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1 Detecting the true extent of introgression during anthropogenic hybridization

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- 3

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9 Abstract

10 Hybridization among naturally separate taxa is increasing due to human impact,

- 11 and can result in taxon loss. Previous classification of anthropogenic
- 12 hybridization has largely ignored the case of bimodal hybrid zones, in which
- 13 hybrids commonly mate with parental species resulting in many backcrossed
- 14 individuals with a small proportion of introgressed genome. Genetic markers can
- 15 be used to detect such hybrids, but until recently too few markers have been
- 16 used to detect the true extent of introgression. Recent studies of wolves and
- 17 trout have used thousands of markers to reveal previously undetectable
- 18 backcrosses. This improved resolution will lead to increased detection of late
- 19 generation backcrosses, shed light on the consequences of anthropogenic

20 hybridization, and pose new management issues for conservation scientists.

22 Anthropogenic hybridization

23 Anthropogenic hybridization (see Glossary), in which human disturbance 24 leads to range overlap and hybridization of previously reproductively isolated 25 populations or species is a growing conservation concern [1-3]. With increased 26 human-generated movement of species into new ranges, there is an increasing 27 number of cases of hybridization between species that were historically 28 **allopatric** [4]. Disturbance of habitats can also result in a breakdown of reproductive isolation between previously isolated, sympatric species [1]. 29 30 **Introgression** is usually hard to detect from phenotypes and there is growing 31 evidence that backcrossing has often proceeded further than is detectable by low 32 density genetic marker panels. In this article we make the case that genomic 33 approaches are essential and increasingly available to disentangle late 34 generation backcrosses from parental populations after introgression has 35 occurred.

36

37 The benefits of anthropogenic hybridization

38 There are possible benefits of anthropogenic hybridization. Policy makers can 39 use hybridization as a management tool to help endangered populations. In 40 'genetic rescue' programs (i.e. breeding programs designed to release small 41 populations from inbreeding depression), individuals from a closely related 42 population or subspecies are introduced to an inbred population to manage 43 inbreeding depression. For example, when Florida panthers (Puma concolor 44 *coryi*) were threatened due to inbreeding depression, eight Texas panthers (*P.* 45 concolor cougaur) were introduced. The hybrid kittens survived better, and the population is now recovering [5]. Approximately 90% of such genetic rescue 46 47 attempts have been successful, showing that anthropogenic hybridization is a 48 viable conservation method [6]. Adaptive introgression ('evolutionary rescue') in 49 which beneficial alleles from an introduced population are selected for in hybrid 50 individuals is another possible benefit of anthropogenic hybridization. For 51 example, a segment of chromosome 15 that has naturally introgressed from 52 Populus balsamifera into P. trichocarpa appears to allow P. trichocarpa to live in 53 colder, drier areas than *P. trichocarpa* individuals without this haplotype [7]. 54 This suggests that there is potential for adaptive introgression to facilitate

- evolutionary rescue of populations at risk of extinction due to climate change [8],
- although such genomic management of at risk populations much enabling
- 57 research, and should be approached with caution [9, 10].
- 58

59 The problems with anthropogenic hybridization

60 Anthropogenic hybridization can cause problems for native species. When no 61 offspring or sterile offspring are produced, reproductive effort is wasted [11]. 62 When fertile F1s are formed, introgression between the two previously diverged 63 species is possible. There are two reasons why even low levels of introgression 64 of non-native alleles are of concern from a conservation perspective. First, if all individuals of a species are hybrids then the species as it was is extinct. This has 65 66 been termed 'extinction by hybridization' [11-15]. Note, however, that there may still be many copies of the native alleles represented in the population, so long as 67 68 the population itself is large enough, and from a 'gene view point' we may be 69 content with this mode of conservation [16].

