detected in the total population increased from 86 in the smallest subset of loci to 94 in the 3
138 larger subsets. We also examined the proportion of loci that were fixed (*i.e.* only a single allele
139 present) across populations. Across subsets, few loci were fixed for a single allele in laboratory-reared
140 (5.6-10.9%), Bt-selected (5.3-7.3%) and field-collected (0-2.4%) populations (Supplementary Table 1).
141 Yet over 80% of loci were fixed in the inbred line following 10 generations of sib-mating
142 (Supplementary Table 1). Of the 125 loci with the fewest missing genotype calls, 86% were fixed in
143 the inbred line. Expanding the number of loci to include those with more missing genotypes ($n = 378,$
144 573, 1231) reduced the percentage of fixed loci in the inbred line by up to 5% (Supplementary Table 1).

145 We then determined the mean number of unique alleles present per locus for each subset of loci
146 (Supplementary Figure 2). In general, we found no within population differences in the mean numbers
147 of unique alleles detected among subsets of loci, and therefore we used a single, conservatively chosen
148 subset of loci ($n = 378$) where at least 10 individuals were genotyped per population per locus for
149 further analysis. The mean numbers of unique alleles per locus were 2.1 for the inbred line, 5.3 for the
150 Bt-selected population, 5.4 and 4.4 for the non-selected, laboratory-reared populations (YDK and
151 BENZ, respectively), and 18.4 and 17 for the field-collected populations (LA and TX, respectively).
152 However, our sample sizes (*i.e.* numbers of individuals sequenced; see Table 1) differed for each
153 population, and it was unclear whether differences between the aforementioned means were caused by

154 sample size or population-level differences. For example, it is to be expected that as sample size
155 increases there will be an increase in the probability of sampling additional, likely rare, alleles.
156 Therefore, we randomly sub-sampled pools of alleles 6, 12, 18, and 24 times without replacement for
157 each population. This allowed us to hold sample sizes constant across populations, and infer whether
158 the mean numbers of unique alleles truly differed by population. As expected, we found that increasing
159 the total number of alleles sampled led to an increase in the mean numbers of unique alleles per locus
160 for all but the inbred line (Figure 1). Yet we also found strong population-level differences.
161 Regardless of the number of alleles sampled, field populations always exhibited the greatest allelic
162 diversity, followed by selected and non-selected colony populations. The lowest allelic diversity was
163 observed in the inbred line.

164 When 18 alleles were randomly sampled per population per locus, we detected an average of
165 just over 1 unique allele per locus in the inbred line, indicating that most loci were fixed for a single
166 allele. For the inbred line, 52 of the 378 loci did not reach fixation. Of these, forty-seven had 2 unique
167 alleles, four had 3 unique alleles, and one had 4 unique alleles when 18 were randomly sampled. On
168 average, Bt-selected and non-selected colony strains each had *ca.* 3 unique alleles per locus, and field-
169 collected populations had *ca.* 9 unique alleles per locus (Figure 1). The majority of unique alleles
170 present in the field-collected populations (70.3% and 68.7% for LA and TX populations, respectively)
171 were observed only once (of 18 alleles; Figure 2). Such low frequency alleles were less common in the
172 Bt-selected and non-selected laboratory populations (35.1% and 21.5%, respectively) and rare (0.7%)
173 in the sib-mated inbred line (Figure 2).

174 We used the same conservative subset of 378 loci, and calculated sample-size corrected S_K
175 (Charlesworth & Charlesworth, 2010), and π (Nei 1978). These two measures are complementary: π is
176 calculated as the proportion of nucleotides that differ per two randomly chosen DNA sequences,

177 averaged across all pairwise comparisons per marker per population, and S_K is calculated as the number
178 of unique single nucleotide variants in a population at a single locus. When averaged across all
179 markers ($n = 378$), the number of variant sites (S_K) per 350 bp marker was 0.15 for the inbred line, and
180 the maximum S_K was 2.95. Bt-selected and non-selected laboratory populations had, on average, just
181 over 1 nucleotide variant per 350 bp locus, with a maximum of *ca.* 6. Field-collected populations had
182 the greatest number of variant sites per 350 bp locus, where the genome-wide average was just over 5
183 nucleotide variants per locus, with a maximum of *ca.* 15. Similar trends were observed for genome-
184 wide and maximum nucleotide diversity (π) values. Relative to the laboratory-reared populations,
185 genome-wide estimates of π were nearly an order of magnitude lower for the inbred line. The genome-
186 wide π estimate for laboratory-reared populations ranged from 4.0×10^{-3} (Bt-selected population) to
187 6.7×10^{-3} (non-selected, YDK population), and 6.2×10^{-4} for the inbred line. Field-collected populations
188 exhibited genome-wide π estimates of 9.4×10^{-3} and 9.2×10^{-3} for the LA and TX populations,
189 respectively. Genome-wide and maximum π and S_K estimates, along with their corresponding 95%
190 non-parametric bootstrapped confidence intervals ($N = 5000$) are reported in Table 2.

191 To further quantify and compare genetic diversity by population, we also examined mean
192 observed heterozygosity in the total population ($n = 192$ total *H. virescens*), as well as within each sub-
193 population. Observed heterozygosity for the total population ($n = 192$) was 0.27 when averaged across
194 the 378 loci, and considerable variation in heterozygosity existed between populations. Mean observed
195 heterozygosities ranged from 0.06 in the inbred line to 0.40-0.46 in field-collected populations. For
196 laboratory-reared populations, mean observed heterozygosity estimates were intermediate to those of
197 the inbred line and field-collected populations, and ranged from 0.15 in the Bt-selected population to
198 0.22-0.25 in the non-selected laboratory populations (Table 2).

199

200 *Genomic divergence among populations*

201 To examine the degree to which the above genetic diversity could be attributed to between
202 population differences, we calculated pairwise estimates of F_{ST} (Table 3) according to Weir and
203 Cockerham (1984). Despite the *ca.* 400 km distance between collection locations for the field
204 populations, very little (0.4%) of the genetic diversity observed in these populations could be attributed
205 to differences between populations. When field populations were compared with non-selected
206 laboratory-reared populations, 16-25% of the genetic variation could be attributed to differences
207 between populations. Additional inbreeding and selection further exacerbated these differences,
208 increasing the percentage of variation attributable to between population differences to over 30%. Of
209 particular interest was the comparison between the ancestral non-selected YDK population, with the
210 more derived Bt-selected (YHD2) and inbred populations. Despite their shared ancestry, a comparison
211 of YDK to YHD2 and the inbred line revealed that 28% and 33% of the existing genetic variation could
212 be attributed to between population differences, respectively.

213

214 *Inbreeding among laboratory-reared H. virescens populations*

215 To determine whether the observed degree of heterozygosity in our inbred line was consistent
216 with that which would be expected following 10 generations of sib-mating, we compared the
217 inbreeding coefficient F , as calculated according to pedigree- (Falconer & Mackay, 1996) and DNA
218 marker-based information (Keller & Waller, 2002; Kim *et al.*, 2007). The expected inbreeding
219 coefficient (F_e), following 10 generations of sib-mating was 0.89. This expected value fell within the
220 bootstrapped 95% confidence intervals for marker-based inbreeding coefficients (F_{IT}) calculated from
221 all subsets of markers. This indicated that there was no significant difference between the expected
222 inbreeding coefficient and the observed inbreeding coefficient calculated using ddRAD-seq marker

223 data. The genome-wide F_{IT} values (95% CIs) were 0.92 (0.88, 0.96), 0.89 (0.86, 0.92), 0.89 (0.87,
224 0.92), and 0.88 (0.86, 0.89), for the inbred line as calculated from 125, 378, 573, and 1231 ddRAD-seq
225 loci, respectively.

226

227 *Linkage mapping*

228 Few genomic resources are available for *H. virescens*. Therefore, we determined the genomic
229 location of loci which were resistant to fixation by generating a dense linkage map. The map was
230 produced via ddRAD-sequencing of the parents and progeny from a male informative cross (reviewed
231 in Baxter *et al.*, 2009). We generated an average of 381,096 sequencing reads (*s.d.* 165,334) per
232 progeny, as well as 493,537 and 762,171 reads per the male and female parents, respectively
233 (Supplementary Figure 3). From this, we produced a linkage map comprised of 659 informative
234 ddRAD-seq loci, plus 3 partial gene sequences of *ABCC2*, *HevCaLP*, and *Desat1*. Adding these partial
235 gene sequences to our linkage map, all with known locations in the *B. mori* genome allowed us to
236 validate marker groupings for our linkage map. All informative ddRAD-seq loci were grouped into 33
237 linkage groups, two more than the expected 30 *H. virescens* autosomes, and one segregating Z
238 chromosome from the hybrid male parent used in our cross. Linkage groups ranged in size from 7cM
239 to 110cM (Figure 3), and yielded a total map length of 1919.5 cM. On average, there were 20
240 ddRAD-seq loci per linkage group, and the average spacing was one locus per 3.5 cM. The smallest
241 and largest linkage groups contained 3 and 53 loci, respectively. The *HevCaLP*, *Desat1*, and *ABCC2*
242 genes were grouped with linkage groups 15, 16, and 22, respectively. These linkage groups
243 corresponded to *B. mori* chromosomes 6, 23, and 15 (Table 4), where these candidate genes are known
244 to reside (Gahan *et al.*, 2001; Gahan *et al.*, 2010; Mita *et al.*, 2004).

245 In total, 99 of the 659 mapped ddRAD-seq loci could be aligned uniquely to a single locus in

246 the *B. mori* genome. Twenty-two linkage groups contained ddRAD-seq loci that could be aligned to a
247 single *B. mori* chromosome (Table 4), while 8 linkage groups did not contain any that could be aligned.
248 Linkage groups 19, 25, and 33 contained ddRAD-seq loci that aligned uniquely to more than one *B.*
249 *mori* chromosome. This was unlikely caused by spurious associations between ddRAD-seq loci;
250 increasing the LOD score to 8 failed to break up associations for those three linkage groups.

251

252 *Identification of genomic regions resistant to fixation*

253 To determine where genetic diversity was being maintained in the genome, we examined
254 observed heterozygosity and nucleotide diversity at the 659 mapped ddRAD-seq loci for one field-
255 collected (LA), the Bt-selected (YHD2), one non-selected (YDK), and inbred lines. Of these 659
256 mapped loci, 302 (46%) were previously included in our population-level analyses, and 357 had not
257 been previously analyzed. For each mapped locus, observed heterozygosity and nucleotide diversity
258 (π) values were only calculated if at least 3 individuals were genotyped per population. Therefore only
259 441 loci were examined for the inbred line, 658 loci were examined for the non-selected (YDK)
260 population, 659 loci were examined for the Bt-selected population, and 546 loci were examined for the
261 field-collected population. In total, 13% ($n = 60$) of mapped loci retained polymorphism in the inbred
262 line, whereas 98% ($n = 645$), 86% ($n = 583$), and 99% ($n = 543$) of mapped loci retained
263 polymorphism in the Bt-selected, non-selected (YDK), and field-collected (LA) populations,
264 respectively. For each of these populations, levels of observed heterozygosity and nucleotide diversity
265 across the genome are compared in Figure 4.

