

1 **Substantial differences in bias between single-digest and double-digest**

2 **RAD-seq libraries: a case study**

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13 **Abstract**

14 The tradeoffs of using single-digest versus double-digest restriction site associated DNA
15 sequencing (RAD-seq) protocols have been widely discussed. However, no direct empirical
16 comparisons of the two methods have been conducted. Here, we sampled a single population of
17 Gulf pipefish (*Syngnathus scovelli*) and genotyped 444 individuals using RAD-seq. Sixty
18 individuals were subjected to single-digest RAD-seq (sdRAD-seq), and the remaining 384
19 individuals were genotyped using a double-digest RAD-seq (ddRAD-seq) protocol. We analyzed
20 the resulting Illumina sequencing data and compared the two genotyping methods when reads
21 were analyzed either together or separately. Coverage statistics, observed heterozygosity, and
22 allele frequencies differed significantly between the two protocols, as did the results of selection
23 components analysis. We also performed an *in silico* digestion of the Gulf pipefish genome and
24 modeled five major sources of bias: PCR duplicates, polymorphic restriction sites, shearing bias,
25 asymmetric sampling (i.e., genotyping fewer individuals with sdRAD-seq than with ddRAD-
26 seq), and higher major allele frequencies. This combination of approaches allowed us to
27 determine that polymorphic restriction sites, an asymmetric sampling scheme, mean allele
28 frequencies, and to some extent PCR duplicates all contribute to different estimates of allele
29 frequencies between samples genotyped using sdRAD-seq versus ddRAD-seq. Our finding that
30 sdRAD-seq and ddRAD-seq can result in different allele frequencies has implications for
31 comparisons across studies and techniques that endeavor to identify genome-wide signatures of
32 evolutionary processes in natural populations.

33

34 **Introduction**

35 Many questions in modern evolutionary biology require genetic information from
36 individuals. Important aspects of the evolutionary process, including adaptive divergence,
37 phylogenetic relationships, mating system dynamics, and the influence of neutral processes on
38 population differentiation, can be estimated only from reliable genotypes of individuals, which
39 often must be compared across populations and species. Consequently, the field of evolutionary
40 biology has eagerly adopted next-generation sequencing technologies and the myriad genotyping
41 techniques that go along with them.

42 One popular genotyping technique is restriction-site associated DNA sequencing (RAD-
43 seq), a reduced-representation approach that targets DNA sequences near restriction sites. This
44 family of techniques allows individuals to be genotyped at the same randomly sampled regions
45 throughout the genome and yields thousands of single-nucleotide polymorphism (SNP)
46 genotypes. Several versions of RAD-seq have been developed, and each one uses slightly
47 different methods to achieve the same purpose. Single-digest RAD-seq (sdRAD-seq) was the
48 original RAD-seq method; it uses one infrequently cutting restriction enzyme plus a sonication
49 step to generate short fragments for sequencing (Baird *et al.* 2008; Miller *et al.* 2007). Double-
50 digest RAD-seq (ddRAD-seq) uses two restriction enzymes and omits the sonication step
51 (Peterson *et al.* 2012). A method called 2bRAD-seq takes advantage of type-IIIB restriction
52 enzymes, which cut twice at a specified distance from the restriction site, releasing a short DNA
53 fragment of around 30 base pairs. Hence, a single type-IIIB restriction enzyme can be used to
54 generate fragments, but they will be very short by today's sequencing standards (Wang *et al.*
55 2012). These methods have been reviewed in detail elsewhere (Andrews *et al.* 2016; Andrews &
56 Luikart 2014; Puritz *et al.* 2014), and the utility of the RAD-seq approach in general for studying

57 adaptive variation has recently been called into question (Lowry *et al.* 2017a,b), although
58 researchers agree that RAD-seq can be a useful tool for molecular ecologists (Catchen *et al.*
59 2017, Lowry *et al.* 2017b, McKinney *et al.* 2017). The benefits of sdRAD-seq versus ddRAD-
60 seq have also been debated in the literature (Andrews *et al.* 2014; Andrews & Luikart 2014;
61 Puritz *et al.* 2014). Ultimately, each method has its own sets of biases and technical issues, and
62 the choice will depend on the focus of the study, the study organism, and the budget allocated to
63 the project (Andrews *et al.* 2016).

64 A major issue in RAD-seq studies is that restriction enzyme cut sites can be polymorphic.
65 Polymorphic restriction sites result in some individuals not being genotyped at particular loci or
66 having a homozygous genotype called when the individual is actually heterozygous (Davey *et al.*
67 2013), a phenomenon known as allelic dropout. These false homozygous calls decrease the
68 observed heterozygosity at loci with polymorphic restriction sites (Andrews *et al.* 2016),
69 resulting in biased summary statistics (Arnold *et al.* 2013). False homozygote calls can lead to
70 high error rates even with sufficient depth of coverage (Henning *et al.* 2014), because
71 polymorphic sites are simply not genotyped (Davey *et al.* 2013). Bias due to allelic dropout may
72 be limited unless effective population sizes are large ($N_e > 10^5$; Andrews *et al.* 2016; Gautier *et*
73 *al.* 2013) or when polymorphism is high (Cariou *et al.* 2016). Although Andrews *et al.* (2016)
74 suggest that loci with null alleles may be identified in a dataset by high variance in coverage
75 depth across samples, this type of filtering step is not typically a part of RAD-seq analyses.
76 Nevertheless, typical filtering steps to retain loci with high coverage across individuals and with
77 allele frequencies above a cutoff may remove many loci experiencing allelic dropout (Andrews
78 *et al.* 2016), and a recent method uses a Bayesian model to identify loci likely suffering from
79 bias due to polymorphic restriction sites and flags them for removal from the analysis (Cooke *et*

80 *al.* 2016). Of the various RAD-seq methods, those using multiple restriction enzymes (e.g.
81 ddRAD-seq) are likely to be more greatly affected by allelic dropout than those using a single
82 restriction enzyme (e.g. sdRAD-seq) because there are at least twice as many potentially
83 polymorphic restriction sites involved (Andrews *et al.* 2016; Arnold *et al.* 2013).

84 Other sources of error in RAD-seq studies emerge from the PCR amplification step. The
85 most important problem is the production of PCR duplicates, which stem from a random allele at
86 a given locus being amplified more than the other allele and result in a falsely homozygous
87 genotype (Andrews *et al.* 2016). These false homozygote calls cause the same biases as allelic
88 dropout and also result in variance in coverage depth at a locus (Andrews *et al.* 2016), making it
89 difficult to differentiate between allelic dropout and PCR duplicates [unless the study design
90 incorporates a way of identifying PCR duplicates, see Casbon *et al.* (2011); Davey *et al.* (2011);
91 Schweyen *et al.* (2014); Tin *et al.* (2015); Andrews *et al.* (2016)]. Although PCR duplicates
92 should impact ddRAD-seq and sdRAD-seq libraries at a similar rate as long as the number of
93 PCR cycles is the same, sdRAD-seq produces fragments with different random break points,
94 permitting PCR duplicates to be removed during filtering steps (Andrews *et al.* 2014). A related
95 problem arises from GC bias during the PCR steps (Andrews *et al.* 2016; Davey *et al.* 2011), and
96 both GC bias and PCR duplicates are minimized by the use of a high-fidelity polymerase such as
97 Phusion (Puritz *et al.* 2014). Finally, PCR preferentially amplifies shorter fragments. This issue
98 will affect ddRAD-seq methods more than sdRAD-seq because the two restriction sites
99 determine the fragment length of RAD loci, whereas in sdRAD-seq the length of RAD loci is
100 randomly determined by shearing (Andrews *et al.* 2016).

101 A final source of variance in coverage depth among loci is a result of the shearing step in
102 sdRAD-seq. Shorter fragments (<10kb) shear less efficiently than longer fragments, resulting in

103 loci from shorter fragments having fewer reads (Davey *et al.* 2013). This phenomenon only
104 affects the sdRAD-seq method, and Andrews *et al.* (2016) suggest that its effects should be
105 minimal, because most sdRAD-seq studies use restriction enzymes whose recognition sites occur
106 rarely in the genome, resulting in mostly large fragments prior to shearing.

107 Although the biases in sdRAD-seq and ddRAD-seq have been deduced and evaluated
108 using simulation models, no study has used both methods and evaluated the impact of using two
109 different RAD-seq library preparation approaches to genotype individuals from a single
110 population. Here, we provide a case study, which demonstrates that the two methods produce
111 different allele frequency distributions for individuals from the same wild-caught population of
112 the Gulf pipefish, *Syngnathus scovelli*.

113 **Methods**

114 *Collection methods*

115 The Gulf pipefish, *S. scovelli*, is a sex-role-reversed marine fish in the family
116 Syngnathidae (seahorses, pipefishes, and seadragons). The species is found in the Gulf of
117 Mexico and along the Atlantic coast of Florida in shallow seagrass beds. Gulf pipefish were
118 collected from the Gulf of Mexico by seine net in Corpus Christi, TX (27°41'33"N,
119 97°10'54"W). Each fish was euthanized in MS-222, preserved in ethanol, and frozen until DNA
120 could be extracted.

121 We extracted DNA from tissue from the adult heads and from entire embryos using the
122 PureGene DNA extraction kit (QIAGEN). Genomic DNA quality was evaluated by visualizing
123 each sample on an agarose gel and each sample was quantified using a Qubit Fluorometer 2.0
124 (Life Technologies).

125 *sdRAD-seq library preparation*

126 We prepared a sdRAD-seq library using DNA from 30 pregnant males and 30 females
127 following the protocols described in Baird *et al.* (2008). From each sample, 1µg of DNA was
128 digested with 100 Units of *Pst*I-HF (New England Biolabs) at 37°C for 90 minutes. The
129 digestions were cleaned up using the DNA Clean & Concentrator-5 (Zymo) kit before the first
130 adapter ligation. These P1 adapters were identical to those used by Baird *et al.* (2008), and each
131 adapter contained a 6-bp barcode and an Illumina sequencing primer. The ligation used 1000U of
132 T4 DNA ligase (New England Biolabs) and NEB Buffer 2 (New England Biolabs) and was
133 incubated at 16°C for 30 minutes before heat inactivation. We pooled 25µL from each of 12
134 samples and sheared these pooled samples using a Bioruptor. Sheared DNA was cleaned up
135 using the DNA Clean & Concentrator-5 kit (Zymo) and eluted in 20µL. Each set of 12 pooled
136 samples remained separate until the final pooling step. This sdRAD-seq library was
137 electrophoresed on a 1.25% agarose gel stained with SafeView (ABMGood) and fragments in
138 the range of 300-700 bp were excised from the gel with a razorblade. DNA was recovered from
139 the gel slices using a Zymoclean Gel DNA Recover Kit (Zymo). The 5' and 3' overhangs were
140 removed using the Quick Blunting Kit (New England Biolabs), and then an adenosine base was
141 added to the 3' end of the fragments in a reaction with Klenow enzyme (New England Biolabs)
142 at 37°C for 30 minutes. The library was subsequently cleaned up using a DNA Clean &
143 Concentrator-5 kit (Zymo) and the P2 adapter, containing only the sequencing adapters and no
144 barcodes, was ligated onto the fragments with T4 DNA ligase (New England Biolabs). After
145 another cleanup step with the DNA Clean & Concentrator-5 kit (Zymo), PCR with Phusion
146 polymerase was run for 18 cycles (cycle conditions: 98°C for 30s; 18 cycles of 98°C for 10s,
147 60°C for 30s, 72°C for 10s; 72°C for 5min) in two separate reactions. Those reactions were

148 pooled and purified with DNA Clean & Concentrator-5 (Zymo). The cleaned PCR products for
149 each set of 12 pooled samples were pooled into a final library whose quality was determined
150 using a Qubit Fluorometer 2.0 (Invitrogen), and the library was sent to the University of Oregon
151 for 100bp single-end Illumina HiSeq 2000 sequencing.