70

71 The second problem with hybridization is that introgression and recombination 72 break up linked gene complexes, and non-native alleles that are favoured (or no 73 longer in linkage with deleterious alleles) can be swept to fixation [17]. While this leads to an initial increase in biodiversity (because alleles from both the 74 75 native and non-native populations are present) as non-native alleles sweep to 76 fixation, native alleles are lost. If we again take a gene view point of biodiversity, 77 any alleles lost from the native population are a loss in biodiversity from the 78 system. For example, non-native alleles at three out of 68 genetic markers have 79 gone to fixation in some populations of California Tiger Salamanders 80 (Ambystoma californiense) after hybridization with Barred Tiger Salamanders (A. 81 *mavortium*) [18]. This has occurred in California Tiger Salamander populations 82 that are nearly 100km from the original Barred Tiger Salamander introduction 83 site, suggesting that these alleles have higher fitness than the native, California 84 Tiger Salamander alleles that they have replaced [18]. 85

86 Goals of studies of anthropogenic hybridization

Studies of anthropogenic hybridization have different goals. A researcher might
be interested to know if hybridization has occurred at all in a population to

- 89 determine whether it should provide the breeding stock for new populations,
- 90 and or whether it should be quarantined because of hybridization. Relatively few
- 91 informative markers are needed to detect individuals of hybrid origin in any
- 92 particular population, as the detection of any non-native allele is a clear
- 93 indication of hybridization [19].
- 94

95 However, if a researcher wishes to understand more about the underlying

- 96 process of hybridization and introgression, then many more markers are
- 97 required. Specific goals might include: to select individuals for breeding
- 98 programs; to understand the relationship between genotype and phenotype; to
- 99 understand the type of hybrid system involved (see next section); and to
- 100 investigate mating patterns and fitness. For any of these goals, it is ideal to
- 101 quantify individual **admixture** accurately, and to do this this hundreds or
- 102 thousands of informative markers may be required (see below).
- 103

104 Classifying hybridization

To assist researchers and policy makers in addressing anthropogenic
hybridization, Allendorf and colleagues [11] categorized hybridization outcomes.
Types 1-3 applied to naturally-occurring hybridization while Types 4-6 applied
to anthropogenic hybridization. Type 4 results in few or sterile F1 hybrids, and is
characterized by wasted reproductive effort. Type 5 results in a hybrid swarm
with widespread introgression into particular populations, but some populations
do not experience hybridization at all. Finally, Type 6 results in a complete

- 112 hybrid swarm following break down of reproductive isolation between species
- 113 across all populations [11].
- 114

115 Three axes of variation determine the outcome of anthropogenic hybridization:

- 116 differences in hybrid fitness, time since **secondary contact**, and mating patterns
- 117 of hybrids. Time since secondary contact and mating patterns of hybrids were
- 118 not explicitly considered in Allendorf et al's original categorization. Type 4
- differs from Types 5 and 6 along an axis of hybrid fitness, where intrinsic post

120 zygotic isolation affects hybrids in Type 4, but not in Types 5 or 6. This results in

- 121 little to no backcrossing in Type 4 hybrid zones, as hybrids are extremely unfit
- 122 compared to parental species. This decrease in hybrid fitness must be extreme,
- as even with a 90% decrease in fitness, the proportion of hybrids in a hybridizing
- 124 population is expected to increase [20].
- 125

126 We suggest that the only difference between Allendorf et al's [11] Type 5 and 127 Type 6 is time since secondary contact. When an F1 reproduces, all of its 128 offspring and descendants are admixed to some extent [20]. If Type 5 129 characterizes a system where only one or few populations have introgression. 130 Type 6 is the logical outcome of this same system, assuming random mating and 131 sufficient time for migration between populations. Thus, we consider Type 5 and 132 Type 6 to be the same, both hybrid swarms with a breakdown of assortative 133 mating, in which hybrids have the same mating success as either of the parental 134 species individuals, and common enough that hybrid x hybrid matings occur.

135

When there is a preference among hybrids for parental species phenotypes, or
hybrids are very rare, we expect a different pattern of introgression.