266 Within the inbred line, mapped loci that retained polymorphism were spread over 16 linkage
267 groups (Figure 4), and often appeared in clusters within a linkage group (Supplementary Figure 4).
268 Linkage groups 7 and 11 had particularly high numbers of polymorphic markers relative to other

269 linkage groups. Therefore, we examined whether clustering of polymorphic markers in our inbred line
270 was greater than could be expected due to random chance. Such clustering may point to the presence
271 of large chromosomal inversions as observed in other insect species (Turissini et al. 2014). A replicated
272 G-test of independence (Sokal and Rohlf, 1995) demonstrated that 8 linkage groups contained
273 significantly more polymorphic loci than would be expected following 10 generations of inbreeding
274 (Supplementary Table 2). Yet based upon the non-significant heterogeneity g-value ($G = 23.87$, $df =$
275 23 , p -value = 0.41) we could not reject the null hypothesis that the distribution of polymorphic loci was
276 homogeneous across linkage groups, and the clustering observed within some linkage groups did not
277 significantly differ from that which could be observed by chance.

278 We also examined whether balancing selection, perhaps due to balanced lethal systems, may be
279 responsible for the residual polymorphism observed in the inbred line. To this end, we calculated and
280 compared Tajima's D values at each polymorphic locus for the inbred line and its ancestral population
281 (YDK). Tajima's D is a statistical test which identifies departures from the neutral model of molecular
282 evolution. A positive Tajima's D value at a locus indicates an excess of intermediate frequency alleles,
283 thereby signifying either a recent population contraction, or balancing selection at that locus. We
284 reasoned that positive Tajima's D values in the inbred line would be much less likely to signify a recent
285 population contraction, and therefore more likely to indicate balancing selection, if these same regions
286 were also significant in their ancestral population (YDK). Indeed, YDK had not undergone any
287 obvious population contractions around the time of this separation 10 generations prior. Therefore, we
288 examined which, if any, of these polymorphic loci in the inbred line shared strongly positive Tajima's D
289 values with YDK. When we calculated Tajima's D values for the 60 polymorphic sites remaining in the
290 inbred line, 21 were significantly positive ($\alpha = 0.05$). Following a Benjamini-Hochberg correction for
291 multiple comparisons (Benjamini and Hochberg, 1995), only two loci remained statistically significant.

292 Furthermore, neither of these two polymorphic loci from the inbred line overlapped with loci showing
293 significantly positive Tajima's D values in YDK. Thus we found little evidence of ongoing balancing
294 selection in our inbred line.

295

296 **Discussion**

297 Here, we examined the degree to which colonization, artificial selection, and intense inbreeding
298 influence genome-wide and fine-scale patterns of diversity. In the absence of a publicly available *H.*
299 *virescens* reference genome, we used ddRAD-seq *de novo* locus construction to identify multiple
300 subsets of polymorphic loci ranging in size from 125 to 1231 markers. Genome-wide measures of
301 allelic diversity, F_{IT} values, and the degree of homozygosity were either unaffected (Supplementary
302 Figure 2), or minimally affected (Supplementary Table 1) by inclusion of markers with high levels of
303 missing genotypic data. Therefore, any biased genotype calls made by the Stacks SNP calling
304 algorithm due to our moderate depth of sequencing coverage had little impact on our overall genome-
305 wide estimates of diversity. Our results demonstrate that moderate coverage ddRAD-seq data can be
306 used with confidence when conducting population genomic comparisons of genome-wide means.

307 We observed a precipitous decline in nucleotide and allelic diversity following long-term
308 laboratory colonization, selection, and inbreeding for *H. virescens*. Despite the decline in genomic
309 diversity for non-selected and Bt-selected laboratory-reared populations, fewer than 10% of loci were
310 fixed. While our Bt-selected population did not retain the level of genetic diversity that their ancestral
311 (YDK) laboratory-reared population did, they consistently had higher measures of genomic diversity
312 than did the non-selected (BENZ) population. Retention of higher levels of polymorphism in our Bt-
313 selected line was likely due to the measures taken during its generation to ensure genomic diversity was
314 maintained in the face of strong selection (Gould, 1995). Alternatively, strong genetic bottlenecks in

315 the non-selected (BENZ) population prior to their use in our study could explain why our Bt-selected
316 line was more genetically diverse than the BENZ non-selected line. Overall, differences among Bt-
317 selected and non-selected laboratory-reared populations were modest; when 18 total alleles were
318 sampled, laboratory-reared populations retained *ca.* 3 alleles per 350-bp locus relative to 9 alleles per
319 350-bp locus present in field-collected *H. virescens*. However, few low frequency alleles remained in
320 the laboratory-reared populations relative to the field-collected populations (Figure 2), which has been
321 observed elsewhere (Munstermann, 1994). For this reason, laboratory-reared populations are generally
322 considered inbred (Roush, 1986). In the case of *H. virescens*, our results clearly show that a great deal
323 of genomic diversity is retained, even following decades in colony. Few genome assembly algorithms
324 accommodate polymorphism well (Kajitani *et al.*, 2014), and it is clear that the reductions in
325 heterozygosity in our inbred line will be useful for production of a high quality *H. virescens* reference
326 genome assembly.

327 To determine the degree to which the genomic diversity detected above could be attributed to
328 differences between populations, we calculated pairwise F_{ST} values according to Weir and Cockerham
329 (Table 3). An F_{ST} value of 0.0004 demonstrated that most of the existing genetic diversity occurred
330 within, as opposed to between, these two field-collected populations of *H. virescens*. This result is
331 similar to that of Groot *et al.* (2011), which described over 98% of the genetic variation detected across
332 North American *H. virescens* to be found within populations. Both studies suggest that extensive gene
333 flow occurs naturally among geographically disparate *H. virescens* populations. Although we did not
334 sample field-collected populations from Yadkin County, NC, nor from Stoneville, MS, the original
335 collection sites of our laboratory-adapted YDK and BENZ populations, our results along with those of
336 Groot *et al.* (2011), suggest that genomic divergence among all four sites would have been low. To
337 examine how decades of laboratory rearing could influence the structure of genomic diversity in

338 laboratory-adapted populations, we compared our YDK and BENZ populations to our field-collected
339 populations. As would be expected, we observed an increase (16-25%) in the percentage of genetic
340 variation that could be attributed to between population differences. This is likely due to the purging
341 of rare alleles (as shown in Figure 2) and the fixation of others as caused by the random process of
342 genetic drift. Further manipulations, like Bt-selection and full-sibling mating increased that percentage
343 of between population genetic variation to well over 30% relative to the field-collected populations.
344 Additional comparisons between ancestral YDK and the more derived Bt-selected and inbred
345 populations further underscore the affects that these population-level manipulations can have on insect
346 colonies in a laboratory setting. Although they were derived from YDK, following Bt-selection and
347 over 2 decades of separation in the laboratory, 28% of the total genetic variability could be attributable
348 to between-population differences. Likewise, 10 generations of full-sibling mating resulted in over
349 32% of the total genetic variability existing between YDK and the inbred line.

350 Following 10 generations of inbreeding, over 80% of markers went to fixation in our sib-mated
351 *H. virescens* population. Indeed, our inbreeding coefficient F_{IT} , as observed from our ddRAD-seq data,
352 met theoretical expectations for all subsets of loci. Our *H. virescens* laboratory population was more
353 amenable to inbreeding than other insect species (Munstermann, 1994; Rumball *et al.*, 1994; Turissini
354 *et al.*, 2014; You *et al.*, 2013), despite their relatively high levels of genomic diversity (Figure 1) and
355 genetic load (Supplementary Figure 5). Higher than expected allelic diversity has been observed in
356 several other insect species following experimental inbreeding attempts (Munstermann, 1994; Rumball
357 *et al.*, 1994; Turissini *et al.*, 2014; You *et al.*, 2013). As one example, only 57% of the *An. gambiae*
358 genome went to fixation, as observed according to SNP markers, following 10 generations of
359 inbreeding (Turissini *et al.*, 2014). Observed differences between our *H. virescens* population and
360 other insects could be species specific, but is more likely related to the proportion of the genome

361 containing balanced lethal systems (Falconer & Mackay, 1996).

362 To determine where heterozygosity was being maintained in our inbred line, we developed a
363 high density genetic linkage map for *H. virescens*. Our map contained 659 newly developed 350 bp
364 ddRAD-seq markers that are long enough for future primer design and direct sequencing. This map
365 represents a new tool for an historically important pest species that lacks genomic resources. No
366 ddRAD-seq markers remained unlinked following mapping, which indirectly speaks to the quality of
367 our linkage map. However, the number of groups in our linkage map was 2 more than expected
368 ($n = 31$). This is likely due to the relatively small mapping population size used in this work; other
369 mapping studies that analyzed segregating populations of a similar size have also reported genetic maps
370 with excess numbers of linkage groups (Pootakham *et al.*, 2015; Singh *et al.*, 2009). Additional
371 explanations for the disparity between our observed and expected number of linkage groups include the
372 uneven distribution of markers over the chromosomes (Paterson, 1996), or recombination ‘hotspots’,
373 which make it difficult to reduce the number of linkage groups to 31.

374 When we applied our linkage map to examine fine-scale patterns of genomic change following
375 10 generations of sib-mating, we found that several linkage groups seemed to contain clusters of loci
376 that retained polymorphism (Figure 4). However, results from a replicated G-test of independence
377 demonstrated that clustering of these polymorphic loci was not significantly different from what could
378 be expected due to chance. This does not preclude the possibility that chromosomal inversions exist
379 among populations of *H. virescens*. Rather it suggests that there were no large inversions responsible
380 for maintaining polymorphisms in our inbred line.

381 Furthermore, only 2 polymorphic loci had significantly positive Tajima's D values, and neither
382 overlapped with those loci that had high D values in YDK. This made it impossible to rule out
383 population contraction as a contributor to the positive Tajima's D values. Unlike the findings for other

384 insects (Mackay et al. 2012, Turissini et al. 2014), the combined results of our F_{IT} test, replicated G-
385 test, and Tajima's D tests suggest that neither balanced lethals nor large chromosomal inversions appear
386 to play a major role in retention of polymorphism for this particular *H. virescens* inbred line.