152 *ddRAD-seq library preparation*

153 We prepared four ddRAD-seq libraries containing a total of 159 pregnant males, 8 non-
154 pregnant males, 160 offspring, and 57 females (384 individuals total). We followed the ddRAD-
155 seq library preparation method from Peterson *et al.* (2012) with several modifications described
156 elsewhere (Flanagan & Jones 2017; Flanagan *et al.* 2016). Briefly, 1 µg of genomic DNA from
157 each individual was digested with 100 units of *Pst*I-HF (New England Biolabs) and 25 units of
158 *Mbo*I (New England Biolabs) in a 3 hour, 37°C incubation. Following purification by AMPure
159 XP beads (Agilent), 250ng of each DNA sample was ligated to barcoded adapters using T4
160 ligase (Epicentre) in a 23°C incubation lasting 30 minutes followed by a 10 minute 65°C heat
161 shock. These adapters were identical to those used in sdRAD-seq library preparation (see above;
162 Baird *et al.* 2008). Ninety-six unique barcodes were used for each ddRAD-seq library, so we
163 pooled the adapter-ligated fragments from 96 individuals after an AMPure XP bead (Agilent)
164 purification. We extracted fragments in the range of 300-700 bp from a 1% agarose gel stained
165 with SafeView (ABMGood). Phusion polymerase (New England Biolabs) was used to amplify
166 the size-selected fragments in four separate rounds of PCR, each using twelve cycles (cycle
167 conditions: 98°C for 30s; 12 cycles of 98°C for 10s, 60°C for 30s, 72°C for 10s; 72°C for 5min).
168 The four PCRs were pooled and cleaned with AMPure XP beads (Agilent). The quality of the
169 final ddRAD-seq library was evaluated by visualizing DNA on a gel and quantifying it with a
170 Qubit Fluorometer 2.0 (Invitrogen). Four libraries, each of which contained 96 barcoded

171 individuals, were sent to the University of Oregon Genomics Core Facility for 100bp single-end
172 Illumina HiSeq 2000 sequencing.

173 *A note on terminology*

174 We conducted several analyses that resulted in 100bp haplotypes, with each haplotype
175 containing at least one SNP, distributed across the genome. Throughout the rest of the
176 manuscript, we will refer to haplotypes derived from an analysis of both sdRAD-seq and
177 ddRAD-seq sequencing reads together as RAD loci, and will refer to this analysis in general as
178 the “combined RAD-seq dataset”, with “sdRAD Together” and “ddRAD Together” specifying
179 individuals in this dataset. Alternatively, sdRAD loci and ddRAD loci are the haplotypes derived
180 from a separate analysis of the sdRAD-seq or ddRAD-seq sequencing reads, respectively. When
181 referring to individuals analyzed separately, they will be labeled as “sdRAD Separate” and
182 “ddRAD Separate”. RAD loci, sdRAD loci, and ddRAD loci all can contain one or more single
183 nucleotide polymorphisms (SNPs), which we will refer to RAD SNPs, sdSNPs, and ddSNPs
184 throughout the manuscript.

185 *Aligning raw reads*

186 The raw reads from each sequencing run were separated by barcode using the
187 process_radtags module of Stacks (Catchen *et al.* 2011; Catchen *et al.* 2013) and each
188 individual’s reads were aligned to the Gulf pipefish genome (Small *et al.* 2016) using Bowtie 2.0
189 (Langmead & Salzberg 2012) with the --sensitive parameters. The genome contains 22 major
190 linkage groups, corresponding to the 22 chromosomes of *S. scovelli*. Approximately 87% of the
191 genome assembly is arranged on these linkage groups. The remaining genomic data are
192 assembled into 1574 scaffolds that have not yet been localized to a chromosome (Small *et al.*
193 2016).

194 *Treating the two datasets as one: genotyping and analysis*

195 Both the ddRAD-seq and sdRAD-seq library preparations used *Pst*I, so the two libraries
196 should share many loci. Therefore, we began by treating them as a single dataset. We used the
197 `ref_map.pl` module in Stacks (Catchen *et al.* 2011; Catchen *et al.* 2013) to identify RAD loci
198 from the aligned reads. For RAD loci to be assembled, we required a minimum of 3 raw reads (-
199 `m 3`), and we allowed 2 mismatches when generating the catalog of RAD loci (`cstacks -n 2`). We
200 then ran the `populations` module in Stacks (Catchen *et al.* 2013), requiring a minimum allele
201 frequency of 0.05, the presence of each locus in at least 50% of the individuals, and that each
202 locus be present in males, females and offspring. We subsequently randomly chose one SNP per
203 RAD locus.

204 Using the resulting `vcf` file, we compared the coverage statistics between sdRAD-seq and
205 ddRAD-seq individuals using Wilcoxon rank-sum tests. Specifically, we calculated per-
206 individual means and variances for each SNP in addition to the total number of reads per
207 individual. Both allelic dropout and PCR duplicates result in uneven coverage of the two alleles
208 at a locus (Andrews *et al.* 2016; Arnold *et al.* 2013; Davey *et al.* 2013; Gautier *et al.* 2013), so
209 we compared the coverage of the reference and alternative alleles for each library preparation
210 method. Specifically, at each SNP we calculated the proportion of reads that belonged to the
211 reference allele in heterozygotes (focusing on heterozygotes controlled for different allele
212 frequencies across SNPs). Larger values therefore represent bias towards the reference allele and
213 smaller values represent bias towards the alternative allele. To determine which loci might have
214 the most extreme coverage, we calculated the mean and variance in coverage for all loci and
215 used those as the null distribution.

216 Another way to investigate the influence of restriction site polymorphisms is to estimate
217 the allelic dropout rates per SNP. We used GBStools (Cooke *et al.* 2016) to estimate SNPs with
218 high rates of restriction site allelic dropout. After adding *MboI* cut sites to the GBStools scripts,
219 we used GBStools to digest the *S. scovelli* genome (Small *et al.* 2016) using both *PstI* and *MboI*
220 (to estimate allelic dropout in the ddRAD dataset) and using only *PstI* (to estimate allelic dropout
221 in the sdRAD dataset). We then used GBStools and python v. 2.7 to estimate allelic dropout in
222 the combined RAD-seq dataset using normalization factors of 1.0.

223 To identify whether signatures of population structure emerged between the sdRAD-seq
224 and ddRAD-seq individuals, we conducted a principal components analysis of population
225 structure using PCAdapt (Luu & Blum 2017). We also calculated F_{ST} values between the
226 sdRAD-seq group and the ddRAD-seq group using using functions in gwscaR
227 (<https://github.com/spflanagan/gwscaR>). For those calculations, $F_{ST} = \frac{H_T - H_S}{H_T}$, where $H_T = 2\bar{p}\bar{q}$
228 and $H_S = \frac{1}{n} \sum_{i=1}^n 2p_i q_i$. In these formulas, the observed allele frequencies are denoted as p and q ,
229 with \bar{p} and \bar{q} representing the mean values across n groups (for this analysis, $n = 2$). Thus, H_T is
230 the expected heterozygosity among populations and H_S is the average expected heterozygosity
231 within populations (Nei 1986; Wright 1943). We also evaluated how imposing a coverage filter
232 (loci with average per-individual coverage between 3x and 50x) impacted the analysis. These
233 analyses were done in R version 3.3.1 (R Core Team 2017).

234 *Treating the two datasets separately: generating two stacks catalogs*

235 To better understand each dataset on its own, we analyzed the ddRAD-seq sequences and
236 the sdRAD-seq sequences separately in two separate runs of ref_map.pl in Stacks (Catchen *et al.*
237 2011; Catchen *et al.* 2013) and then ran populations once for each dataset to generate vcf files
238 using three populations: males, females, and offspring. The same parameter settings were used as

239 above (minimum stack depth of 3, 2 mismatches allowed, minimum allele frequency of 0.05, loci
240 present in 50% of individuals). We then repeated the analyses described above: compared
241 coverage per individual and coverage per locus, calculated our allelic imbalance metric,
242 estimated restriction site polymorphism using GBStools (Cooke *et al.* 2016), conducted a
243 principal components analysis with PCAdapt (Luu & Blum 2017), and compared allele
244 frequencies using F_{ST} .

245 In addition to comparing the two different library preparation methods to each other, we
246 compared the assembly methods (whether the ddRAD-seq and sdRAD-seq reads were analyzed
247 together or separately in Stacks). These comparisons were done using ANOVA on coverage
248 statistics and F_{ST} values, with the statistic as the response variable and the library preparation and
249 assembly methods as explanatory variables. To further investigate the relationship between
250 variation in coverage and F_{ST} values, we binned loci into six coverage categories (3-5x, 5-10x,
251 10-20x, 20-30x, 30-50x, and >50x) and used the R package lattice (Sarkar 2008) to visualize the
252 mean F_{ST} values for loci in each category.

253 The number of individuals sequenced using sdRAD-seq was much smaller than the
254 number sequenced using ddRAD-seq. To ensure the patterns we observed in our F_{ST} results were
255 not due to sample size, we randomly chose 60 individuals from the ddRAD-seq dataset to
256 compare to the 60 sdRAD-seq individuals. Additionally, we compared 60 ddRAD-seq
257 individuals to a different set of 60 ddRAD-seq individuals as a control. This analysis was done
258 using the sdSNPs and ddSNPs generated from the separate analyses, and F_{ST} values were
259 compared using ANOVAs where the sample size or the comparison (sdRAD to ddRAD or
260 ddRAD to ddRAD) and the analysis approach (separate filtered; separate unfiltered; together
261 filtered; or together unfiltered) were explanatory variables.

262 *Impact of library preparation on selection components analysis*

263 To fully assess the impact of different RAD-seq library preparation methods on the
264 results of an empirical study, we performed an F_{ST} -based selection components analysis.
265 Selection components analysis identifies signatures of selection on the genome by comparing
266 allele frequencies in individuals from a single population at different life history stages
267 (Christiansen & Frydenberg 1973; Flanagan & Jones 2015; Monnahan *et al.* 2015). In a previous
268 analysis of the ddRAD-seq individuals from this study, we demonstrated that signatures of sex-
269 biased viability selection and sexual selection are distributed across the genome in *S. scovelli*,
270 and that more loci show signatures of sex-biased viability selection than sexual selection
271 (Flanagan & Jones 2017). In that analysis, we inferred maternal alleles from the 130 father-
272 offspring combinations and compared the inferred maternal alleles to the females collected in the
273 population to identify putative signatures of sexual selection. Collected adult male and adult
274 female allele frequencies were compared to identify sex-biased viability selection (Flanagan &
275 Jones 2017).

276 Because the sdRAD-seq individuals do not include any offspring, we could not repeat the
277 sexual selection component of the analysis for sdRAD-seq individuals. However, we used the
278 combined RAD-seq dataset (ddRAD individuals and sdRAD individuals analyzed together) and
279 evaluated the results in comparison to an analysis based only on the ddRAD-seq data.
280 Additionally, we compared allele frequencies in males and females in all three datasets (RAD-
281 seq, ddRAD-seq, and sdRAD-seq).