138 Backcrossing into the parental species leads to an increasingly large number of 139 individuals with a small proportion (<10%) of their genome that is from the 140 opposite species. As backcrossing continues, morphological differences between 141 parental species and backcrossed individuals lessen, making it more and more 142 difficult to detect a backcross using only phenotypic traits. This results in many 143 hybrid individuals with very small proportions of another genome, although 144 with a maintained bi-modal distribution of trait values between the two parental 145 species (Figure 1). From a conservation perspective we consider this to be a 146 worst-case scenario as these introgressed individuals are very difficult to detect. 147 This can be contrasted with a general lack of assortative mating, in which hybrid 148 individuals are as likely to breed with other hybrid individuals as with parental 149 species (leading to a hybrid swarm), or, in the unlikely event of true assortative 150 mating, where hybrid individuals preferentially breed with each other, which 151 would lead to the eventual formation of a hybrid species e.g. [21]. The contrast 152 between hybrid zones with unimodal distributions of traits and admixture

scores and those with bimodal distributions has previously been described in the
context of naturally occurring hybrid zones [22], but does not yet seem to inform
studies of anthropogenic hybridization.

156

157 The distribution of hybrid scores in a system at equilibrium varies depending on 158 ecological factors that can affect hybrid fitness, and hybrid encounter rate. 159 Extrinsic post zygotic isolation can vary according to ecological factors, affecting 160 the ability of hybrids to successfully mate and reproduce [23]. Further, stochastic 161 factors, particularly when hybrids are rare, or management might alter the 162 reproductive success of hybrid individuals in wild systems. However, if hybrids are fertile, the proportion of hybrid individuals in all populations should increase 163 164 [20], leading to the extreme end points of majority hybrid populations which either follow a hybrid swarm or **bimodal hybrid zone** distribution. 165

166

167

168 Key considerations for genetic analyses of anthropogenic

169 hybridization

170 Published studies of anthropogenic hybridization generally follow a similar 171 protocol. Researchers use codominant marker genotypes to estimate divergence 172 between the two species [24] and then use a clustering approach such as 173 STRUCTURE [25-28], or ADMIXTURE [29, 30] to partition individuals into 174 different genetic groups (K). Those individuals with an admixture score (Q) 175 intermediate to the extreme admixture scores associated with parental species 176 individuals are designated hybrids. Many studies then use HYBRIDLAB [31, 32] 177 or similar methods to simulate hybrid genotypes from the sampled genotypes to 178 assess the **efficiency** (i.e. type II error rate, rate of assigning hybrid individuals 179 as parental species), and **accuracy** (i.e. type I error rate, rate of erroneously 180 assigning parental species individuals as hybrids; [33]). The 'overall 181 performance' of an analysis is the product of efficiency and accuracy and this 182 performance can be used to assess the reliability of the study itself [33]. Here we 183 outline some best practices and points to consider in order to avoid 184 underestimation of the extent of hybridization. 185

186

187 **Divergence between parental species**

188 It is highly relevant to have an estimate of divergence between the focal species 189 in the absence of hybridization. **F**_{st} is often reported in studies of anthropogenic 190 hybridization, but is rarely used to motivate the marker density deployed for 191 estimates of individual admixture, typically because the same markers are used 192 to determine both Fst and individual Q estimates. Simulations have clearly shown 193 that species (or subspecies) with lower divergence will require more markers to 194 accurately estimate admixture, because of shared polymorphisms between them, 195 leading to fewer **diagnostic markers** [33]. While it might not be practical to use 196 markers to estimate F_{st} and then determine how many markers are needed to 197 estimate individual admixture scores, an initial assessment of F_{st} will hint at how 198 much power a system has to detect advanced backcrosses.

199

200 Historical admixture

201 Many systems have a history of repeated secondary contact and hybridization. 202 Documenting historical admixture using genomic resources can determine 203 whether the introgression found is due to recent, anthropogenic forces, or to 204 natural causes, which will change the conservation status of the situation [34, 205 35]. There are techniques for detecting historical admixture. For example, the 206 ABBA-BABA test can be used to determine if there has been historical 207 introgression from a third species or population into each of two closely related 208 sister taxa, to explain variation that is not well explained by a null assumption of 209 bifurcating phylogeny [36]. This technique can be applied to either sequences of 210 single individuals from each population, or to multiple individuals from each 211 population [37], and can be used to indicate historical (hundreds to thousands of 212 generations before present) admixture. Similarly, $\delta a \delta i$ analyses can be used to 213 determine how well different demographic models fit the pattern of variation in 214 the data, where demographic models can include admixture at different time 215 points [38]. For example demographic modeling was used to demonstrate that 216 hybridization between golden-winged (Vermivora chrysoptera) and blue winged 217 warblers (V. cyanoptera) has probably been occurring since the original species 218 split, and is not solely due to anthropogenic forces [39]. Finally, researchers can