387

388 **Conclusions**

389 This work serves as one of the most thorough attempts to quantify the effects of genomic
390 responses to selection and inbreeding in a non-model insect species. We demonstrated that laboratory-
391 reared *H. virescens* have reduced allelic and nucleotide diversity relative to field-collected populations,
392 and that inbreeding further diminishes genetic diversity. Although we identified several loci that did
393 not go to fixation in *H. virescens* following 10 generations of inbreeding, our ddRAD-seq marker-based
394 F_{IT} values met theoretical expectations. This work demonstrates the difficulty involved in producing
395 fully homozygous insect strains, which are currently critical to producing high-quality, complete
396 reference genomes.

397

398 **Methods**

399 *Field-collected H. virescens*

400 Adult male moths were collected from Bossier Parish, Louisiana, and College Station, Texas
401 using pheromone-baited live traps. Collections took place in LA from May through September, 2012,
402 and in TX from May through October, 2012. Moths from each collection date were immediately
403 placed in bottles of 95% ethanol for long-term storage. All bottles were held at -20 °C until DNA
404 isolations took place.

405

406 *H.virescens colonies*

407 *H. virescens* were collected from Yadkin County, NC in 1988 (Gould *et al.*, 1995). This original
408 population founded two of the colonies used in this study, each of which had been reared in the
409 laboratory for *ca.* 290 generations. YHD2 was selected for high levels of Bt resistance for 4 years (up
410 to 48 generations) on MVP- treated (0.864 mg/mL diet; Mycogen, San Diego, CA) corn-soy diet
411 (Gould *et al.*, 1995), whereas a non-selected population (YDK) was reared on corn-soy diet alone. A
412 third population (BENZ) originating from Stoneville, MS, was acquired from Benz Research
413 Incorporated (Carlisle, PA) and had been reared in the laboratory for over 10 years (120 generations).
414 BENZ *H. virescens* were acquired in their pupal stage, and newly eclosed adults were used for
415 population-level comparisons. To produce an inbred population, single pair matings (SPMs) were set
416 up between YDK siblings for 10 generations. An initial 37 SPMs were used to establish 29 lineages in
417 filial generation one (8 single pair matings did not produce progeny). When SPMs failed to produce
418 offspring, likely due to inbreeding depression, surviving lineages were expanded (Supplementary
419 Figure 5). This was done to extend inbreeding for as many generations as possible, thus promoting as
420 complete a reduction in heterozygosity as possible. Adult males from each laboratory-reared
421 population were killed by freezing (-20 °C), and stored at -80 °C until DNA isolation.

422

423 *Mapping cross*

424 A non-selected female from the BENZ population was crossed to a Bt-selected (YHD2) male in
425 a single pair mating. One hybrid male offspring was then back-crossed to a Bt-selected (YHD2)
426 female, and their progeny were reared to adulthood on untreated corn-soy diet according to Joyner and
427 Gould (1985). Of the 120 progeny, 97 reached adulthood. Parents and their 97 adult progeny were
428 killed by freezing and stored until DNA isolation as described above.

429

430 *Genomic DNA library preparation*

431 All DNA was isolated from the adult thorax using a Qiagen Dneasy Blood and Tissue Kit
432 (Qiagen, Inc., Valencia, CA, U.S.A.). Genomic DNA samples were prepared for Illumina sequencing
433 according to the Poland *et al.*, (2012) protocol with minor modifications. Two-hundred ng of DNA per
434 individual were digested with EcoRI and MspI. For each individual, the overhang sites were ligated to
435 standard Truseq Universal adapters (Illumina, Inc. San Diego, CA). Adapters ligated to EcoRI
436 overhang sites contained one of 48 unique barcodes (Elshire *et al.*, 2011; Supplementary Table 3).
437 DNA fragments from each individual were assigned a unique barcode, and individuals were combined
438 into pools of no more than 48 individuals. A Pippin Prep (Sage Science, Inc., Beverly, MA) was used
439 to select adapter-ligated DNA fragments ranging from 450-650 bp from each pool. Size-selected DNA
440 fragments were amplified in a Peltier PTC200 thermalcycler (here and throughout) using Illumina
441 primers (Supplementary Table 4) under the following conditions: 72 °C for 5min, 18 cycles of 98 °C
442 for 30 sec, 65 °C for 20 sec, 72 °C for 30 sec, followed by 72 °C for 5 min. For each pool, 1 of 4
443 Illumina indices was added via PCR to the MspI adapter. Therefore, sequences from each individual
444 could be identified by the unique combination of barcode and index. A complete list of barcodes and
445 indices used in this study can be found in the Supplementary Tables 3 and 4, respectively. Amplified
446 libraries were pooled, cleaned with a Qiaquick PCR Purification Kit (Qiagen, Inc., Valencia, CA,
447 U.S.A.), and diluted to 4nM prior to sequencing. Prepared genomic DNA libraries constructed from
448 303 *H. virescens* individuals were spread across 9 full and partial Illumina MiSeq runs. The MiSeq
449 reagent kit v. 2 was used for initial preparation of the mapping family and the inbred line. All
450 subsequent preparations, including re-runs of the mapping family and inbred line were prepared with
451 the MiSeq reagent kit v. 3.

452

453 *De novo marker formation*

454 Overlapping paired-end reads were merged with FLASH (Magoc & Salzberg, 2011), and Stacks
455 v. 1.09 (Catchen *et al.*, 2011; 2013) was used for demultiplexing and *de novo* formation of loci.
456 Merged paired-end reads were filtered for quality using the process_radtags script. Further quality
457 filtering entailed removal of reads when: 1) they did not have an intact EcoRI cut site, 2) had a quality
458 score < 30, or 3) were smaller than 350 bp. We did not allow process_radtags to rescue reads where
459 barcode sequences contained an error. All remaining merged reads were truncated at a length of 350
460 bp, and fed into the Stacks pipeline.

461

462 *Stacks parameter settings*

463 Reads from all individuals were run through ustacks with the following parameter settings: -m
464 3, -M 14 (allowing for 5% nucleotide mismatch rate between alleles per individual), -max_locus_stacks
465 2, --alpha 0.05. A consensus catalog of loci was first formed using the parents of the mapping cross
466 with cstacks, where the -n 14 parameter allowed for a 5% between individual nucleotide mismatch rate.
467 For the mapping family, genotype calls were made using sstacks prior to field- and colony-strain alleles
468 being added to the catalog. Progeny genotypes were automatically corrected using the Stacks
469 genotypes script. Twenty-four individuals of each colony and field-collected strain collected in 2012
470 were later added to the catalog, and all field-collected and laboratory-reared populations were also
471 genotyped using sstacks.

472

473 *Data analyses*

474 All population genomic and linkage analysis were conducted in R version 3.1.2 (R core team,
475 2014).

476

477 *Genomic diversity among H. virescens populations*

478 In total, we sequenced the 13 surviving males from an inbred line subjected to 10 generations of
479 sib-mating, 42-46 males per colony strain, and 30 males per field-collected population (Table 1). Prior
480 to running sequence data through the Stacks pipeline, we checked individual read counts across
481 populations to ensure uniformity (Supplementary Figure 1). Twelve of the 204 individuals sequenced
482 had too few (< 90,000) or too many reads (> 688,000) and were removed from the dataset prior to
483 analysis, following Bi *et al.* (2013). From our Stacks output, we constructed 4 different sets of
484 consensus loci present across populations. These subsets, containing a core overlapping set of 125 loci,
485 and increasing in size from 125 to 1231, consisted of marker sets with varying percentages of missing
486 genotype calls (range = 11.2-29.5%) (Supplementary Table 1). We used these 4 different subsets to
487 examine and compare changes in genomic diversity across populations.

488 We estimated the mean number of unique alleles present per locus, and corresponding 95% non-
489 parametric bootstrapped confidence intervals (N = 5000) across populations using a custom-written R
490 script. Each allele represented the accumulation of SNPs within a 350bp locus, analogous to a
491 haplotype. At a given locus, and within each population of size N (see Table 1 for sample sizes), alleles
492 (total = 2N) were randomly sampled without replacement either 6, 12, 18, or 24 times. Then the
493 number of unique alleles were counted for each sampling regime. Due to their small sample size
494 resulting from intensive inbreeding, we only sampled 6, 12, and 18 alleles per locus for the inbred line.
495 Our analysis focused primarily on the subset of loci containing 378 consensus loci because genotype
496 calls were present for at least 10 individuals per population. We also calculated two measures of
497 nucleotide diversity per 350 bp locus using the R package, pegas (v. 0.6; Paradis, 2010): π (Nei, 1987)
498 and S_K corrected for sample size (Charlesworth & Charlesworth, 2010; Watterson, 1975). We then

499 generated population-level genome-wide means and 95% non-parametric bootstrapped (N = 5000)
500 confidence intervals for each metric (Table 2).

501

502 *Genetic divergence between populations*

503 To determine the degree of genetic diversity accounted for by differences between our field-
504 collected, laboratory-reared, Bt-selected, and inbred populations, we calculated Weir and Cockerham's
505 F_{ST} (Weir and Cockerham, 1984) along with corresponding 95% bootstrapped confidence intervals (N =
506 5000). Calculations were carried out using the R package, *diveRsity* (v. 1.9.89; Keenan *et al.*, 2013).

507

508 *Estimating the inbreeding coefficient*

509 To estimate our marker-based inbreeding coefficient, we examined multiple sets of loci
510 (Supplementary Table 1) and found that trends across all datasets were similar (data not shown).
511 However, we reported F_{IT} values from a set of 378 loci because this reduced dataset contained few
512 missing genotypes per population, while still making inferences from several hundred loci. We
513 calculated F_{IT} for the inbred line relative to the non-selected (YDK) population after Keller and Waller
514 (2002), where $(1-F_{IS})(1-F_{ST}) = 1-F_{IT}$. F_{IS} was the level of inbreeding within the inbred line, calculated
515 as $F_{IS} = 1-(H_O/H_E)$, where H_O and H_E were calculated for each locus using the R package *adegenet* (v.
516 1.4-2; Jombart, 2008). F_{ST} was the accumulated effect of inbreeding over time, calculated as 1-
517 $(H_E(\text{inbred line})/H_E(\text{YDK}))$.

518

519 *Amplification and genotyping of PCR-based markers for the mapping family*

520 Progeny from the mapping family were genotyped at three additional loci, and these loci were
521 mapped alongside our ddRAD-seq markers to validate our linkage groupings. We targeted the

522 previously described *H. virescens* genes *DesatI*, *ABCC2*, and *HevCaLP* via PCR followed by gel
523 electrophoresis, or direct sequencing. Amplification and genotyping protocols were as follows.