282 To perform the selection components analysis, we used the same procedure as described
283 in Flanagan & Jones (2017), converted into R code, which we have made available in a package
284 called gwscaR (<https://github.com/spflanagan/gwscaR>). For the sexual selection analysis, we

285 first inferred maternal alleles by subtracting the paternal allele from each offspring's genotype.
286 We then calculated F_{ST} values between the inferred maternal alleles and the collected females,
287 using the same methods described above. The value of $2NF_{ST}(k - 1)$ has a χ^2 distribution with $(k$
288 $- 1)(n - 1)$ degrees of freedom, where k is the number of alleles, N is the total number of
289 individuals, and n is the number of populations sampled (Waples 1987; Workman & Niswander
290 1970). We applied this calculation to every locus in our analysis to calculate p -values and then
291 applied the Benjamini and Hochberg (1995) false discovery rate to identify significant loci at the
292 level of $\alpha = 0.05$.

293 This analysis was performed in the ddRAD-seq dataset with 130 father-offspring
294 combinations and in the combined RAD-seq dataset with 153 father-offspring combinations. The
295 comparison of males and females involved 57 females and 159 males in the ddRAD-seq dataset,
296 30 males and 30 females in the sdRAD-seq dataset, and 87 females and 189 males in the RAD-
297 seq dataset. In each of the comparisons, we used a single SNP from each RAD locus, and each
298 locus was required to be present in at least 50% of the individuals in each group. We also
299 retained only SNPs with a minor allele frequency of at least 0.05 and with an average coverage
300 value between 5x and 20x. For each of the two selection components, we used ANOVA to
301 compare the F_{ST} values using the type of analysis method as the explanatory variable.

302 *Comparing the results to samtools*

303 Bias resulting from the Stacks analysis may have been a result not of the sequencing
304 methods *per se*, but rather an artifact of the Stacks pipeline. Therefore, we used samtools (Li *et*
305 *al.* 2009; Li 2011) to create consensus loci using both the 60 sdRAD individuals and the subset
306 of 60 ddRAD individuals. We subsequently used bcftools to call variant SNP sites. In calling
307 SNPs, we required that 50% of the individuals had data for a particular SNP (-d 0.5) and

308 excluded sites where all samples were phased with the phase bit set at 0.05 (-p 0.05 -P full).
309 Using vcftools (Danecek *et al.* 2011), we further filtered SNPs to remove indels and retain
310 biallelic SNPs with a minor allele frequency of at least 0.05, and in R (R Core Team 2017) we
311 removed loci with an average per-individual coverage below 3 reads per locus. To identify
312 whether the quality scores were meaningful in the context of reducing bias, we also filtered the
313 dataset to include only loci with a quality score ≥ 30 . We used the same samtools-bcftools-
314 vcftools pipeline on the 60 sdRAD individuals and the subset of 60 ddRAD individuals
315 separately. Shared loci between these two separate assemblies were identified using custom
316 scripts in R (R Core Team 2017).

317 To identify whether similar patterns emerged using the samtools assembly as in the
318 Stacks assembly, we calculated per-locus coverage statistics for the samtools datasets and the
319 quality-filtered samtools datasets. We also calculated F_{ST} values between ddSNPs and sdSNPs in
320 four different datasets: (1) analyzed together by samtools; (2) analyzed together by samtools and
321 filtered based on quality scores; (3) analyzed separately by samtools; and (4) analyzed separately
322 by samtools and filtered based on quality scores.

323 *In silico digestion of the reference genome*

324 To model the impact of the different sources of error, we wrote a C++ program
325 (https://github.com/spflanagan/SCA/tree/master/programs/insilico_radseq) to perform an *in*
326 *silico* digestion of the reference genome sequences and model shearing bias, polymorphic
327 restriction sites, PCR bias, and uneven coverage. The program performs both a single-digest and
328 a double-digest of the reference. When modeling polymorphic restriction sites, we used an
329 approach similar to that of Gautier *et al.* (2013). We assumed that 10% of the restriction sites
330 would be constant, and the constant loci were chosen randomly. The loci that were polymorphic

331 had an expected proportion of nucleotides with a segregating mutation of $\theta = 4N_e\mu$, and θ was
332 the same for both the single digestion and double digestion but each locus had its own μ . We
333 altered the value of θ by changing both the effective population size (N_e) to be 5,000, 10,000,
334 and 20,000, and by drawing μ from a uniform distribution of either $[10^{-9}, 10^{-8}]$ or $[10^{-8}, 10^{-7}]$.

335 This program parses the fasta file containing the reference genome and finds the
336 restriction enzyme recognition sites. When conducting a single digest, the length of the fragment
337 determines how many sheared fragments are generated. Because the sdRAD-seq library
338 preparation sheared to an average fragment size of 500bp, the fragment was sheared s times
339 based on the length of the fragment, l , at s random locations on the fragment, according to the
340 formula: $s = l/500$. The resulting RAD loci from either end of the fragment (any extra sheared
341 parts from the middle of the fragment were discarded) were kept if their length was between 250
342 and 700 bp. If shearing bias was modeled, as each fragment was generated, it was only kept if
343 adding that fragment to the set of processed fragments maintained an average fragment length
344 above 500bp, biasing the shearing towards longer fragments. To model the double digestion,
345 only fragments with both restriction enzyme sites that were 250-700bp in length were kept.

346 Once the *in silico* digestion was complete, a simulated population needed to be sampled
347 at both sdRAD and ddRAD loci generated by the *in silico* digestion. Each locus was given a
348 population-level allele frequency, either a random uniformly distributed number in the range of
349 $[0,1)$ or drawn from a normal distribution centered around 0.8 with a standard deviation of 0.13
350 ('skewed'). This skewed distribution used the mean and standard deviation of the SNPs from our
351 combined dataset. If the locus was shared between the single and double digest, it had the same
352 population-level allele frequency in both. We modeled one biallelic SNP per locus, and
353 individuals were randomly assigned genotypes based on the population-level allele frequencies.

354 We then determined whether polymorphic restriction sites or PCR duplication affected the
355 genotype at each locus for each individual. If the locus could have a polymorphic restriction site
356 (i.e., it was not one of the 10% of sites that were constant), a number was drawn from a Poisson
357 distribution with a mean $\theta_i * l_{RS}$, where θ_i is the proportion of nucleotides with a mutation for
358 locus i (see above) and l_{RS} is the length of the restriction site (6 for *PstI* and 4 for *MboI*). If the
359 Poisson distribution returned a 1, then one of the alleles was randomly chosen to be dropped and
360 the genotype for that locus became homozygous for the selected allele. If the Poisson distribution
361 returned 2 or higher, then both alleles were dropped and the locus was missing for that
362 individual. For the individuals sampled at the *in silico* ddRAD loci, the locus was evaluated
363 twice in this manner since either restriction site could be polymorphic.

364 To model PCR duplication events, a Poisson distribution was used with a mean
365 representing the percentage of reads per PCR cycle that would be duplicated multiplied by the
366 number of PCR cycles. This approach assumed that more PCR cycles would result in higher
367 duplication rates. If the Poisson distribution returned a 1 or higher, one of the two alleles was
368 randomly chosen as the duplicated allele and replaced the genotype at the second allele. We
369 varied the PCR duplication rate from 0 to 5% per cycle, and the number of cycles was 12 for the
370 ddRAD loci and 20 for the sdRAD loci, mirroring the number of cycles we used in the library
371 preparation steps (see above).

372 Once the genotypes were assigned and affected (or not) by polymorphic restriction sites
373 and PCR duplications, we calculated F_{ST} at the loci shared by the single and double digests
374 between the individuals sampled by *in silico* ddRAD and *in silico* sdRAD. We sampled a total of
375 400 individuals, and either had a symmetric sampling scheme ($n_{sd} = n_{dd} = 200$) or asymmetric
376 ($n_{sd} = 60, n_{dd} = 340$). F_{ST} was calculated as $F_{ST} = (H_T - H_S)/H_T$, where H_S is the weighted

377 average expected heterozygosity in each sub-population and H_T is the expected heterozygosity in
378 the entire population. We ran the *in silico* digestion with different restriction site mutation rates,
379 PCR duplication rates, and population-level allele frequencies and compared the resulting F_{ST}
380 values to each other and to the observed values from the empirical library preparation.

381 **Results**

382 *Assembly statistics*

383 The analysis treating sdRAD-seq and ddRAD-seq reads together included data from 444
384 individuals, 60 of which were prepared using the sdRAD-seq method. After the pruning imposed
385 by the populations module of Stacks (minor allele frequency ≥ 0.05 , SNPs present in 50% of
386 females, males, and offspring; Catchen *et al.* 2013), 84,851 SNPs from 36,007 RAD loci were
387 retained. The analysis of only the 60 sdRAD-seq individuals resulted in 250,425 sdSNPs from
388 115,708 sdRAD loci. The ddRAD-seq dataset contained 69,109 ddSNPs from 31,956 ddRAD
389 loci. The sdRAD and ddRAD datasets shared 49,893 SNPs on 23,396 RAD loci.

390 *Differences in coverage when assembled together versus separately*

391 All of the coverage data were skewed, so we used the natural log to transform the
392 following coverage metrics. Total coverage was affected by the interaction of the library
393 preparation method (sdRAD-seq vs ddRAD-seq) and whether the data were analyzed together or
394 separately ($F_{1,169977} = 304.2$, $p = 2 \times 10^{-16}$). The sdRAD-seq individuals had more reads than
395 ddRAD-seq individuals when they were analyzed separately (TukeyHSD $p < 0.0001$) but not
396 together (TukeyHSD $p = 0.58$; Fig. 1). When the sdRAD-seq reads were analyzed together with
397 the ddRAD-seq reads, the average number of reads per individual was lower than coverage for
398 sdRAD individuals when the two library preparation methods were analyzed separately

399 (TukeyHSD $p < 0.0001$; Fig. 1). The ddRAD-seq individuals had the same coverage regardless
400 of whether they were analyzed together or separately (TukeyHSD $p = 0.16$; Fig. 1).

401 The log transformed per-SNP, per-individual coverage was higher in ddRAD than
402 sdRAD when analyzed separately (TukeyHSD $p < 0.0001$) and when analyzed together
403 (TukeyHSD $p < 0.0001$). When sdRAD-seq reads and ddRAD-seq reads were analyzed together,
404 sdSNPs had higher coverage than sdSNPs from the separate analysis (TukeyHSD $p < 0.0001$),
405 but ddRAD showed no difference (Tukey HSD $p = 0.9996$; Fig. 1).

406 One indication that loci may be biased due to restriction site polymorphism or PCR
407 duplicates is if one allele has higher coverage than the other allele at a given locus. We found
408 that both library preparation methods and analysis approach impacted the proportion of reference
409 reads in heterozygotes ($F_{1,211838} = 20.78$, $p = 5.16 \times 10^{-6}$). The ddRAD-seq individuals had higher
410 proportions than the sdRAD-seq individuals when analyzed together (TukeyHSD $p < 0.0001$) but
411 not when analyzed separately (TukeyHSD $p = 0.0685$). The proportion of reference reads in
412 heterozygotes was lower in ddRAD when analyzed separately from sdRAD (TukeyHSD $p <$
413 0.0001), whereas sdRAD proportions were consistent regardless of analysis method (TukeyHSD
414 $p = 0.8934$; Table 1).

415 The GBStools analysis estimated the number of dropped alleles per individual per SNP.
416 The estimated dropped allele count was influenced by both the library preparation (ddRAD-seq
417 vs sdRAD-seq) and the analysis method ($F_{1,101116} = 85.93$, $p < 2 \times 10^{-16}$; Table 1). The sdRAD-
418 seq library displayed more evidence for dropped alleles than ddRAD-seq library when they were
419 analyzed alone (Tukey HSD, $p < 0.0001$) or together (Tukey HSD, $p < 0.0001$). The ddRAD-seq
420 showed more dropouts when analyzed together than alone (Tukey HSD, $p < 0.0001$), and we saw
421 the same pattern for sdRAD-seq (TukeyHSD, $p < 0.0001$).