- examine the length of haplotype blocks that are identical by descent, as linkage
- disequilibrium decays over time due to recombination [40, 41]. The distribution
- of haplotype block lengths should follow a Poisson distribution [41] and
- 222 deviation from this distribution can be used to infer population admixture over
- both short (tens of generations) [42] and long time spans [41]. These and other
- techniques for disentangling historical and contemporary admixture are
- 225 reviewed in [43].
- 226

227 Generations since secondary contact and recombination rates

228 It is important to estimate the number of generations since secondary contact to 229 estimate the potential number of backcross generations in a system. This 230 estimate might have substantial uncertainty, but in many cases of anthropogenic 231 hybridization there are historical records that suggest when a non-native species 232 was first introduced or sighted that can be combined with typical generation 233 times for the taxa involved. The expected proportion of invasive genome in a 234 backcrossed individual halves with each successive generation of backcrossing 235 [44].

236

237 Recombination each generation leads to less linkage disequilibrium between 238 non-native loci, which means that genotype at a species-specific marker in one 239 position is less informative about surrounding, un-sampled loci. For example, 240 genomic regions with high recombination rates were found to be associated with 241 more introgression of the non-native genome in replicate swordtail (Xiphophorus 242 *birchmanni and X. malinche*) hybrid zones [17]. Due to obligatory crossing over, 243 which is expected to occur once per chromosome arm [45], at least twice as 244 many markers as there are chromosome arms are needed to cover each 245 independent section of the genome. In some cases, there is a species-specific 246 estimate of recombination (e.g. [46]), or one can refer to taxon-specific patterns. 247 For example, there is as much as 10 times more recombination in avian genomes 248 than in mammalian genomes [47]. Additionally, information on recombination 249 rate can be combined with genomic methods examining haplotype block lengths 250 to date introgression events (as discussed above). We discuss how many 251 markers are needed further in Box 1.

252

253 Assessing the power of markers

254 Many studies of anthropogenic hybridization assess the power of genetic 255 markers used by simulating hybrid genotypes and then determining the power 256 the markers have to detect these hybrid genotypes [48]. When assessing the 257 power of markers in this way, it is important to ensure that the biology of the 258 system is reflected in the simulation. In particular, if the two species of interest 259 have been in contact for many generations and F1s are thought to be fertile 260 (Figure 1), then simulations should account for the possibility of many 261 generations of backcrossing. This is rarely done in conservation genetic studies -262 many studies simulate backcrosses to assess the power of their markers, and 263 find low power to detect even first generation backcrosses, for example finding 264 less than 80% of first generation backcrosses are properly assigned [49, 50]. 265 Further information obtained from laboratory or field studies, such as 266 asymmetry in hybrid fertility (e.g. between sexes, Haldane's Rule [51] or 267 according to the species of the mother of the F1, Darwin's Corollary [52]), should 268 also be included in simulations. For example, if previous laboratory work has 269 established that backcrossing is largely unidirectional because of decreased 270 fitness of hybrid individuals in the opposite direction (as expected by Darwin's 271 Corollary) or due to the relative abundance of the parental species, then 272 mitochondrial markers should be integrated into future analyses to add power to 273 detect hybrids.

274

275 **Defining hybrid individuals**

276 To be defined as a hybrid, a focal individual must be genetically differentiated 277 from both parental species. Parental species are assumed to have an admixture 278 (Q) score of 0 or 1, although because of error (e.g. non-diagnostic markers, 279 genotyping errors), very few individuals will have an estimated score of exactly 0 280 or 1. Any score in between indicates a hybrid [25]. It is typical for a researcher to 281 set a Q score as a cut-off for hybrid individuals, so any individual above (or 282 below) this score is considered parental. Thresholds are determined either by 283 power, specifically, at what level can the markers differentiate between hybrids 284 and parental species, or by the number of acceptably mis-matched markers, e.g.