524 A 468 bp fragment from *DesatI* was amplified in a 30 μ L reaction with forward and reverse
525 primers [5'-TGAGGGACCATCGTCTCCAT-3'] and [5'-CACTGCTACATTTTGGGCAG-3'],
526 respectively (Ward, 2009). Each reaction contained 6 μ L of 5 \times GoTaq buffer (ProMega), 29 μ M per
527 dNTP, 92 ng per primer, 0.75 U GoTaq polymerase, and *ca.* 1 μ g genomic DNA. Sample DNA was
528 amplified alongside a negative control (here and throughout), where pcr-grade H₂O was substituted for
529 genomic DNA. Reactions were incubated at 95°C for 1 min followed by 35 cycles of 95°C for 1min,
530 52°C for 1min, and 72°C for 2 min. PCR products were purified using a standard ethanol precipitation,
531 and directly sequenced on an ABI3730xl (Applied Biosystems, San Francisco, CA). A single
532 nucleotide polymorphism (cytosine to tyrosine substitution) at bp 36 was found in the YHD2 parent of
533 the mapping cross. Offspring were genotyped at this locus using PolyPhred (Nickerson *et al.*, 1997),
534 and genotype calls were visually confirmed using consed (Gordon *et al.*, 1998).

535 An intronic region of the *ABCC2* gene previously described by Gahan *et al.* (2010) was
536 amplified using primers Hs-ABC2dU02-F1 [5' – TGGTTACAAGAAATAGAAAATGCAAC-3'] and
537 Hs-ABC2eU03-R2 [5' – CTTTCAAACCTGAACCGCATCAC – '3]. Each 30 μ L reaction volume
538 consisted of 6 μ L of 5 \times GoTaq buffer, 29 μ M per dNTP, 73 ng per primer, 0.75 U GoTaq polymerase,
539 and 1 μ g genomic DNA. Reactions were held at 95°C for 2 min followed by 30 cycles of 95°C for 30
540 sec, 58°C for 30 sec, and 72°C for 40 sec, and the resulting products were cleaned via ethanol
541 precipitation. Following sequencing on an ABI3730xl, chromatogram files were visualized using
542 FinchTV (version 1.3.1, PerkinElmer, Inc., Seattle, WA). As described by Gahan *et al.* (2010), the
543 YHD2 parent was homozygous for a 22 bp deletion, whereas the F₁ parent was heterozygous for this
544 deletion. Therefore, we examined the segregation of this deletion in the mapping family offspring,

545 which was detectable by the presence of a TAT sequence near amplicon bp 40.

546 Finally, the *HevCaLP* locus described by Gahan *et al.* (2007) was amplified in a multiplexed
547 reaction using three primers: the universal reverse primer [5'-
548 ATACGAGCTGACGACACGCTGGGAGA-3'], one forward primer that targets a retrotransposon
549 insertion conferring resistance to *Bacillus thuringiensis* [5' –
550 CGCAACGCGCGATCTACTCTTGTCACC – 3'], and another forward primer that targets wild-type
551 sequence [5' – AAGTGTCCCAGTCGATGCTGAA – 3']. An initial 20- μ l reaction contained 4 μ l 5 \times
552 GoTaq buffer, 29 μ M per dNTP, 56 ng per primer, 0.5 U GoTaq polymerase, and 1 μ g genomic DNA.
553 Reactions were incubated at 95°C for 2 min, followed by 30 cycles of 95°C for 30 sec, 58°C for 20 sec,
554 and 72°C for 40 sec. A reconditioning reaction, aimed at reducing heteroduplex formation, was set up
555 as above, but incubated for 3 rather than 30 cycles. These reactions were capable of producing two
556 amplicons, which differed in length by 76 bp. The YHD2 parent was homozygous for the long
557 amplicon (*ca.* 800 bp) containing the insertion that confers resistance to *Bacillus thuringiensis*, whereas
558 the F₁ parent was heterozygous for a long and short amplicon. PCR products from mapping family
559 offspring were run on a 2% agarose gel alongside Hyperladder I (Bioline, Taunton, MA) for
560 visualization and genotype scoring.

561

562 *Linkage mapping*

563 Double-digest RAD-seq markers present in fewer than 75% of the mapping family offspring
564 were filtered out, and the remainder were checked using a chi-square test for Mendelian segregation (α
565 = 0.01). PCR-based markers, as well as those ddRAD-seq markers that segregated in a mendelian
566 fashion were assigned to linkage groups (LOD = 5, maximum recombination fraction = 0.3) using the
567 onemap package (Margarido *et al.*, 2007) in R. We validated groupings by aligning all markers to the

568 *Bombyx mori* genome using Blastn in Kaikobase version 3.2.2 (<http://sgp.dna.affrc.go.jp/KAIKObase/>).
569 Furthermore, we confirmed that the locations of the *DesatI*, *ABCC2*, and *HevCaLP* pcr-based markers,
570 as well as ddRAD-seq markers found in their respective linkage groups aligned to the same *B. mori*
571 chromosomes (Table 4). Markers on each linkage group were ordered using the recombination
572 counting and ordering algorithm (RECORD; Van Os *et al.*, 2005). RECORD was chosen based upon
573 previous studies demonstrating the reliability of its performance (Collard *et al.*, 2009, Mollinari *et al.*,
574 2009). Recombination fractions were converted to centiMorgan distances using the Kosambi mapping
575 function (Kosambi, 1944). The final linkage map was drawn using Genetic Mapper version 0.5.

576

577 *Assessment of fine-scale differences in nucleotide diversity across laboratory-reared populations*

578 Mapped markers were examined for observed heterozygosity (H_o) and nucleotide diversity (π),
579 as above, for the inbred line, the Bt-selected line, one non-selected line (YDK), and one field-collected
580 population (LA), and these values were displayed for visual comparison in Figure 4. For each
581 population, at least 3 individuals must have been genotyped for a marker to be included in the analysis.
582 Then, we examined the distribution of mapped markers that retained polymorphism in the inbred line to
583 determine whether heterogeneity, or significant clustering, could be observed. Using a replicated G-
584 test of independence (Sokal and Rohlf, 1995), the distribution of polymorphic loci across all linkage
585 groups that contained five or more markers (24 of the 33 total linkage groups) was examined. Under
586 assumptions of homogeneity, we expected that the ratio of polymorphic to fixed loci across linkage
587 groups in the inbred line would follow Hartl and Clark's (2007):

$$588 H_T = H_o(1 - 1/2^{*N_e})^T, \text{ where}$$

589 H_o = initial observed heterozygosity in the population, which we set equal to the observed

590 heterozygosity of its ancestral YDK population.

591 N_e = the number of breeding adults in each generation, which we set equal to 2 according to our sib-
592 mating design.

593 T = the number of generations

594 According to this equation, the expected frequency of polymorphic loci was 0.015 within a linkage
595 group following 10 generations of sib-mating. Each linkage group was first examined for deviation
596 from this expected frequency of polymorphic loci using a G-test of independence with a Bonferroni-
597 adjusted alpha value ($\alpha = 0.002$) to account for multiple comparisons. Individual uncorrected G-values
598 produced for each of the 24 linkage groups were added together to generate a “pooled” G-value, and a
599 “total” G-value was calculated according to Sokal and Rohlf (1995). Finally, a heterogeneity G-value,
600 which tested the hypothesis that polymorphic loci were significantly more clustered within linkage
601 groups than due would be due to random chance, was calculated by taking the difference between total
602 and pooled G-values, then comparing it to a X^2 distribution with 23 degrees of freedom and an a priori
603 alpha value of 0.05.

604 Each marker that retained polymorphism in the inbred line was then examined for an excess of
605 intermediate frequency alleles, which could indicate either a population contraction or ongoing
606 balancing selection near that locus. For this, we used a Tajima's D test as calculated by the R package,
607 pegas (v. 0.6; Paradis, 2010). To tease apart the effects of population demographic changes and forces
608 of natural selection, we compared the Tajima's D values at each locus in the inbred line to those of their
609 ancestral population (YDK). We reasoned that significantly positive Tajima's D values that overlapped
610 between the two lines would be less likely due to the population demographic changes created by sib-
611 mating, and more likely due to selection. A Benjamini-Hochberg adjustment was applied to all
612 Tajima's D p-values to account for multiple hypothesis tests using the R package fdrtool (v.1.2.15;
613 Strimmer 2008).

614

615 **Acknowledgements**

616 Thanks to Dr. R. Whetten of North Carolina State University, and Dr. J. Schaff of the NCSU
617 Genomic Sciences Lab for their insightful suggestions on ways to improve our methods. Dr. S.
618 Micinski, Dr. J. Lopez, and Dr. J. Westbrook collected the moths used in this project. R. Waples
619 provided one of the custom scripts used in our data pipeline. This project is supported by the
620 Biotechnology Risk Assessment Program competitive grant number 2012-33522-19793 from the
621 USDA - National Institute of Food and Agriculture.