422 PCR duplicates are expected to result in variance in coverage depth at a locus (Andrews
423 et al. 2016). Increasing the number of PCR cycles will increase the number of PCR duplicates in
424 the sequencing library, so we expected that the sdRAD-seq dataset would suffer more from the
425 problem of PCR duplicates. We compared the average variance in coverage across all SNPs
426 between the two library preparation methods and between the two analysis approaches. We log-
427 transformed the variance in coverage for each SNP and found that the library preparation method
428 and the analysis method interact to influence variance in coverage ($F_{1,169977} = 474.5, p < 2 \times 10^{-16}$;
429 Table 1). The sdSNPs had higher variance in coverage than ddSNPs both when analyzed
430 together (TukeyHSD $p < 0.0001$) and separately (TukeyHSD $p < 0.0001$). Within the sdSNPs,
431 the variance in coverage was higher when assembled together than alone (TukeyHSD $p <$
432 0.0001), whereas ddSNPs had higher variance in coverage when assembled alone (TukeyHSD p
433 < 0.0001).

434 *Observed heterozygosity*

435 Observed heterozygosity is expected to be affected by polymorphic restriction sites,
436 leading to decreased observed heterozygosity at some loci. In contrast to our expectations, the
437 ddRAD-seq dataset had higher proportions of heterozygotes than sdRAD-seq regardless of
438 whether they were analyzed together (TukeyHSD $p < 0.0001$) or separately (TukeyHSD $p <$
439 0.0001), although the interaction term was significant ($F_{1,169977} = 128.10, p < 2 \times 10^{-16}$; Table 1),
440 indicating that the library preparation method impacted heterozygosity in different ways
441 depending on the way the data were analyzed. The subset of 60 ddRAD individuals used in the
442 F_{ST} analysis also had a higher mean proportion of heterozygotes than the sdRAD individuals
443 (Wilcoxon signed-rank test $W = 1033200000, p < 2.2 \times 10^{-16}$).

444 *Analysis of population structure using principal components analysis*

445 When we applied the principal components approach of PCAdapt (Luu & Blum 2017) to
446 identify population structure in the dataset with ddRAD-seq and sdRAD-seq individuals
447 combined, we found that 24.6% of the variation was explained by library preparation method
448 (Fig. 2). Alternatively, in the analysis of the separate ddRAD-seq and sdRAD-seq datasets,
449 individuals did not sort based on library preparation method, and the first axis of variation only
450 explained 6.5% of the variation, suggesting that differences between the two sets of individuals
451 were small.

452 *Comparison of allele frequencies in ddRAD-seq and sdRAD-seq*

453 We compared allele frequencies between the ddRAD-seq and sdRAD-seq individuals in
454 two ways: when they were analyzed separately (as if comparing results from separate studies)
455 and when they were analyzed together (treating them as if they were one dataset). When the
456 individuals were analyzed together, many more loci were fixed for one allele or the other in both
457 the ddRAD-seq and the sdRAD-seq datasets, but the overall distributions of allele frequencies
458 were similar between ddRAD individuals and sdRAD individuals (Supplemental Fig. 1). The
459 assembly method significantly impacted F_{ST} values between ddRAD and sdRAD individuals
460 ($F_{1,74171} = 228.3, p < 2 \times 10^{-16}$; Fig. 3, Supplemental Fig. 2).

461 When the ddRAD and sdRAD individuals were analyzed separately (14,324 shared
462 SNPs), the mean major allele frequency was significantly higher in sdSNPs ($\mu = 0.7901$) than in
463 ddSNPs ($\mu = 0.7888$; one-sided paired Wilcoxon signed rank test $V = 49279000, p = 0.00015$).
464 This pattern was reflected in the F_{ST} values, which had a mean of 0.00307 and ranged from 0 to
465 0.4883 (Fig. 3, Supplemental Fig. 2). When the two sets of individuals were analyzed together
466 (27,334 SNPs), the mean major allele frequency was higher in ddRAD individuals than sdRAD

467 individuals ($\mu_{\text{ddRAD}} = 0.8211$, $\mu_{\text{sdRAD}} = 0.7038$; one-sided paired Wilcoxon signed rank test $V =$
468 379090000 , $p < 2.2 \times 10^{-16}$), and mean F_{ST} was 0.00604 and ranged from -1.6309 to 0.9576. This
469 significantly higher mean (TukeyHSD $p < 0.0001$) did not include the 4,440 SNPs that were
470 fixed for different alleles in the combined analysis. Any similar sites in the combined analysis
471 would not have been polymorphic in at least one of the separate datasets and so those loci were
472 not retained.

473 One approach to ameliorate the bias of RAD-seq, particularly the bias due to PCR
474 duplicates, is to remove SNPs from the analysis with high per-SNP, per-individual coverage
475 (Schweyen *et al.* 2014). Therefore, we imposed a filter to retain SNPs with an average coverage
476 between 3x and 50x on both datasets, and this filter significantly impacted the F_{ST} values ($F_{1,74171}$
477 $= 292.2$, $p < 2 \times 10^{-16}$; Fig. 3, Supplemental Fig. 2), though its effect interacted with the analysis
478 approach ($F_{1,74171} = 133.7$, $p < 2 \times 10^{-16}$). In the comparison of allele frequencies at sdSNPs and
479 ddSNPs generated from separate analyses, the filter removed 4,363 SNPs (9,961 SNPs were
480 retained) and changed the mean F_{ST} to 0.00266, though this was not a significant change
481 (TukeyHSD $p = 0.94$). The filter did significantly decrease the mean F_{ST} to -0.0033 for the
482 analysis of the SNPs together (TukeyHSD $p < 0.0001$). Filtering for coverage removed 16,778
483 loci from the combined analysis, with 10,556 SNPs retained. The filter removed most of the
484 SNPs with fixed alleles between sdRAD and ddRAD, with only 301 such SNPs remaining.
485 Keeping loci with coverage between 3x and 50x in both the separate and combined analyses
486 removed any difference in the mean between the two sets of F_{ST} values (TukeyHSD $p = 0.637$).
487 The importance of coverage filters is emphasized when looking at a heatmap of mean F_{ST} values
488 for SNPs binned by coverage (Fig. 4). This figure shows that the most extreme F_{ST} values

489 emerge when one group has mid-range coverage (e.g., 10-20x) but the other group has
490 exceedingly high coverage (>50x).

491 To test for the effects of the number of sampled individuals, we also compared allele
492 frequencies between all 60 sdRAD individuals and 60 randomly selected ddRAD individuals and
493 between 60 ddRAD individuals and a separate randomly chosen set of 60 ddRAD individuals.
494 For each of these comparisons, we performed the same four F_{ST} comparisons as above (analyzed
495 together; analyzed together with a coverage filter; analyzed separately; and analyzed separately
496 with a coverage filter). We found that sampling fewer individuals in the ddRAD-seq dataset
497 resulted in higher F_{ST} values ($F_{1,121180} = 4.997$, $p = 0.0254$; Fig. 3, Supplemental Fig. 2),
498 although the type of analysis interacted with the effect of sample size ($F_{3,121180} = 6.298$, $p =$
499 0.0003 ; Fig. 3, Supplemental Fig. 2). The comparison of 60 ddRAD-seq individuals to 60 other
500 ddRAD-seq individuals resulted in overall lower F_{ST} values than the corresponding comparisons
501 of sdRAD-seq and ddRAD-seq individuals ($F_{1,179742} = 13.04$, $p = 0.0003$; Fig. 3, Supplemental
502 Fig. 2), although the analysis type interacted with the effect of whether ddRAD individuals were
503 being compared to sdRAD or other ddRAD individuals ($F_{3,179742} = 15.72$, $p = 3.21 \times 10^{-10}$; Fig. 3,
504 Supplemental Fig. 2).

505 *Differences in the outcome of selection components analysis*

506 For the selection components analysis, we filtered loci to retain only those with coverage
507 between 5x and 20x. After the filtering step, we compared allele frequencies in males and
508 females using 28,230 SNPs, 48,743 ddSNPs, and 92,710 sdSNPs. The comparison of maternal
509 alleles to collected females could only be conducted in the combined RAD dataset and the
510 ddRAD dataset, since no offspring were genotyped using sdRAD-seq. The inference of maternal
511 alleles reduced the number of loci used to 16,099 SNPs and 35,666 ddSNPs, because not all

512 father-offspring genotype combinations facilitate inference of the maternal allele (Flanagan &
513 Jones 2017). However, it is worth noting that the inference of maternal alleles does not bias
514 allele frequencies unless error rates are high (Flanagan & Jones 2017).

515 The comparison of allele frequencies between males and females to test for sex-biased
516 viability selection yielded dramatically different results depending on which sequencing and
517 analysis methods were used. The males-females analysis using sdRAD-seq had higher F_{ST} values
518 than the equivalent comparison in the ddRAD-seq analysis (ANOVA $F_{2,169680} = 4567$, $p < 2 \times 10^{-16}$;
519 Fig. 5, Supplemental Fig. 3) and higher values than the analysis of both ddRAD-seq and
520 sdRAD-seq individuals together (TukeyHSD $p < 2 \times 10^{-16}$), although the combined analysis had
521 higher F_{ST} values than the ddRAD comparison. The combined analysis also identified 4,315
522 significant SNPs after correcting for multiple comparisons whereas the ddRAD analysis
523 identified only 58 significant ddSNPs (Fig. 5, Supplemental Fig. 3). Note that this number is
524 different from the number identified in Flanagan & Jones (2017) because our previous analysis
525 used slightly more stringent filtering methods than those used here.

526 To ensure that the differences between the ddRAD-seq and sdRAD-seq selection
527 components analysis comparing males and females was not simply driven by the different
528 sample sizes, we randomly sampled 30 males and 30 females from the ddRAD-seq dataset and
529 compared their allele frequencies. The smaller sample size yielded significantly higher F_{ST}
530 values than the full ddRAD-seq dataset (one-sided Wilcoxon signed-rank test, $W = 1435300000$,
531 $p < 2.2 \times 10^{-16}$), but the F_{ST} values were still significantly lower than the sdRAD-seq dataset (one-
532 sided Wilcoxon signed-rank test, $W = 1967400000$, $p < 2.2 \times 10^{-16}$).

533 The test for sexual selection yielded similar results to that of males versus females.
534 Again, the combined analysis had many more significant SNPs. The selection components

535 analysis of the combined dataset identified 125 significant SNPs after Benjamini and Hochberg
536 (1995) false discovery rate correction, whereas only 16 ddSNPs were significant in the ddRAD
537 selection components analysis. However, the ddRAD-seq analysis had a higher mean F_{ST} value
538 than the combined analysis in this case (ANOVA $F_{1,51763} = 8.668$, $p = 0.00324$; Fig. 5), although
539 the range of F_{ST} values was larger in the combined analysis (0.0000 – 0.2915 in combined,
540 0.0000 – 0.1966 in ddRAD).

541 *Comparison to samtools analysis*

542 To verify that the results were due primarily to differences in the underlying data rather
543 than to artifacts in the analysis methods, we re-analyzed the sdRAD individuals and the subset of
544 60 ddRAD individuals with samtools, bcftools, and vcftools. The samtools analysis of sdRAD
545 and ddRAD individuals together resulted in 133,946 variant SNPs, after filtering for coverage.
546 The samtools analysis of sdRAD individuals separately resulted in 232,101 variant SNPs and the
547 samtools analysis of ddRAD individuals separately yielded 63,934 variant SNPs, with 53,057
548 shared variant SNPs between the two analyses.