285 one allele indicative of the other species might be an error, but two markers 286 suggest hybridization [53]. These thresholds can range widely between studies, 287 from 0.8 [54] to 0.999 [30] in relation to a parental species score of 1.0. 288 Determination of the threshold is a balancing act between Type I and Type II 289 errors, in which the researcher must decide whether it is better to mistakenly 290 assign a parental species individual as a hybrid (Type I; too low 'accuracy'; [33]) 291 or assign hybrid individuals as parental types (Type II; too low 'efficiency'; [33]). 292 If the researcher accepts a higher level of Type II errors, they consider advanced 293 backcrosses as parental species. For example, an admixture score threshold of 294 0.8 would include most second-generation backcrosses (87% of the genome is 295 species A, 13% of the genome is species B on average) as parental species. 296 Similarly, with a Q score of 0.9, third-generation backcrosses (average of 93% 297 species A) would be included as parental species individuals.

298

299 There are two ways to ameliorate error introduced in species assignment using 300 thresholds. One obvious way is to employ more markers (Box 1), which 301 increases the power of a study and allows the setting of thresholds approaching 302 0 and 1. Studies that have used thousands of markers use the most stringent 303 thresholds e.g. [30]. A second solution to the threshold problem is to do away 304 with them entirely. Rather than assigning individuals to species classes based on 305 point estimates, it is more appropriate to use **credible** or **confidence intervals** 306 around point estimates which capture uncertainty in the marker system 307 appropriately (Box 2). In this scenario any individual with a credible interval 308 overlapping 0 or 1 is considered a parental species and all others are considered 309 hybrids.

310

An additional problem in separating hybrid individuals from parental species is that some hybrids, particularly later generation of backcrosses, will be homozygous for all sampled diagnostic loci by chance. This is due to increased variation around the proportion of genome inherited from each parental species with each generation of backcrossing ([44]; Box 1). The hybrid nature of these individuals will be undetectable, and they will be classified as parental species, even though unmarked genome regions may be introgressed. Increasing the

number of markers increases the probability of sampling a hybrid individual at
loci that are heterozygous or homozygous for alleles representative of both
parental species (Box 1).

321

322 Higher density markers to identify bimodal hybrid zones

323 When researchers apply higher density marker panels to examples of 324 anthropogenic hybridization, they generally uncover more backcrossed 325 individuals compared to studies using low-density panels, and can draw more 326 accurate conclusions about the system. These newly-detected backcrosses are 327 often genetically very similar to the parental species, with less than 10% 328 introgression, indicative of a bimodal hybrid zone. For example, in a study of 329 Italian wolves that hybridize with domestic dogs, use of 170,000 SNPs found that 330 hybridization had occurred 3 -5 generations prior to sampling [30]. This multi-331 generation backcrossing was not detectable in the population when 18 332 microsatellite markers were used [49]. Further, while very few individuals were 333 found to have Q scores between 0.25 and 0.75, as would be expected in a hybrid 334 swarm with a complete breakdown of reproductive isolation, 62% of sampled 335 Eurasian wolves had a small proportion (<5%) of admixture with domestic dogs 336 [55]. The Eurasian wolf – domestic dog system has the distribution of admixture 337 scores and phenotypes that characterizes a bimodal hybrid zone with some 338 degree of mating preference for parental phenotypes, or rare intermediate 339 hybrids. In this system, most individuals are either phenotypically dog-like with 340 extreme Q scores at one end of the distribution, or phenotypically wolf-like with 341 Q scores at the other end of the distribution. There are few individuals with 342 intermediate Q scores and phenotypes. This can be contrasted with the 343 westslope cutthroat (Oncorhynchus clarki lewisi) – rainbow trout (O. mykiss) 344 system, which has also recently been genotyped using 3180 diagnostic SNPs [56]. While the increase in number of markers did lead to increased detection of 345 advanced backcrosses, there were also many individuals with intermediate Q 346 347 scores and phenotypes [56, 57]. This suggests that the westslope cutthroat-348 rainbow trout system is a hybrid swarm that has little assortative mating. 349