622

623

624

625 **References**

- 626
- 627 Baeshen, R., Ekechukwu, N., Toure, M., Paton, D., Coulibaly, M., Traoré, S. and Tripet, F. (2014).
628 Differential effects of inbreeding and selection on male reproductive phenotype associated with the
629 colonization and laboratory maintenance of *Anopheles gambiae*. *Malar J*, 13(1), p.19.
- 630 Baxter, S.W., McMillan, O., Chamberlain, N., French-Constant, R.H., Jiggins, C.D. (2009). Prospects
631 for Locating Adaptive Genes in Lepidopteran Genomes. From Molecular Biology and Genetics of the
632 Lepidoptera. CRC Press, Boca Raton, FL, USA. ed. by Goldsmith MR, Marec F. pp. 105-118.
- 633 Benjamini, Y., and Hochberg, Y. (1995). Controlling the False Discovery Rate: A practical and
634 powerful approach to multiple testing. *Journal of the Royal Statistical Society. Series B*
635 (*Methodological*), 57(1), pp. 289-300.
- 636 Bi, K., Linderoth, T., Vanderpool, D., Good, J., Nielsen, R. and Moritz, C. (2013). Unlocking the vault:
637 next-generation museum population genomics. *Molecular Ecology*, 22(24), pp.6018-6032.
- 638 Blanco, C. (2012). *Heliothis virescens* and Bt cotton in the United States. *GM Crops & Food*, 3(3),
639 pp.201-212.
- 640 Boller, E. (1972). Behavioral aspects of mass-rearing of insects. *Entomophaga*, 17(1), pp.9-25.
- 641 Catchen, J., Amores, A., Hohenlohe, P., Cresko, W., Postlethwait, J. and De Koning, D. (2011). Stacks:
642 Building and Genotyping Loci De Novo From Short-Read Sequences. *G3: Genes Genomes Genetics*,
643 1(3), pp.171-182.
- 644 Catchen, J., Hohenlohe, P., Bassham, S., Amores, A. and Cresko, W. (2013). Stacks: an analysis tool set
645 for population genomics. *Molecular Ecology*, 22(11), pp.3124-3140.
- 646 Charlesworth, B., Charlesworth, D. (2010). Elements of Evolutionary Genetics. Roberts and Company
647 Publishers, Greenwood, CO, USA. pp. 29.
- 648 Charlesworth, D., and Willis, J.H. (2009). The genetics of inbreeding depression. *Nature Reviews*
649 *Genetics*, 10, pp.783-796.
- 650 Collard, B., Mace, E., McPhail, M., Wenzl, P., Cakir, M., Fox, G., Poulsen, D., Jordan, D. (2009). How
651 accurate are the marker orders in crop linkage maps generated from large marker datasets? *Crop and*
652 *Pasture Science*, 60(4), pp.362-372.
- 653 Collins A.M. Artificial selection of desired characteristics in insects. Advances and Challenges in
654 Insect Rearing, Eds. EG King and NC Leppla. USDA Technical Bulletin. Jan. 1984 pp. 9-19.
- 655 Davey, J.W., Cezard, T., Fuentes-Utrilla, P., Eland, C., Gharbi, K., and Blaxter, M.L. (2012). Special
656 features of RAD sequencing data: implications for genotyping. *Molecular Ecology*, 22, pp.3151-3164.
- 657 Dekker, T., Ibba, I., Siju, K.P., Stensmyr, M.C., Hansson, B.S. (2006). Olfactory shifts parallel
658 superspecialism for toxic fruit in *Drosophila melanogaster* sibling, *D. sechellia*. *Current Biology*,
659 16(1), pp.101-109.
- 660 de Valdez, M.F.W., Nimmo, D., Betz, J., Gong, H.F., James, A.A., Alphey, L. and Black, W.C. (2011).
661 Genetic elimination of dengue vector mosquitoes. *Proceedings of the National Academy of Sciences*,

- 662 USA, 108(12), pp.4772-4775.
- 663 Dobzhansky, T., and Spassky, N. (1954). Environmental modification of heterosis in *Drosophila*
664 *pseudoobscura*. *Proceedings of the National Academy of Sciences, USA*, 40(6), pp.407-415.
- 665 Elshire, R.J., Glaubitz, J.C., Sun, Q., Poland, J.A., Kawamoto, K., Buckler, E.S., Mitchell, S.E. (2011).
666 A robust, simple genotyping-by-sequencing (GBS) approach for high diversity species. *PLoS ONE*,
667 6(5), pp. e19379.
- 668 Etzel, L.K., and Legner, E.F. (1999). Culture and Colonization, Chapter 7 (pp. 125-176) in Handbook
669 of Biological Control, Academic Press, San Diego, CA, USA. ed. Bellows TS and Fisher TW.
- 670 Falconer, D.S., and Mackay TFC (1996) Introduction to Quantitative Genetics. Fourth Edition,
671 Longman Group Ltd. Essex, England. pp. 89.
- 672 Fox, C.W., Scheibly, K.L., Smith, B.P., Wallin, W.G. (2007). Inbreeding depression in two seed-feeding
673 beetles *Callosobruchus maculatus* and *Stator limbatus* (Coleoptera: Chrysomelidae). *Bulletin of*
674 *Entomological Research*, 97(1), pp.49-54.
- 675 Fritz, M.L., Walker, E.D., Miller, J.R., Severson, D.W., Dworkin, I. (2015). Divergent host preferences
676 of above- and below-ground *Culex pipiens* mosquitoes and their hybrid offspring. *Medical and*
677 *Veterinary Entomology*, 29(2), pp. 115-123.
- 678 Gahan, L.J., Gould, F., Heckel, D.G. (2001). Identification of a gene associated with Bt resistance in
679 *Heliothis virescens*. *Science*, 293, pp.857-860.
- 680 Gahan, L.J., Pauchet, Y., Vogel, H., Heckel, D.G. (2010). An ABC transporter mutation is correlated
681 with insect resistance to *Bacillus thuringiensis* Cry1Ac toxin. *PLoS Genetics*, 6(12), pp. e1001248.
- 682 Gerloff, C.U., Ottmer, B.K., Schmid-Hempel, B. (2003). Effects of inbreeding on immune response and
683 body size in a social insect, *Bombus terrestris*. *Functional Ecology*, 17(5), pp. 582-589.
- 684 Goldman, I. F., Arnold, J., and Carlton, B. C. (1986). Selection for resistance to *Bacillus thuringiensis*
685 subspecies israelensis in field and laboratory populations of the mosquito *Aedes aegypti*. *Journal of*
686 *Invertebrate Pathology*, 47(3), pp. 317-324.
- 687 Gordon, D., Abajian, C., Green, P. (1998). Consed: a graphical tool for sequence finishing. *Genome*
688 *Research*, 8, pp. 195-202.
- 689 Gould, F., Anderson, A., Reynolds, A., Bumgarner, L., Moar, W. (1995). Selection and genetic analysis
690 of a *Heliothis virescens* (Lepidoptera: Noctuidae) strain with high levels of resistance to *Bacillus*
691 *thuringiensis* toxins. *Journal of Economic Entomology*, 88(6), pp.1545-1559.
- 692 Groot, A., Gemenio, C., Brownie, C., Gould, F., and Schal, C. (2005). Male and female antennal
693 responses in *Heliothis virescens* and *H. subflexa* to conspecific and heterospecific sex pheromone
694 compounds. *Environmental Entomology*, 34(2), pp. 256-263.
- 695 Groot, A.T., Classen, A., Inglis, O., Blanco, C.A., Lopez, J., Teran Vargas, A., Schal, C., Heckel, D.G.,
696 and Schofl, G. (2011). Genetic differentiation across North America in the generalist moth *Heliothis*
697 *virescens* and the specialist *H. subflexa*. *Molecular Ecology*, 20, pp. 2676-2692.
- 698 Hartl, D.L., and Clark, A.G. (2007). Principles of Population Genetics, 4th ed. Sunderland, MA:

- 699 Sinauer Associates, Inc.
- 700 Hoy, M.A. (1990). Genetic improvement of parasites and predators. FFTC-NARC International
701 Seminar on 'The use of parasitoids and predators to control agricultural pests', Tukuha Science City,
702 Ibaraki-ken, 305 Japan, October 2-7, 1989. pp.15
- 703 Huettel, M.D. (1976). Monitoring the quality of laboratory-reared insects: A biological and behavioral
704 perspective. *Environmental Entomology*, 5(5), pp. 807-814.
- 705 Jombart, T. (2008). adegenet: a R package for the multivariate analysis of genetic markers.
706 *Bioinformatics*, 24, pp. 1403-1405.
- 707 Joyner, K. and Gould, F. (1985). Developmental consequences of cannibalism in *Heliothis zea*
708 (Lepidoptera: Noctuidae). *Annals of the Entomological Society of America*, 78, pp. 24-28.
- 709 Kajitani, R., Toshimoto, K., Noguchi, H., Toyoda, A., Ogura, Y., Okuno, M., Yabana, M., Harada, M.,
710 Nagayasu, E., Maruyama, H., Kohara, Y., Fujiyama, A., Hayashi, T., and Itoh, T. (2014). Efficient *de*
711 *novo* assembly of highly heterozygous genomes from whole-genome shotgun short reads. *Genome*
712 *Research*, 24, pp. 1384-1395.
- 713 Keenan, K., McGinnity, P., Cross, T.F., Crozier, W.W., and Prodöhl, P.A. (2013). diveRsity: An R
714 package for the estimation of population genetics parameters and their associated errors. *Methods in*
715 *Ecology and Evolution*, doi: 10.1111/2041-210X.12067.
- 716 Keller, L.F., and Waller, D.M., (2002). Inbreeding effects in wild populations. *Trends in Ecology and*
717 *Evolution*, 17(5), pp. 230-241.
- 718 Kim, S.H., Cheng, K.M., Ritland, C., Ritland, K., and Silversides, F.G., (2007). Inbreeding in the
719 Japanese quail estimated by pedigree and microsatellite analyses. *Journal of Heredity*, 98(14), pp. 378-
720 381.
- 721 Kosambi, D.D. (1944). The estimation of map distance from recombination values. *Annals of*
722 *Eugenics*, 12, pp. 172-175.
- 723 Li, R., Li, Y., Fang, X., Yang, H., Wang, J., Kristiansen, K., and Wang, J. (2009). SNP detection for
724 massively parallel whole-genome resequencing. *Genome Research*, 19, pp. 1124-1132.
- 725 Mackauer, M. (1976). Genetic problems in the production of biological control agents. *Annual Review*
726 *of Entomology*, 21, pp. 369-385.
- 727 Mackay, T.F.C., Richards, S., Stone, E.A., Barbadilla, A., Ayroles, J.F., Zhu, D., *et al.*, (2012) The
728 *Drosophila melanogaster* genetic reference panel. *Nature*, 482, pp. 173-178.
- 729 Magoc, T., and Salzberg, S. (2011). FLASH: Fast length adjustment of short reads to improve genome
730 assemblies. *Bioinformatics*, 27(21), pp. 2957-2963.
- 731 Margarido, G. R. A., Souza, A. P. and Garcia, A. A. F. (2007). OneMap: software for genetic mapping
732 in outcrossing species. *Hereditas*, 144, pp. 78-79.
- 733 Mita, K., Kasahara, M., Sasaki, S., Nagayasu, Y., Yamada, T., Kanamori, H., Namiki, N., Kitagawa, M.,
734 Yamashita, H., Yasukochi, Y., Kadono-Okuda, K., Yamamoto, K., Ajimura, M., Ravikumar, G.,
735 Shimomura, M., Nagamura, Y., Shin-i, T., Abe, H., Shimada, T., Morishita, S., and Sasaki, T. (2004)