549 These samtools results share some similarities with the Stacks analysis of the sdRAD-seq
550 individuals and a subset of 60 of the ddRAD-seq individuals. Average coverage depth per locus
551 per individual (total depth at the locus/the number of genotyped individuals) was determined by
552 an interaction between library preparation method (ddRAD-seq vs sdRAD-seq) and analysis
553 approach (analyzed together or separately; $F_{1,563923} = 1526.5$, $p < 2 \times 10^{-16}$), with both sdSNPs
554 and ddSNPs analyzed together having lower average coverage than when analyzed separately
555 (sdRAD TukeyHSD $p < 0.0001$; ddRAD TukeyHSD $p < 0.0001$). When analyzed separately, the
556 sdRAD-seq dataset had higher coverage than the ddRAD-seq dataset (TukeyHSD $p < 0.0001$;
557 Table 2). Similar to Stacks, when analyzed separately, the ddRAD-seq dataset had higher

558 coverage. However, in samtools the average coverage was lower when the datasets were
559 analyzed together, whereas coverage increased for sdRAD but not ddRAD in the Stacks analysis.
560 Note that samtools output locus-wide coverage statistics instead of per-locus, per-individual
561 coverage statistics, so we were unable to detect such tradeoffs in the samtools dataset.

562 The proportion of heterozygotes in the samtools analysis was determined by the
563 interaction between library preparation method and whether the data were analyzed together or
564 separately ($F_{1,563923} = 24.667$, $p < 2 \times 10^{-16}$), similar to the Stacks analysis. When analyzed
565 separately, the ddRAD-seq dataset had a higher proportion of heterozygotes than the sdRAD-seq
566 dataset (TukeyHSD $p < 0.0001$). The sdRAD-seq dataset had a higher proportion of
567 heterozygotes than when it was analyzed with the ddRAD-seq data (TukeyHSD $p < 0.0001$),
568 whereas the ddRAD-seq dataset had a higher proportion of heterozygotes when it was analyzed
569 without the sdRAD individuals (TukeyHSD $p < 0.0001$). These patterns are consistent with the
570 overall analysis with Stacks (Table 1) and with the analysis of the Stacks dataset with a subset of
571 60 ddRAD individuals (Table 2).

572 The samtools analysis also resulted in large F_{ST} values between the two datasets. We
573 compared F_{ST} values between ddSNPs and sdSNPs with a basic coverage filter (equivalent to the
574 baseline filtering done in the Stacks analysis), but we also used the samtools quality scores to
575 impose an additional filter. F_{ST} values between datasets were dependent on both the analysis
576 method and whether the additional quality filter was imposed ($F_{1,360635} = 211.4$, $p < 2 \times 10^{-16}$).
577 Without the filter, the F_{ST} values were not significantly different when the datasets were
578 analyzed together or separately (TukeyHSD $p = 0.6053$), but after including the quality filter, the
579 separate analysis had lower F_{ST} values than the analysis of the data together (TukeyHSD $p <$

580 0.0001). For both analysis approaches, including the quality filter reduced the F_{ST} values
581 (separate TukeyHSD $p < 0.0001$; together TukeyHSD, $p < 0.0001$).

582 *Analysis of the in silico digestion*

583 The *in silico* digestions of the reference genome demonstrated the importance of PCR
584 duplications, restriction site polymorphism, shearing bias, mean allele frequency skew, and
585 asymmetric sampling schemes for allele frequency estimation in studies employing ddRAD-seq
586 or sdRAD-seq. We began by isolating one or two sources of bias at a time. Both the percentage
587 of reads resulting from PCR duplicates ($F_{1,84148} = 38.4$, $p < 5.82 \times 10^{-10}$) and whether the allele
588 frequencies were skewed towards large major allele frequencies affected F_{ST} values, although the
589 effect of a skewed allele frequency spectrum had a more dramatic effect ($F_{1,84148} = 2056.9$, $p =$
590 2×10^{-16}) and the interaction was not significant ($F_{1,84148} = 0.019$, $p = 0.89$; Table 3).

591 Unsurprisingly, higher rates of polymorphic restriction sites ($\theta = 4N_e\mu$) impacted the
592 differentiation between sdRAD and ddRAD loci ($F_{1,42015} = 18.5$, $p = 1.7 \times 10^{-5}$), although this
593 effect was primarily driven by altering the N_e value ($F_{1,42013} = 40.963$, $p = 1.57 \times 10^{-10}$) rather than
594 the mutational distribution ($F_{1,42013} = 1.096$, $p = 0.295$; Table 3).

595 One source of bias that could create differences between sdRAD loci and ddRAD loci is
596 shearing bias. However, shearing bias did not significantly affect the distribution of F_{ST} values
597 ($F_{1,32658} = 0.179$, $p = 0.671898$). More important factors were the symmetry of the sampling
598 scheme ($F_{1,32658} = 212.879$, $p < 2 \times 10^{-16}$) and whether the average allele frequency was set at 0.8
599 instead of 0.5 ($F_{1,32658} = 836.636$, $p < 2 \times 10^{-16}$), two factors which had a significant interaction
600 ($F_{1,32658} = 11.528$, $p = 0.000686$; Table 3).

601 In an actual study, all of these sources of bias are expected to be present. Therefore, we
602 ran the *in silico* digestion with all of the biases, and changed one bias at a time to see how

603 different parameters interacted in a complex system of bias. The variables that increased the F_{ST}
604 values the most were skewed allele frequencies ($t = 11.756, p < 2 \times 10^{-16}$) and the symmetry of
605 the sampling scheme ($t = 5.679, p = 1.37 \times 10^{-8}$; Table 3). Although shearing bias alone did not
606 inflate F_{ST} values (see above), when combined with other factors shearing bias had a significant
607 effect on F_{ST} ($t = 2.037, p = 0.0417$; Table 3).

608 **Discussion**

609 Here we present a case study, which involves sampling a single population using both
610 sdRAD-seq and ddRAD-seq. This analysis provides a unique empirical opportunity to
611 investigate sources of bias in two different RAD-seq methods. Pairing our empirical analysis
612 with an *in silico* digestion of the Gulf pipefish genome allows us to assess the importance of
613 factors such as shearing bias, polymorphic restriction sites, PCR duplicates, allele frequencies,
614 and sampling schemes on differentiation between individuals genotyped using sdRAD-seq and
615 those genotyped by ddRAD-seq.

616 The analysis approach played an important role in the outcomes of the data analysis at
617 every step, from the coverage of loci, to restriction site dropout, to differences between ddRAD-
618 seq and sdRAD-seq allele frequencies. We analyzed the datasets using both Stacks and samtools,
619 and identified similar trends with a few small differences. In both analyses, the sdRAD-seq
620 dataset had higher per-individual, per-SNP coverage (Fig. 1, Table 2), an unsurprising result
621 since the 60 sdRAD individuals were sequenced in one lane, whereas the ddRAD individuals
622 were pooled 96 per lane. Analyzing the data with both Stacks and samtools resulted in a higher
623 proportion of heterozygotes in the ddRAD-seq datasets than the sdRAD-seq datasets, especially
624 when the data were treated separately. Finally, both the Stacks and samtools analyses resulted in
625 large F_{ST} values between the sdRAD-seq and ddRAD-seq individuals, suggesting that the

626 patterns we observed are due to underlying differences in the dataset, not due to bias arising from
627 differences between the analysis pipelines. In both cases, however, it is clear that more bias is
628 introduced to the analysis when ddRAD-seq and sdRAD-seq are analyzed together as one
629 dataset. For this, we can offer a clear suggestion to researchers: if dealing with data generated by
630 different methods, analyze them separately and then identify overlap between the datasets.

631 Despite the fact that all sampled Gulf pipefish came from a single population, we
632 observed significantly different allele frequencies between individuals that were genotyped using
633 sdRAD-seq and those genotyped using ddRAD-seq. If coverage filters were not in place, the F_{ST}
634 values ranged up to 0.9788 when the two types of libraries were analyzed together and 0.4883
635 when analyzed separately. These extreme values were much larger than the maximum values
636 observed between geographically distinct populations of *S. scovelli* (Flanagan *et al.* 2016). The
637 difference between genotyping methods was not simply due to the fact that we sampled 60
638 individuals using sdRAD-seq and 384 using ddRAD-seq, because a comparison of 60 randomly
639 selected ddRAD individuals to the 60 sdRAD individuals yielded F_{ST} values that were
640 substantially higher than a comparison of 60 ddRAD individuals to another 60 ddRAD
641 individuals (Fig. 3). However, skewed sample sizes did exaggerate differences between the two
642 datasets in our *in silico* digestion of the reference genome (Table 3).

643 The differences between the datasets generated by different RAD-seq methods were
644 partly a result of the differences in coverage we observed between the datasets. Our sdRAD-seq
645 dataset had higher per-individual coverage than the ddRAD-seq dataset primarily because the
646 sdRAD-seq dataset contained only 60 individuals sequenced in one Illumina lane, whereas the
647 ddRAD-seq dataset comprised data from four lanes of Illumina sequencing, each with 96
648 individuals. Similarly, the ddRAD-seq dataset had fewer SNPs (31,956 ddSNPs compared to

649 113,166 sdSNPs used in the analyses; Fig. 1). Variance in coverage and allelic dropout can result
650 from pooling of individuals right after the ligation rather than immediately before sequencing
651 (daCosta & Sorenson 2014), but since both the sdRAD-seq and ddRAD-seq individuals were
652 pooled at the same step in the present analysis, this factor is unlikely to explain the differences
653 between the two datasets. Regardless of the source of the variance in coverage, differences in
654 coverage between the two datasets appear to have driven extreme F_{ST} values (Fig. 3, 4).
655 Additionally, the results from our *in silico* digestion suggest that loci with major allele
656 frequencies skewed toward a value of one result in elevated F_{ST} (Table 3). Since F_{ST} is
657 calculated from allele frequencies, it is unsurprising that we found increased F_{ST} values between
658 methodologies when the mean allele frequency was elevated in our *in silico* digestion (see
659 Jakobsson *et al.* 2013; Jost 2008). However, our results are a reminder to researchers that inflated
660 allele frequencies due to bias in RAD-seq can yield inflated estimates of F_{ST} .

661 Polymorphic restriction sites were found to play a major role in bias in this RAD-seq
662 study. In our *in silico* digestion, increasing the rate of mutations at restriction sites significantly
663 increased F_{ST} values between sdRAD-seq and ddRAD-seq samples, although this effect was less
664 pronounced when combined with other sources of bias (Table 3). Skewed coverage of reference
665 and alternative alleles is expected because of allelic dropout, primarily due to polymorphic
666 restriction sites (Andrews *et al.* 2016; Gautier *et al.* 2013). In our sequencing data, when the
667 datasets were analyzed separately, we found that the difference in skew in coverage towards the
668 reference allele in ddRAD-seq SNPs and sdRAD-seq SNPs was less pronounced (Table 1). More
669 important than the means is the range and the variance of the skew, since the skew should occur
670 in either direction. Although the ddRAD-seq individuals had different mean proportions of
671 reference reads in heterozygotes than sdRAD-seq when analyzed together, the two did not differ

672 when analyzed separately. In fact, the sdRAD-seq dataset analyzed separately had a larger range
673 of values (0.951) compared to the ddRAD-seq (0.903; Table 1), indicating that sdRAD-seq might
674 be experiencing more allelic dropout. In addition, the GBStools analysis indicates that the
675 sdRAD-seq dataset has more dropout alleles than the ddRAD-seq dataset (Table 1). This result is
676 surprising because the ddRAD-seq dataset includes more restriction sites and is expected to be
677 more impacted by polymorphic restriction sites than sdRAD-seq (Andrews *et al.* 2016; Arnold *et*
678 *al.* 2013). Additionally, polymorphic restriction sites are expected to falsely inflate
679 homozygosity (Davey *et al.* 2013), and our ddRAD-seq dataset had higher proportions of
680 heterozygous individuals than the sdRAD-seq dataset (Table 1), a pattern which was not due to
681 the larger sample size and supports the finding that our sdRAD-seq dataset seems to be more
682 greatly affected by polymorphic restriction sites than our ddRAD-seq dataset.

683 We expected PCR duplicates to be an important source of bias in our RAD-seq studies
684 (Andrews *et al.* 2016; Davey *et al.* 2011; Hoffberg *et al.* 2016), especially since several recent
685 methods have emerged to reduce their impact (Ali *et al.* 2016; Hoffberg *et al.* 2016), including
686 the use of paired-end sequencing to identify and remove PCR duplicates from datasets
687 (Hohenlohe *et al.* 2013; Schweyen *et al.* 2014; Smith *et al.* 2014; Tin *et al.* 2015). Our *in silico*
688 digest results suggested that PCR duplicates do affect F_{ST} values, but do not have a marked
689 effect when combined with other sources of bias (Table 3). In our empirical data, we found that
690 our sdRAD-seq SNPs had higher variance in coverage than our ddRAD-seq SNPs, which we
691 expected because we used more PCR cycles in the preparation of the sdRAD-seq library. One
692 suggestion for ameliorating the bias due to PCR duplication is to remove loci with incredibly
693 high coverage (i.e., > 50X; Schweyen *et al.* 2014). When we performed this filtering step, the
694 F_{ST} values shifted towards zero, but only significantly so for the dataset containing sdRAD-seq

695 and ddRAD-seq reads analyzed together. Therefore, filtering to remove high-coverage loci can
696 remove some of the bias between sdRAD-seq and ddRAD-seq datasets but possibly only if the
697 data are pooled. Our results are therefore mixed with regard to PCR duplicates, because the *in*
698 *silico* digestion showed that they may not play a major role in causing differences between
699 sdRAD-seq and ddRAD-seq, but we found some evidence of PCR duplicates in our dataset.

700 Andrews *et al.* (2016) suggested that shearing bias would not greatly impact sdRAD-seq
701 datasets, but our *in silico* digestion suggests that shearing bias may be an important factor. We
702 found that shearing bias alone did not substantially alter allele frequencies, but it interacted with
703 other sources of bias, including asymmetric sampling schemes and skewed allele frequencies
704 (Table 3). These results highlight the importance of considering all sources of bias when
705 evaluating RAD-seq datasets, as the effects of various sources of bias may interact to alter allele
706 frequencies.

707 The bias caused by genotyping some individuals with sdRAD-seq and some with
708 ddRAD-seq strongly affected the results of selection components analysis. In the comparison of
709 males and females, the F_{ST} values were significantly larger when the ddRAD-seq and the
710 sdRAD-seq individuals were analyzed together, and a greatly inflated number of loci had
711 significant p -values after the false discovery rate correction in both the comparison of males and
712 females and the comparison of inferred maternal alleles and females (Fig. 5). While the sexual
713 selection analysis could not be conducted in the sdRAD-seq dataset alone, the comparison of
714 males and females using only sdRAD-seq data resulted in much larger F_{ST} values than other
715 analyses. In addition, no loci were identified as significant targets of selection in this analysis.
716 These comparisons imply that selection components analysis may produce different results
717 depending on the choice of ddRAD-seq or sdRAD-seq.

718 Our comparison of ddRAD-seq and sdRAD-seq was based on different individuals that
719 were genotyped by the two methods. This dataset was originally collected for a study of selection
720 components analysis in this population of pipefish, and it was due to logistical constraints
721 (particularly the shearing step) that we switched from sdRAD-seq to ddRAD-seq after
722 genotyping only 60 of the 444 collected individuals (see Flanagan & Jones 2017 for the analysis
723 of the ddRAD-seq individuals). Our comparison of two sub-samples of 60 ddRAD-seq
724 individuals indicate that the differences between sdRAD-seq and ddRAD-seq too large to be
725 explained by sampling error, and consequently must stem from differences in the methods. A
726 more rigorous comparison of the bias emerging from sdRAD-seq and ddRAD-seq would involve
727 genotyping the same individuals using both methods, but we believe that our analysis reveals
728 several noteworthy patterns and points of caution, especially as syntheses and meta-analyses of
729 RAD-seq studies are conducted.

730 In conclusion, we have shown that simply using different genotyping methods can result
731 in different allele frequencies, some of which are at the scale of differentiation measured
732 between populations. The major sources of this bias are polymorphic restriction sites, small and
733 asymmetric sampling schemes, and to some extent PCR duplicates. These results suggest that
734 bias could jeopardize comparisons of different datasets by inflating observed differentiation,
735 potentially obscuring true evolutionary processes. Encouragingly, researchers are identifying
736 ways to minimize bias (Ali *et al.* 2016; Hoffberg *et al.* 2016), and it may be possible to
737 incorporate sources of bias into genotyping methods using Bayesian statistics. In the meantime,
738 it is important for researchers to be cognizant of the factors causing different allele frequencies
739 between datasets, and to recognize that not all of these differences are the result of evolutionary
740 processes.

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749 **Data Accessibility**

750 Sequence data will be submitted to the NCBI sequence read archive. Data are archived on Dryad
751 at doi:10.5061/dryad.qf916. The gwscaR code is available from GitHub
752 (<https://github.com/spflanagan/gwscaR>). All other programs and scripts used in the analyses can
753 be downloaded from GitHub (<https://github.com/spflanagan/https://github.com/spflanagan/SCA>)
754 or can be obtained by contacting the authors.

755 **Conflict of Interest**

756 The authors have no conflicts of interest to declare.

Tables

Table 1.

The impact of analyzing ddRAD-seq and sdRAD-seq reads together or separately on variables related to allelic dropout and PCR duplications. These are the results of the analysis using Stacks. Shown for each variable are the mean, variance (in parentheses), and range (in square brackets: [min. – max.]). When analyzed separately, ddRAD-seq has higher per-SNP coverage, but the pattern is reversed when the datasets are analyzed together. Regardless of analysis approach, the ddRAD-seq dataset has slightly higher heterozygosity compared to the sdRAD-seq dataset. A decrease in heterozygosity may reflect allelic dropout as a consequence of polymorphic restriction sites. The sdRAD-seq dataset has higher mean variance in coverage than the ddRAD-seq dataset, possibly reflecting the impact of PCR duplicates (note that the values presented in the table for the coverage variability are the summary statistics for the coverage standard deviations, rather than the variances). The proportion of reference reads in heterozygotes indicates whether one allele was overrepresented at a given locus. The ddRAD-seq library had higher proportions of reference reads in heterozygotes than the sdRAD-seq library, and the proportion was higher in the ddRAD-seq library when it was analyzed together with the sdRAD-seq library than when it was analyzed separately. The number of dropped alleles are estimated by GBStools. Analyzing the dataset together results in more dropout alleles being maintained in the dataset, and sdRAD-seq has a higher number of dropout alleles than ddRAD-seq, regardless of analysis method.

	sdRAD separately	ddRAD separately	sdRAD together	ddRAD together
Average Coverage Per SNP mean (sd) [min - max]	11.55 (78.26) [4.06-20075.63]	13.26 (29.43) [3.76-2440.11]	15.12 (173.16) [4.2-20075.63]	13.44 (31.08) [4.43-2222.23]

Coverage Variability (in sd)	6.76 (39.14)	6.69 (21.14)	8.59 (83.67)	6.52 (19.27)
mean (sd) [min - max]	[1.43-9542.12]	[1.68-2405.49]	[1.35-9542.12]	[1.68-1491.7]
Proportion Heterozygotes	0.2 (0.14)	0.24 (0.16)	0.22 (0.14)	0.24 (0.15)
mean (sd) [min - max]	[0-1]	[0-1]	[0-1]	[0-1]
Proportion of Reference Reads in Heterozygotes	0.4998 (0.0609)	0.5008 (0.0543)	0.4995 (0.0669)	0.5034 (0.0697)
mean (sd) [min - max]	[0.0231-0.9741]	[0.0067-0.9094]	[0.0073-0.9741]	[0.0067-0.9745]
Number of Dropped Alleles	0.15 (0.23)	0.07 (0.13)	0.23 (0.35)	0.11 (0.19)
mean (sd) [min - max]	[0-1.01]	[0-0.95]	[0-1.92]	[0-1.4]

Table 2.

Analyzing the datasets with Stacks and samtools results in similar differences in coverage, proportion of heterozygotes, and extreme F_{ST} values. This table presents the results of analyzing the 60 sdRAD-seq individuals and a random subset of 60 ddRAD-seq individuals either together or separately. In both cases, the reads were aligned to the *S. scovelli* genome prior to analysis. For each variable displayed below, we report the mean, variance (in parentheses), and range (in square brackets: [min. – max.]). In both analyses, coverage differed between sdRAD SNPs and ddRAD SNPs. The samtools dataset had less extreme coverage values and variances, likely because samtools removes indels, which Stacks does not. This additional filter likely removed the most problematic loci, which in Stacks were those with extremely high coverage. In both the Stacks and samtools analyses, the ddRAD-seq dataset had a higher proportion of heterozygotes than the sdRAD-seq dataset when they were analyzed separately. Both Stacks and samtools resulted some extreme F_{ST} values between ddRAD-seq and sdRAD-seq datasets, even after filtering for coverage thresholds (Stacks analysis) or using quality filters (samtools).

	Stacks				samtools			
	sdRAD separately	ddRAD separately	sdRAD together	ddRAD together	sdRAD separately	ddRAD separately	sdRAD together	ddRAD together
Average Coverage Per SNP; mean (variance) [range]	11.55 (6,124.26) [4.06-20,075.63]	13.8 (1,364.11) [4-2,870.55]	15.12 (29,985.22) [4.2-20,075.63]	13.05 (775.57) [3.68- 2,145.45]	8.95 (35.41) [3-253.95]	10.55 (63.27) [3-251.53]	8.13 (55.69) [3-252.74]	8.13 (55.69) [3-252.74]
Proportion Heterozygotes; mean (variance) [range]	0.2 (0.02) [0-1]	0.25 (0.02) [0-1]	0.22 (0.02) [0-1]	0.21 (0.03) [0-1]	0.25 (0.02) [0-1]	0.27 (0.02) [0-1]	0.26 (0.02) [0-1]	0.26 (0.07) [0-1]
F_{ST} ; mean (variance) [range]	0.01 (0.0007) [-0.56-0.66]		0.01 (0.0005) [0-0.79]		0.01 (0.0009) [0-0.77]		0.01 (0.0016) [0-0.81]	
Filtered F_{ST} ; mean (variance) [range]	0.01 (0.0006) [-0.56-0.66]		0.01 (0.0003) [0-0.66]		0 (0.0005) [0-0.56]		0.02 (0.0013) [0-0.71]	

Table 3.

Results of the *in silico* digestion. The table is divided into four sections: (1) the effects of PCR duplication rates ("PCR Dup.") and a skewed allele frequency spectrum on F_{ST} values ("Mean AF"); (2) influence of restriction site mutation rates on F_{ST} by changing the effective population size (N_e) and by changing the average per-nucleotide mutation rate ("Restriction Site Mut."); (3) the effects of skewed allele frequencies (mean $p = 0.5$ vs mean $p = 0.8$), shearing bias, and asymmetric sampling ($n_{sd} = 60$, $n_{dd} = 340$) on F_{ST} ; and (4) how F_{ST} is affected by multiple sources of bias. F_{ST} is significantly affected by asymmetric sampling, which when combined with PCR bias actually elevated F_{ST} values. Skewed allele frequencies also resulted in greater differentiation between simulated sdRAD and ddRAD loci. We present the mean and standard error for F_{ST} as well as the maximum F_{ST} value ("Max. F_{ST} ") for each parameter combination.