- 736 The Genome Sequence of Silkworm, *Bombyx mori*. *DNA Research*, 11(1), pp. 27-35
- 737 Mollinari, M., Margarido, G.R., Vencovsky, R., Garcia, A.A. (2009). Evaluation of algorithms used to
738 order markers on genetic maps. *Heredity*, 103(6), pp. 494-502.
- 739 Mukhopadhyay, J., Rangel, E.F., Ghosh, K., Munstermann, L.E. (1997). Patterns of genetic variability
740 in colonized strains of *Lutzomyia longipalpis* (Diptera: Psychodidae) and its consequences. *American*
741 *Journal of Tropical Medicine and Hygiene*, 57(2), pp. 216-221.
- 742 Munstermann, L.E. (1994). Unexpected genetic consequences of colonization and inbreeding: allozyme
743 tracking in Culicidae (Diptera). *Annals of the Entomological Society of America*, 87(2), pp. 157-164.
- 744 Nei M (1987) *Molecular evolutionary genetics*. New York: Columbia University Press.
- 745 Nickerson, D.A., Tobe, V.O., and Taylor, S.L. (1997). PolyPhred: automating the detection and
746 genotyping of single nucleotide substitutions using fluorescence-based resequencing. *Nucleic Acids*
747 *Research*, 25(14), pp. 2745-2751.
- 748 Nielsen, R., Paul, J.S., Albrechtsen, A., Song, Y.S. (2011). Genotype and SNP calling from next-
749 generation sequencing data. *Nature Reviews Genetics*, 12, pp. 443-451.
- 750 Norris, D.E., Shurtleff, A.C., Toure, Y.T., Lanzaro, G.C. (2001). Microsatellite DNA polymorphism and
751 heterozygosity among field and laboratory populations of *Anopheles gambiae* s.s. (Diptera: Culicidae).
752 *Journal of Medical Entomology*, 38(2), pp. 336-340.
- 753 Oppenheim, S.J., Gould, F., Hopper, K.R. (2012). The genetic architecture of a complex ecological
754 trait: host plant use in the specialist moth, *Heliothis subflexa*. *Evolution*, 66(11), pp. 3336-3351.
- 755 Paradis, E. (2010). pegas: an R package for population genetics with an integrated-modular approach.
756 *Bioinformatics*, 26, 419-420.
- 757 Paterson, A.H. (1996). Making genetic maps. A.H. Paterson (Ed.), *Genome Mapping In Plants*, R. G.
758 Landes Company, San Diego, California, pp. 23–29
- 759 Peterson, B.K., Weber, J.N., Kay, E.H., Fisher, H.S., Hoekstra, H.E. (2012). Double digest RADseq:
760 An inexpensive method of *de novo* SNP discovery and genotyping in model and non-model species.
761 *PLoS ONE*, 7(5), e37135.
- 762 Poland, J.A., Brown, P.H., Sorrells, M.E., Jannink, J.L. (2012). Development of high-density genetic
763 maps for barley and wheat using a novel two-enzyme genotyping-by-sequencing approach. *PLoS ONE*,
764 7(2), pp. e32253.
- 765 Pootakham, W., Jomchai, N., Ruang-areerate, P., Shearman, J. R., Sonthirod, C., Sangsrakru, D.
766 Tragoonrun, S., Tangphatsornruang, S., (2015). Genome-wide SNP discovery and identification of QTL
767 associated with agronomic traits in oil palm using genotyping-by-sequencing (GBS). *Genomics*,
768 105(5), pp. 288-295.
- 769 Pradeep, A. R., Chatterjee, S. N., & Nair, C. V. (2005). Genetic differentiation induced by selection in
770 an inbred population of the silkworm *Bombyx mori*, revealed by RAPD and ISSR marker systems.
771 *Journal of Applied Genetics*, 46(3), pp. 291.
- 772 R core team (2014). R: A language and environment for statistical computing. R Foundation for

- 773 Statistical Computing, Vienna, Austria. URL <http://www.R-project.org/>.
- 774 Raulston, J.R. (1975). Tobacco budworm: observations on the laboratory adaptation of a wild strain.
775 *Annals of the Entomological Society of America*, 68, pp. 139-142.
- 776 Roush, R.T. (1986). Inbreeding depression and laboratory adaptation in *Heliothis virescens*
777 (Lepidoptera: Noctuidae). *Annals of the Entomological Society of America*, 79, pp. 583-587.
- 778 Rumball, W., Franklin, I., Frankham, R., Sheldon, B. (1994). Decline in heterozygosity under full-sib
779 and double first-cousin inbreeding in *Drosophila melanogaster*. *Genetics*, 136, pp. 1039-1049.
- 780 Shaw, K.L. (2000). Interspecific genetics of mate recognition: inheritance of female acoustic preference
781 in Hawaiian crickets. *Evolution*, 54(4), pp.1303-1312.
- 782 Sheck, A. L., and Gould, F. (1995). Genetic analysis of differences in oviposition preferences of
783 *Heliothis virescens* and *H. subflexa* (Lepidoptera: Noctuidae). *Environmental Entomology*, 24(2), pp.
784 341-347.
- 785 Sheck, A. L., and Gould, F. (1996). The genetic basis of differences in growth and behavior of
786 specialist and generalist herbivore species: selection on hybrids of *Heliothis virescens* and *Heliothis*
787 *subflexa* (Lepidoptera). *Evolution*, 50(2), pp. 831-841.
- 788 Sheck, A. L., Groot, A. T., Ward, C. M., Gemeno, C., Wang, J., Brownie, C., Schal, C., and Gould, F.
789 (2006). Genetics of sex pheromone blend differences between *Heliothis virescens* and *Heliothis*
790 *subflexa*: a chromosome mapping approach. *Journal of Evolutionary Biology*, 19(2), pp. 600-617.
- 791 Singh, R., Tan, S.G., Panandam, J.M., Rahman, R.A., Ooi, L.C.L., Low, E.L., Sharma, M., Jansen, J.,
792 and Cheah, S. (2009). Mapping quantitative trait loci (QTLs) for fatty acid composition in an
793 interspecific cross of oil palm. *BMC Plant Biology*, 9, 114.
- 794 Sokal, R.R., and Rohlf, F.J. (1995). *Biometry: The Principles and Practices of Statistics in Biological*
795 *Research*. 3rd. ed. New York: W.H. Freeman and Company.
- 796 Sokolowski, M.B. (1980). Foraging strategies of *Drosophila melanogaster*: A chromosomal analysis.
797 *Behavioral Genetics*, 10(3), pp. 291-302.
- 798 Strimmer, K. (2008). *fdrtool*: a versatile R package for estimating local and tail area-based false
799 discovery rates. *Bioinformatics Applications*, 24(12), pp. 1461-1462.
- 800 Tajima, F. (1989) Statistical method for testing the neutral mutation hypothesis by DNA polymorphism.
801 *Genetics*, 123, pp. 585-595.
- 802 Taylor, M.F.J., Heckel, D.G., Brown, T.M., Kreitman, M.E., Black, G. (1993). Linkage of pyrethroid
803 resistance to a sodium channel locus in the tobacco budworm. *Insect Biochemistry and Molecular*
804 *Biology*, 23(7), pp. 763-775.
- 805 Tomaru, M., Doi, M., Higuchi, H., Oguma, Y. (2000). Courtship song recognition in the *Drosophila*
806 *melanogaster* complex: heterospecific songs make females receptive in *D. melanogaster*, but not in *D.*
807 *sechellia*. *Evolution*, 54(4), pp. 1286-1294.
- 808 Turissini, D.A., Gamez, S., White, B.J. (2014). Genome-wide patterns of polymorphism in an inbred
809 line of the African malaria mosquito, *Anopheles gambiae*. *Genome Biology and Evolution*, 6(11), pp.

810 3094-3104.

811 Van Os, H., Stam, P., Visser, R.G.F., Van Eck, H.J. (2005). RECORD: a novel method for ordering loci
812 on a genetic linkage map. *Theoretical and Applied Genetics*, 112, pp. 30-40.

813 Ward, M.D. (2009). Genetics of Sex Pheromones: Mapping Desaturase Genes in *Heliothis* Species
814 (Unpublished Master's Thesis). North Carolina State University, Raleigh, NC.

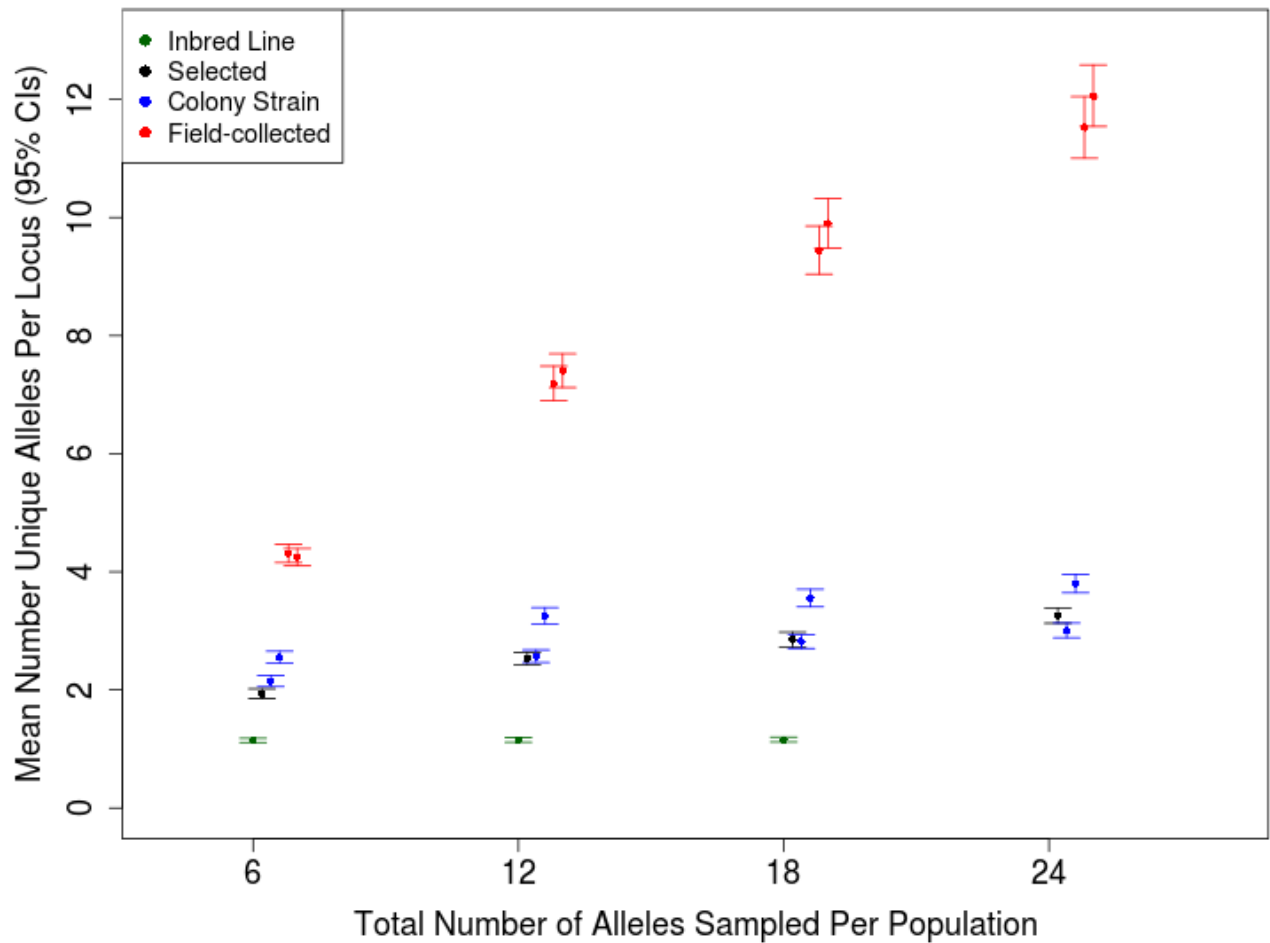
815 Watterson, G.A. (1975). On the number of segregating sites in genetical models without recombination.
816 *Theoretical Population Biology*, 7, pp. 256-276.