	PCR Dup.	n_{sd}	n_{dd}	Mean AF	Shearing Bias	Restriction Site Mut.	N_e	Mean \pm SE F_{ST}	Max. F_{ST}
PCR Duplication and Skewed AF	0	200	200	0.5	NotBiased	0	10000	0.00056 \pm 0.000013	0.01679
	1	200	200	0.5	NotBiased	0	10000	0.00054 \pm 0.000012	0.01309
	2	200	200	0.5	NotBiased	0	10000	0.00058 \pm 0.000013	0.01529
	3	200	200	0.5	NotBiased	0	10000	0.00059 \pm 0.000012	0.01268
	4	200	200	0.5	NotBiased	0	10000	0.00061 \pm 0.000013	0.01438
	5	200	200	0.5	NotBiased	0	10000	0.00063 \pm 0.000013	0.01349
	1	200	200	0.8	NotBiased	0	10000	0.00097 \pm 0.000017	0.01569
	2	200	200	0.8	NotBiased	0	10000	0.00096 \pm 0.000017	0.01730
	3	200	200	0.8	NotBiased	0	10000	0.00095 \pm 0.000017	0.01603
	4	200	200	0.8	NotBiased	0	10000	0.00103 \pm 0.000018	0.01708

	5	200	200	0.8	NotBiased	0	10000	0.00100	± 0.000017	0.01963
Restriction site polymorphisms	0	200	200	0.5	NotBiased	10^{-7} to 10^{-8}	5000	0.00054	± 0.000012	0.01698
	0	200	200	0.5	NotBiased	10^{-7} to 10^{-8}	10000	0.00056	± 0.000012	0.01398
	0	200	200	0.5	NotBiased	10^{-7} to 10^{-8}	20000	0.00063	± 0.000014	0.01809
	0	200	200	0.5	NotBiased	10^{-8} to 10^{-9}	5000	0.00053	± 0.000011	0.01072
	0	200	200	0.5	NotBiased	10^{-8} to 10^{-9}	10000	0.00056	± 0.000013	0.01861
	0	200	200	0.5	NotBiased	10^{-8} to 10^{-9}	20000	0.00060	± 0.000013	0.01426
Skewed AF, shearing bias, and asymmetric sampling	0	200	200	0.5	NotBiased	0	10000	0.00056	± 0.000013	0.01679
	0	200	200	0.5	Biased	0	10000	0.00051	± 0.000031	0.01386
	0	200	200	0.8	Biased	0	10000	0.00085	± 0.000037	0.01338
	0	200	200	0.8	NotBiased	0	10000	0.00093	± 0.000016	0.01625
	0	340	60	0.5	Biased	0	10000	0.00040	± 0.000022	0.00649
	0	340	60	0.5	NotBiased	0	10000	0.00040	± 0.000009	0.01042
	0	340	60	0.8	Biased	0	10000	0.00082	± 0.000034	0.00863
	0	340	60	0.8	NotBiased	0	10000	0.00071	± 0.000012	0.01137
Multiple sources of bias	1	200	200	0.8	Biased	10^{-8} to 10^{-9}	10000	0.00098	± 0.000040	0.00934
	1	340	60	0.5	Biased	10^{-8} to 10^{-9}	10000	0.00041	± 0.000022	0.00804
	1	340	60	0.8	Biased	10^{-7} to 10^{-8}	10000	0.00072	± 0.000033	0.01037
	1	340	60	0.8	Biased	10^{-8} to 10^{-9}	10000	0.00076	± 0.000033	0.00765
	1	340	60	0.8	NotBiased	10^{-8} to 10^{-9}	10000	0.00072	± 0.000012	0.01084
	2	200	200	0.8	Biased	10^{-8} to 10^{-9}	10000	0.00107	± 0.000048	0.01244
	2	340	60	0.5	Biased	10^{-8} to 10^{-9}	10000	0.00048	± 0.000027	0.00911
	2	340	60	0.8	Biased	10^{-7} to 10^{-8}	10000	0.00075	± 0.000032	0.00898
	2	340	60	0.8	Biased	10^{-8} to 10^{-9}	10000	0.00077	± 0.000034	0.01008
	2	340	60	0.8	NotBiased	10^{-8} to 10^{-9}	10000	0.00074	± 0.000013	0.01122
	3	200	200	0.8	Biased	10^{-8} to 10^{-9}	10000	0.00106	± 0.000044	0.01189

3	340	60	0.5	Biased	10^{-8} to 10^{-9}	10000	0.00049	\pm 0.000025	0.00688
3	340	60	0.8	Biased	10^{-7} to 10^{-8}	10000	0.00072	\pm 0.000029	0.00823
3	340	60	0.8	Biased	10^{-8} to 10^{-9}	10000	0.00080	\pm 0.000036	0.00940
3	340	60	0.8	NotBiased	10^{-8} to 10^{-9}	10000	0.00076	\pm 0.000013	0.01384

Figures

Figure 1.

Comparison of locus coverage statistics in sdRAD-seq and ddRAD-seq datasets when analyzed together and separately. The sdRAD-seq dataset had a higher average number of reads per individual when assembled separately from the ddRAD-seq dataset, whereas the ddRAD-seq loci had higher per-individual coverage when assembled together with the sdRAD-seq reads (top row). The ddRAD-seq dataset had more reads per individual per locus than the sdRAD-seq dataset (bottom row)

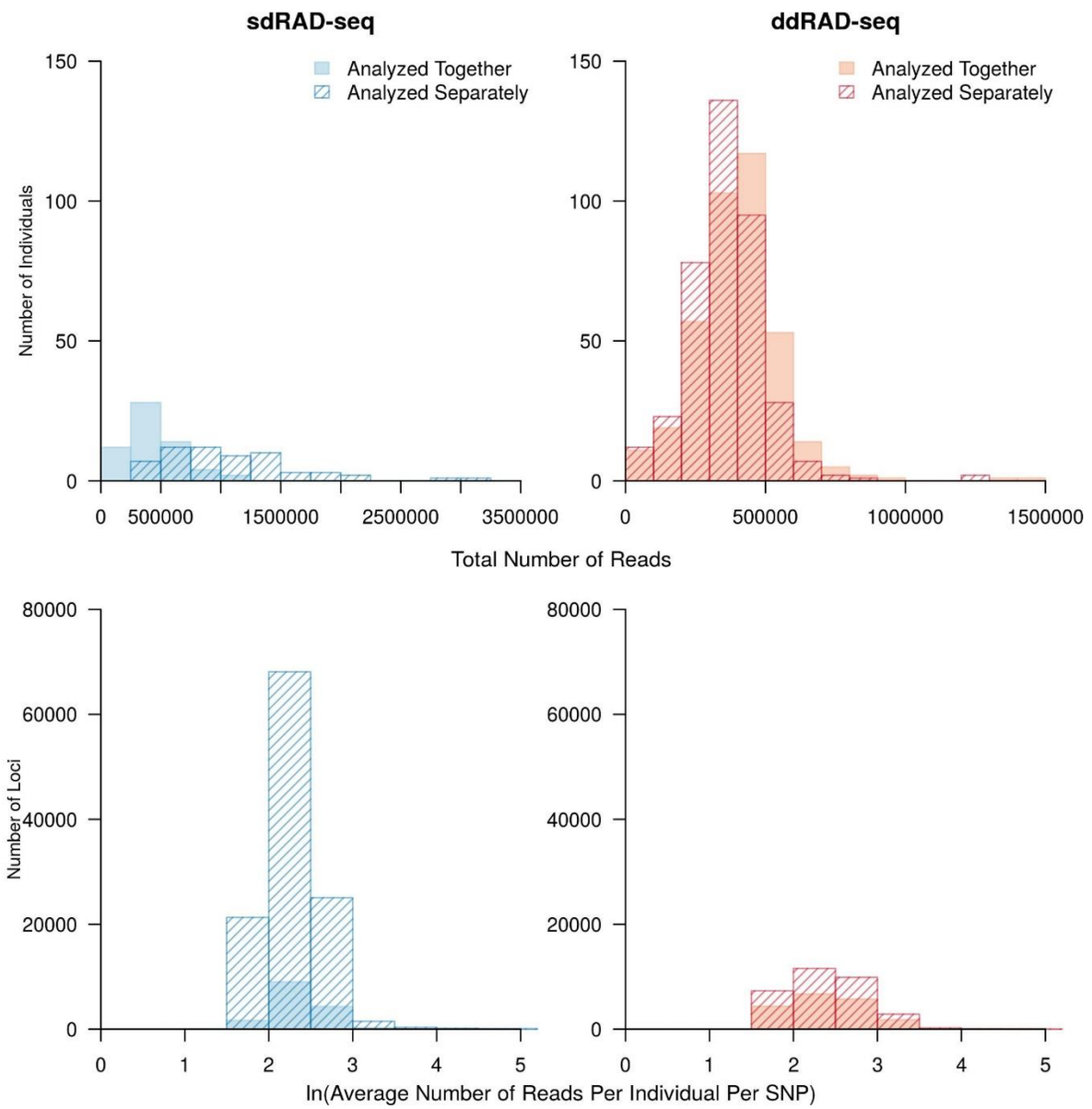


Figure 2.

Principal components analysis of the sdRAD-seq and ddRAD-seq datasets when analyzed together (left; light blue and orange) or separately (right; dark blue and dark red). When analyzed together the primary axis of variation in the dataset separates the individuals sequenced by sdRAD and ddRAD, whereas when the data were analyzed separately the sdRAD and ddRAD individuals overlap.

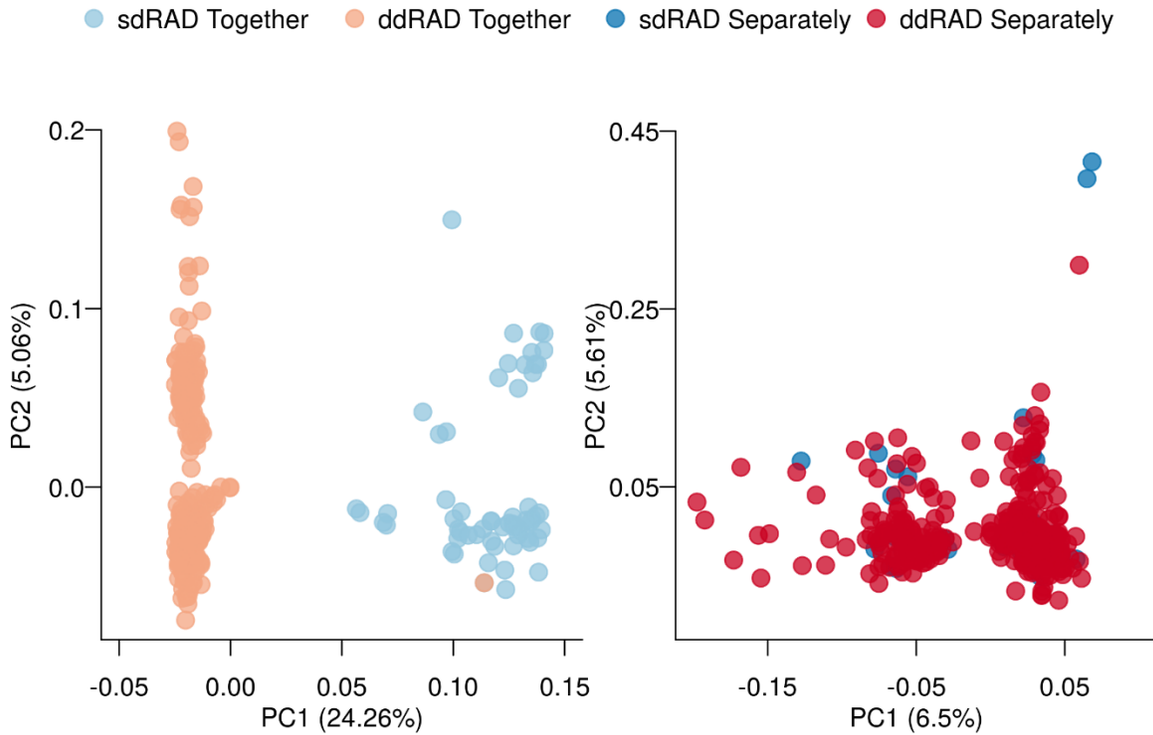


Figure 3.

Comparison of sdRAD-seq and ddRAD-seq allele frequencies using F_{ST} . We analyzed the complete datasets (left column), the complete sdRAD dataset and a subset of the ddRAD dataset (middle column), and two subsets of the ddRAD dataset (right column) to evaluate the impact of sample size on the allele frequencies. For each of the comparisons, we analyzed the sdRAD-seq and ddRAD-seq data in two separate runs of Stacks (Catchen *et al.* 2013; Catchen *et al.* 2011), which are presented in the top two rows. We also pooled the sdRAD-seq and ddRAD-seq data and analyzed them in a single analysis in Stacks (Catchen *et al.* 2013; bottom two rows; Catchen *et al.* 2011). For each of the analyses, we also investigated the effect of imposing a coverage filter to remove SNPs with average per-SNP coverage $\leq 3X$ or $>50X$. The number of SNPs and the mean F_{ST} values for the analyses are presented in each panel.

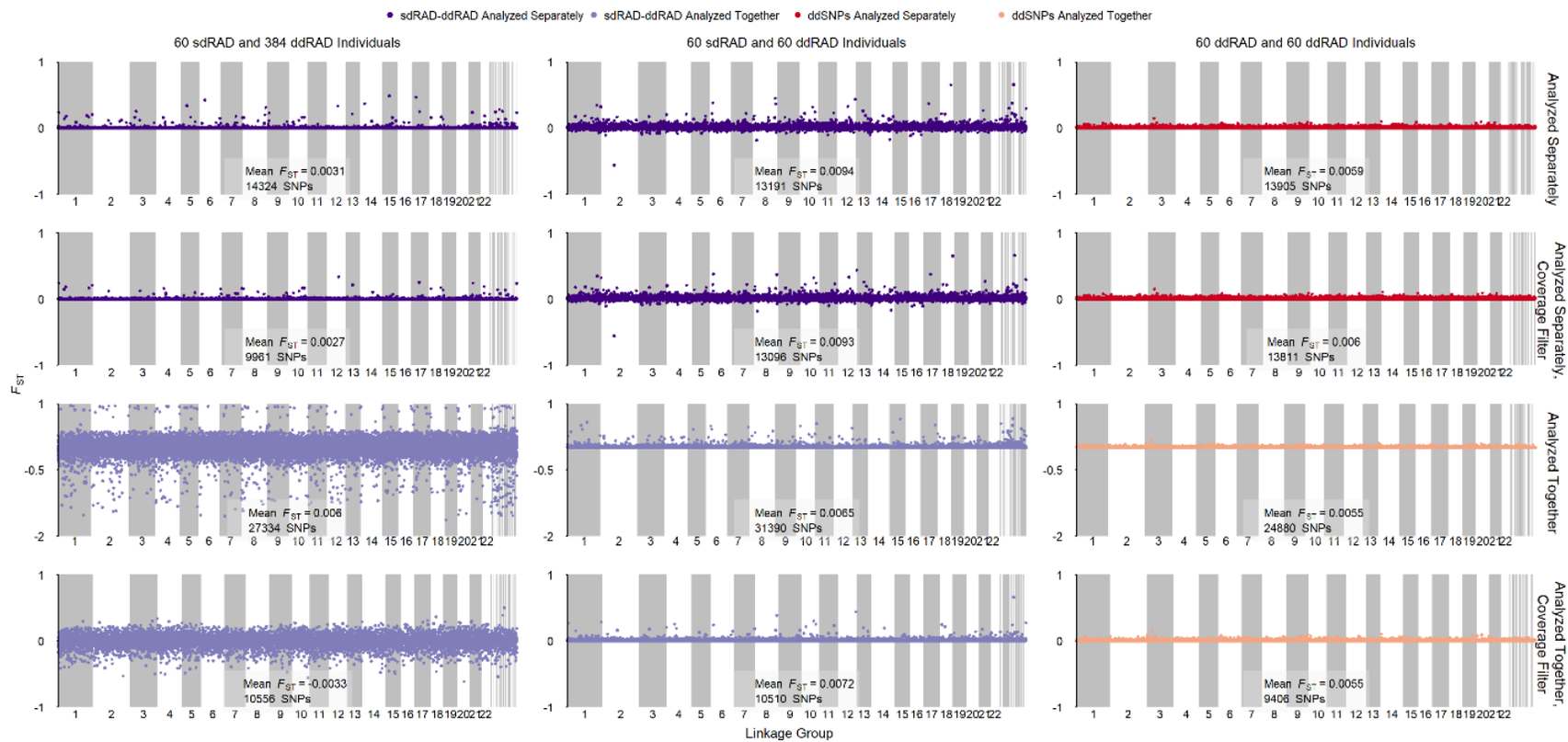


Figure 4.

The impact of sdRAD and ddRAD coverage on F_{ST} values when the datasets were analyzed separately (left) and together (right). The values presented here are mean F_{ST} values for each coverage category without any coverage filters imposed. When analyzed separately, the highest F_{ST} values (light grey) occurred when one dataset had medium coverage (5-20x) and the other had high coverage (30+). When analyzed together, many of the F_{ST} values were negative, which is indicative of major issues with the dataset. These extreme values occurred when ddRAD had high coverage ($\geq 50x$) and sdRAD had medium coverage (10-20x). Note that the two panels are on different scales.

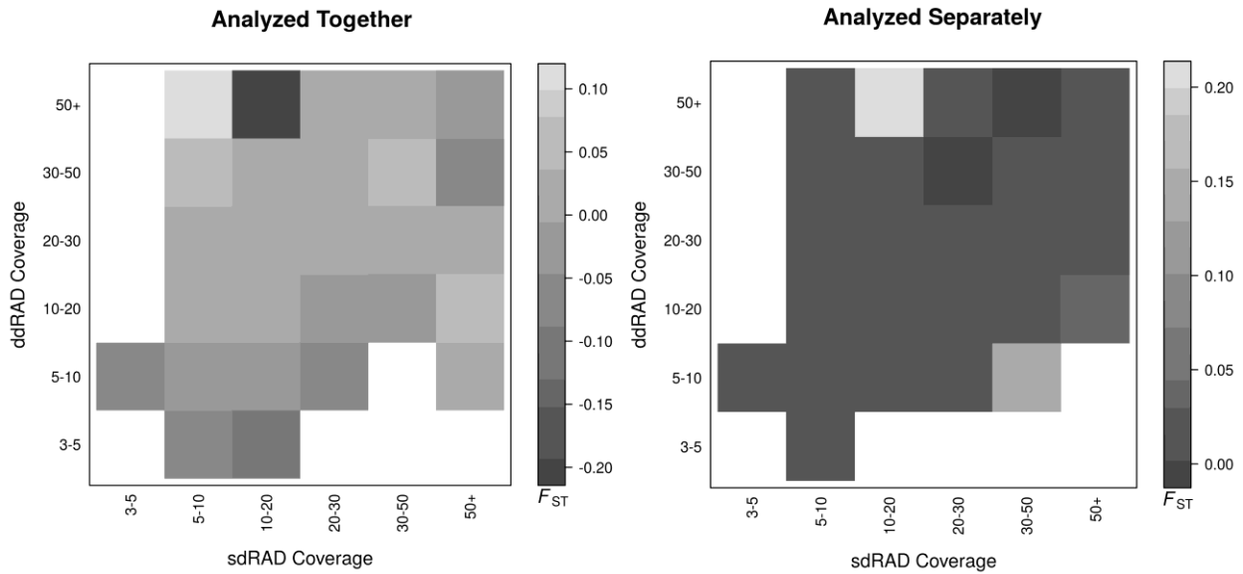
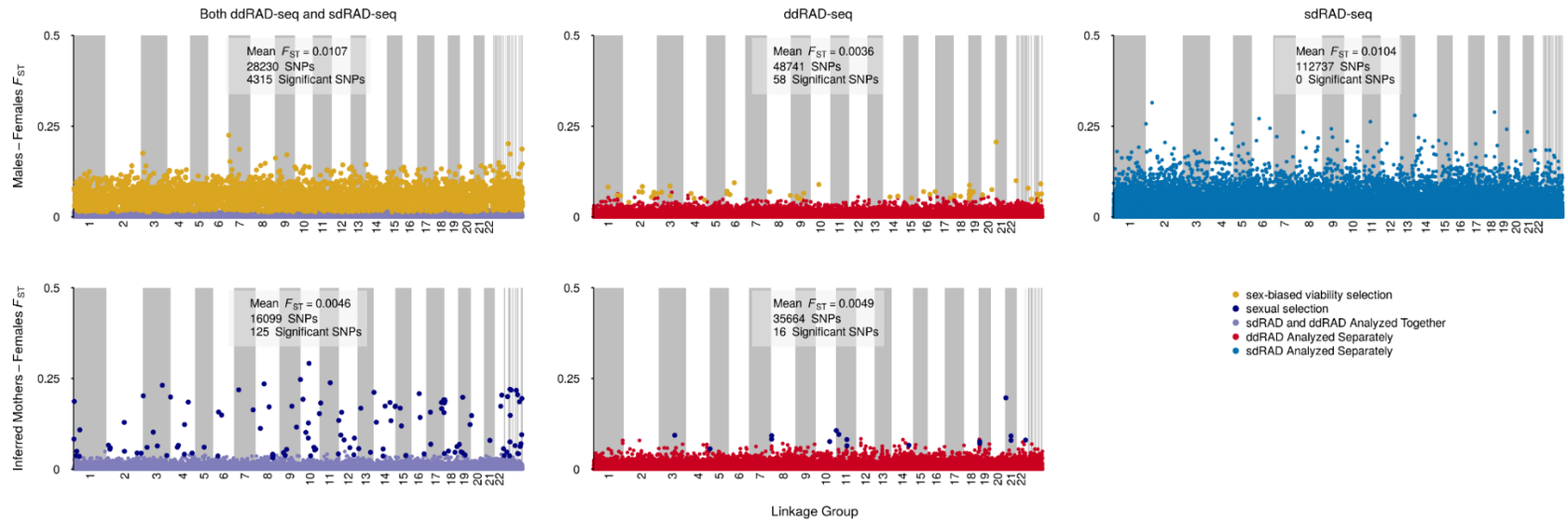


Figure 5.

The type of sequencing method and the data analysis method greatly impacted the results of the selection components analysis. Analyzing the ddRAD-seq and sdRAD-seq datasets together (left column) yielded higher F_{ST} values than when using only ddRAD-seq (middle column). Analyzing only the sdRAD-seq dataset (right column) resulted in higher F_{ST} values between males and females than both the combined analysis and the ddRAD analysis. The colors represent the different types of analyses (sdRAD and ddRAD analyzed together in purple, ddRAD analyzed separately in red, and sdRAD analyzed separately in blue) and the different types of inferred selection (sexual selection in dark blue and sex-biased viability selection in yellow).



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