817 Weir, B.S., and Cockerham C.C. (1984). Estimating F-statistics for the analysis of population structure.
818 *Evolution*, 38(6), pp. 1358-1370.

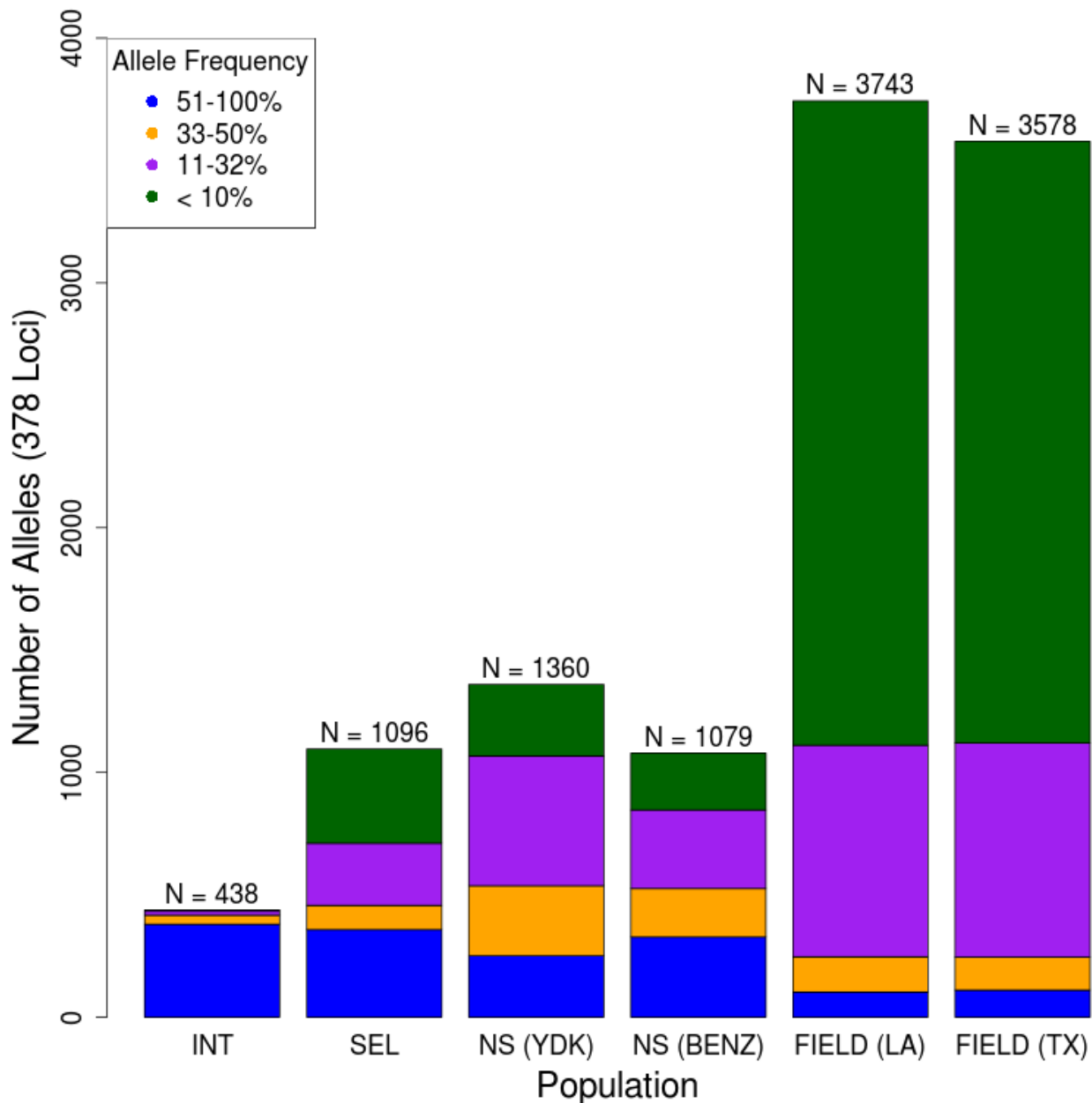
819 Xu, P., Xu, S., Wu, X., Tao, Y., Wang, B., Wang, S., Qin, D., Lu, Z., Li, G. (2014). Population genomic
820 analyses from low-coverage RAD-Seq data: a case study on the non-model cucurbit bottle gourd. *The*
821 *Plant Journal*, 77, pp. 430-442.

822 You, M., Yue, Z., He, W., Yang, X., Yang, G., Xie, M., Zhan, D., Baxter, S.W., et al. (2013). A
823 heterozygous moth genome provides insights into herbivory and detoxification. *Nature Genetics*, 45,
824 pp. 220-225.

825 **Figures**
826

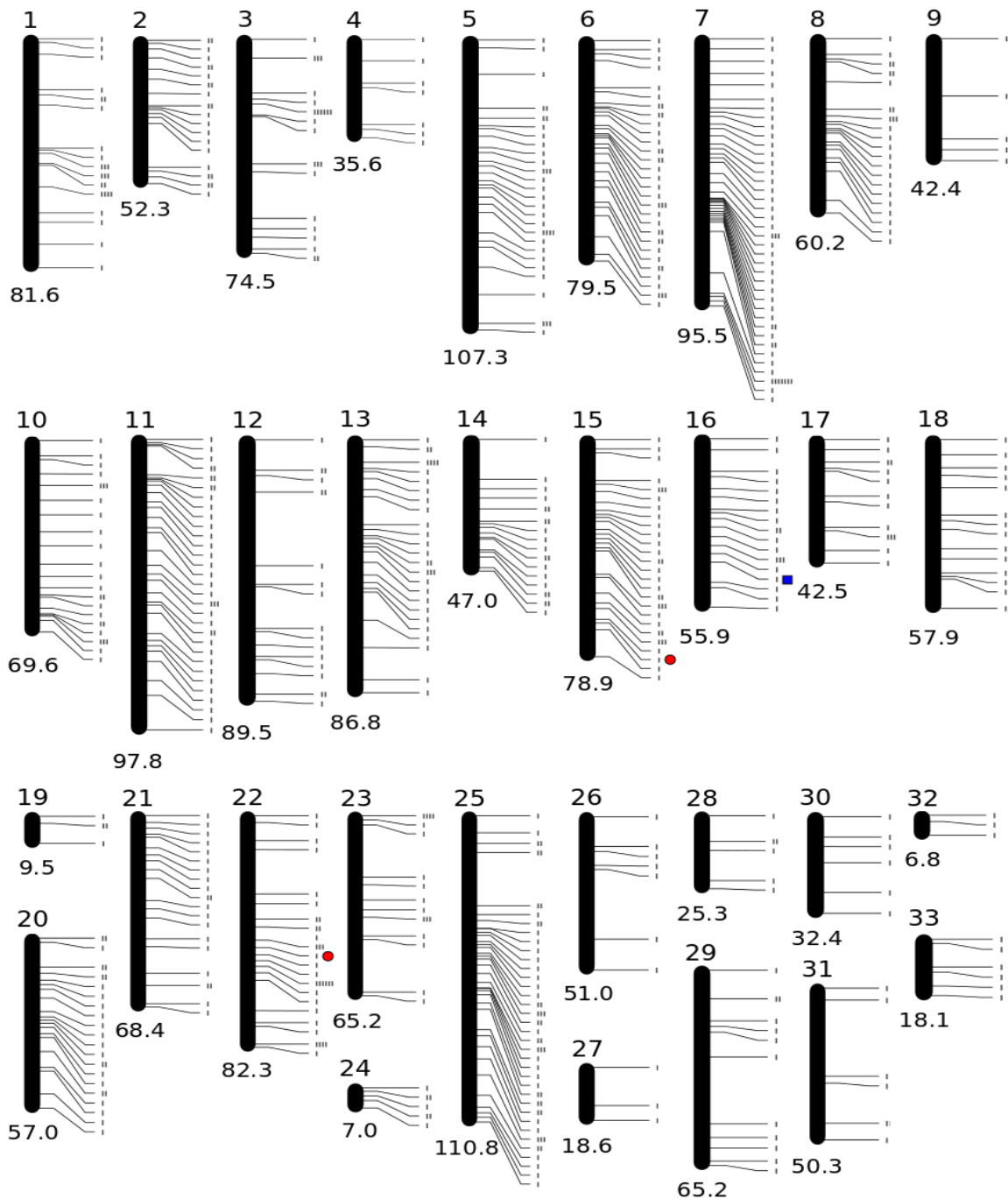


828 **Figure 1** – Mean numbers of unique alleles detected among 378 loci depends upon the number of
829 alleles sampled per population. For all but the inbred line (in green), as sample size increases, so does
830 the number of unique alleles detected per locus. Due to low survivorship in the inbred line, no mean
831 was computed for an allelic sample size per locus of 24.
832
833

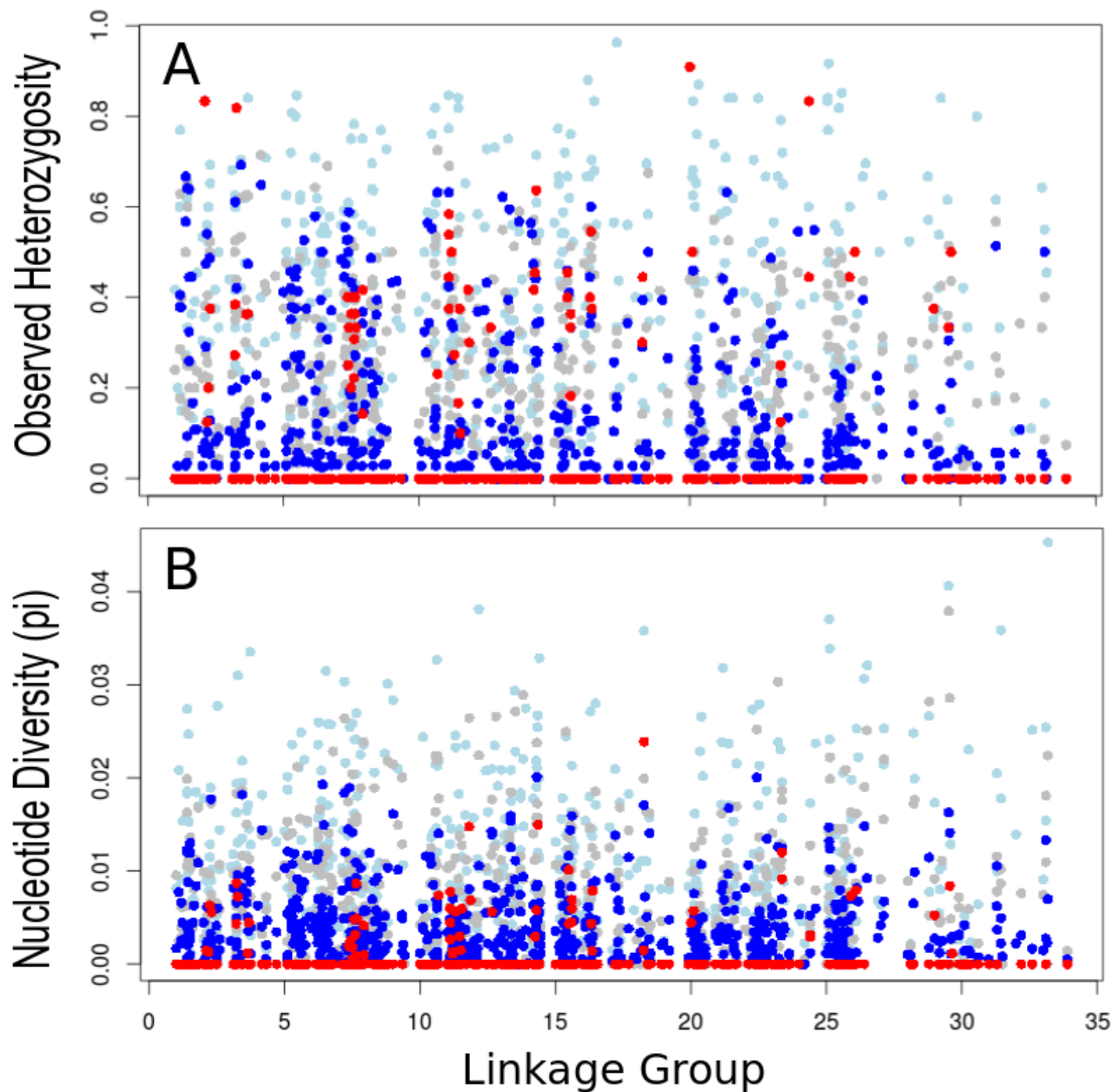


835 **Figure 2** – Total numbers of unique alleles detected in sib-mated (INT), Bt-selected (SEL), non-
836 selected (NS), and field-collected (FIELD) populations based upon random sampling of 18 haplotypes
837 per population per locus. Alleles were binned and color-coded according to the frequencies at which
838 they were present out of 18 total haplotypes. Numbers above each bar represent the total number of
839 unique alleles found per population out of 378 loci.

840
841



843 **Figure 3** – *Heliothis virescens* linkage map with a total length of 1919.5 cM. Centimorgan lengths are
844 below each linkage group. Each tick mark represents an individual marker that mapped to a particular
845 position in the linkage group. Red circles next to linkage groups 15 and 22 represent the positions of
846 the *HevCaLP*, and the *ABCC2*, respectively. The blue square represents the position of the delta-11-
847 desaturase on linkage group 16.



849 **Figure 4** – Observed heterozygosity (A) and nucleotide diversity (B) estimates per mapped marker site.
850 For comparative purposes, estimates for the Louisiana field population were included (light blue).
851 Grey and blue circles represent estimates in non-selected (YDK), and Bt-selected (YHD2) populations,
852 respectively. Red circles represent estimates in the inbred line following 10 generations of sib-mating.

853 **Tables**

854

855 **Table 1** - Population history, sample sizes (N) and ddRAD-seq read summary data. Filtered N refers to
856 the population sample size following removal of individuals with low read counts.

857

Population	Population History	Total N	Filtered N	Mean Number Reads per Individual
Inbred Line	10 generations of sib-mating	13	13	334689
YHD2	Bt-selected	43	41	346264
BENZ	Unselected	42	40	318636
YDK	Unselected	46	44	335728
LA2012	Field-collected	30	27	398336
TX2012	Field-collected	30	28	306164

859

860

861

862

863

864

865

866

867

868

869

870

871

872

873

874

875 **Table 2** – Genome-wide nucleotide diversity values per 350 bp locus across populations. All values were calculated using a conservatively
 876 chosen set of 378 loci for which at least 10 individuals were genotyped within each population. Genome-wide values represent population-
 877 level π and S_K averaged across all loci. The abbreviations Bt-sel and NS stand for Bt-selected and non-selected, respectively.
 878

Population	Mean Observed Heterozygosity	Genome-wide π (2.5, 97.5% CIs)	Max π	Genome-wide S_K (2.5, 97.5% CIs)	Max S_K
Inbred line	0.06	0.0006 (0.0004, 0.0008)	0.013	0.15 (0.10, 0.19)	2.95
Bt-Sel (YHD2)	0.15	0.0040 (0.0035, 0.0043)	0.020	1.67 (1.54, 1.81)	6.53
NS (YDK)	0.26	0.0067 (0.0061, 0.0073)	0.028	1.66 (1.53, 1.80)	6.20
NS (BENZ)	0.22	0.0051 (0.0046, 0.0056)	0.026	1.35 (1.24, 1.47)	5.46
Field (LA)	0.46	0.0094 (0.0087, 0.0100)	0.034	5.35 (5.03, 5.68)	15.55
Field (TX)	0.40	0.0092 (0.0085, 0.0099)	0.033	5.02 (4.71, 5.35)	15.63

879

880 **Table 3** – Pairwise estimates of genetic divergence across 378 ddRAD-seq loci calculated according to Weir and Cockerham's F_{ST} . Estimates
 881 of F_{ST} are above the diagonal, while corresponding bootstrapped confidence intervals (2.5%, 97.5%) are presented below.
 882

	Inbred Line	Bt-Sel (YHD2)	NS (YDK)	NS (BENZ)	Field (LA)	Field (TX)
Inbred Line	-	0.6378	0.3272	0.6209	0.3873	0.3899
Bt-Sel (YHD2)	(0.5479, 0.7342)	-	0.2818	0.4854	0.3137	0.3175
NS (YDK)	(0.3096, 0.3465)	(0.1823, 0.3796)	-	0.3863	0.1656	0.1664
NS (BENZ)	(0.5705, 0.6645)	(0.3643, 0.5922)	(0.3345, 0.4288)	-	0.2552	0.2565
Field (LA)	(0.3663, 0.4078)	(0.2380, 0.3930)	(0.1521, 0.1850)	(0.2134, 0.2931)	-	0.0004
Field (TX)	(0.3681, 0.4106)	(0.2403, 0.3971)	(0.1529, 0.1861)	(0.2148, 0.2951)	(-0.0115, 0.0156)	-

883

884 **Table 4** – Linkage group (LG) correspondence with *B. mori* chromosome (Chr). Linkage groups with an asterisk contained one or more
885 markers that aligned uniquely to an unmapped *B. mori* sequence. Where linkage groups contained markers that aligned to more than one *B.*
886 *mori* chromosome, italicized marker names correspond to the italicized *B. mori* chromosome.
887
888

LG	<i>B.mori</i> Chr	Number Markers Aligned to <i>B.mori</i> Chr	Total markers in LG	LG Length (cM)	Average Marker Spacing (cM)	Names of Markers Aligned to <i>B.mori</i> Chr
1	25	3	25	81.6	3.3	19754, 29329, 20282
2	8	6	23	52.3	2.3	22394, 27499, 21475, 17667, 17852, 25500
3*	17	3	24	74.5	3.1	376, 21931, 25112
4	-		7	35.6	5.1	
5*	4	5	36	107.3	3.0	2556, 29595, 22095, 8695,15160
6	5	6	40	79.5	2.0	18575, 4268, 21041, 23679, 66, 5723
7	22	9	49	95.5	1.9	13710, 2328, 1443, 3654, 13880, 22897, 3123, 2430, 19858
8*	11	9	25	60.2	1.7	22584, 939, 3060, 18820, 23156, 17932, 202, 22290, 21173
9	23	2	7	42.4	6.1	4275, 1286
10	3	5	26	69.6	2.7	5015, 5329, 4265, 4545, 23262
11	9	8	37	97.8	2.6	19938, 6113, 16969, 19392, 1572, 89, 22588,19343
12	-		18	89.5	5.0	
13	13	6	31	86.8	2.8	2074, 15962, 26388, 1193, 16684,18825
14	19	1	21	47.0	2.2	14242
15	6	7	35	78.9	2.3	25943, 4579, 25083, 1752, 3385, 2784, 18178
16	23	4	20	55.9	2.8	2485, 16350, 806, 20599
17	7	2	14	42.5	3.0	29343, 343
18	26	1	14	57.9	4.1	1113
19	11, 22	2	4	9.5	2.4	3536, <i>16185</i>
20*	10	6	25	57.0	2.3	17820, 2978, 18970, 12277, 16316, 12984

890 **Table 4 continued.**

891

892

LG	<i>B.mori</i> Chr	Number Markers Aligned to <i>B.mori</i> Chr	Total markers in LG	LG Length (cM)	Average Marker Spacing (cM)	Names of Markers Aligned to <i>B.mori</i> Chr
21	21	3	21	68.4	3.3	26113, 20542, 7624
22	15	2	33	82.3	2.5	21411, 5773
23	14	1	17	65.2	3.8	4156
24			7	7.0	1.0	
25	12,1	4	53	110.8	2.0	20280, 9211,12382, 23604
26			7	51.0	7.3	
27	2	1	3	18.6	6.2	5781
28			6	25.3	4.2	
29	16	1	12	65.2	5.4	23481
30			6	32.4	5.4	
31			7	50.3	7.2	
32			3	6.8	2.3	
33	14, 28	2	6	18.1	3.0	6601, 20184

894

895

896

897

898