350 **Designing an ideal study of an anthropogenic hybrid zone**

351 When embarking on a study of anthropogenic hybridization, there are many 352 considerations in deciding on the genetic resources to be used (Box 1). As whole 353 genome sequencing (WGS) becomes cheaper [58], conservation biologists should 354 consider whether WGS is the best way forward. Firstly, WGS data allows for 355 detection of heterogeneity of introgression across the genome. If conservation 356 biologists truly adopt a 'gene view point' of hybridization [16] then individuals 357 ought to be classified based on whether they carry specific alleles at identified 358 loci, rather than by overall Q scores (but see [10] for a discussion of the difficulty 359 of implementing this approach). Secondly, WGS enables the researcher to 360 distinguish between historical and contemporary introgression. Finally, we anticipate that the use of WGS will result in more diagnostic or **ancestry** 361 362 informative markers being detected, and thus make studies more powerful. 363 Researchers will be more confident in their estimates of individual admixture, 364 and will report the power and confidence associated with their analyses (Box 2). 365 While the bioinformatics skills required to assemble a genome and call SNPs may 366 seem intimidating, we believe that 1) these are skills are now routinely taught in 367 universities and 2) WGS presents an additional opportunity for conservation 368 biologists to collaborate with speciation geneticists (Box 3). Another 369 consideration is that high quality DNA is needed for the most accurate 370 assemblies, although progress is being made towards high quality sequences 371 from poor quality samples (e.g. [59]). While the use of WGS is more expensive 372 than microsatellite marker studies, when the cost of microsatellite markers, 373 including the cost of labour, was compared to the use of SNP markers in 374 European wolves, SNPs were less expensive if at least 24 samples were 375 genotyped [60]. This suggests that the use of thousands of variable genome wide 376 markers (e.g. from ddRAD [61]) may represent a practical middle ground for 377 conservation biologists, depending on the history and biology of the system. 378 Taken together, we believe that the best way forward to accurately detect 379 backcrossing in studies of anthropogenic hybrid zones is to routinely use higher 380 density markers, including WGS when possible. 381

382 Concluding Remarks

383 Advanced backcrosses are unlikely to have been detected with many of the 384 methods that biologists studying anthropogenic hybridization have used to date. 385 Most studies of anthropogenic hybridization have used fewer than 20 markers 386 [13], too few to reliably detect individuals that are more than two generations 387 backcrossed [33], unless markers are perfectly species diagnostic [44]. For this 388 reason, it is rare for studies to consider backcrossed individuals past the second 389 generation of backcrossing, regardless of the number of generations that have 390 passed since secondary contact. Here, we suggest that studies should attempt to 391 go much further. By accounting for the number of generations since secondary 392 contact and increasing the density of genetic markers accordingly, many more 393 backcrossed individuals will become distinguishable from the parental 394 populations. We echo the call for more genetic markers to be used in these 395 studies to allow for higher accuracy and efficiency [1, 3, 13, 33, 62], particularly 396 since we have now entered the genomics era, making tens or hundreds of 397 thousands of markers obtainable even in non-model systems [58]. It seems likely 398 that anthropogenic hybridization will only increase in frequency and result in 399 increased gene flow between previously isolated species [1]. The increase in 400 number of markers and associated power will also open up the opportunity to 401 ask new questions in these systems, parallel to those speciation biologists 402 explore in natural hybrid zones (Box 3). There are new challenges with 403 increased marker density, but a genomic approach to studying these systems will 404 help researchers to detect backcrosses and make the best policy 405 recommendations. 406

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411 Additional Elements

412

413 Glossary

414	٠	Anthropogenic hybridization: the breakdown of reproductive isolation
415		between two species due to human action, including but not limited to,
416		species introduction, habitat disturbance or escape of domestic species.
417	•	Accuracy: the proportion of identified hybrids that are actually of hybrid
418		ancestry [33]. A low accuracy suggests a high rate of type I errors, in
419		which parental species individuals are erroneously assigned as hybrids.
420	٠	Admixture: the mixing of genomes from structured or diverged
421		populations
422	٠	Allopatry: species in non-overlapping ranges
423	٠	Ancestry informative markers: genetic markers with substantial allele
424		frequency differences across populations, which can be used to assign
425		individuals to each population [63]
426	•	Bimodal hybrid zone: a hybridizing population in which preference for
427		parental phenotypes, or scarcity of hybrids with which to mate, results in
428		a population that includes few F1 hybrids, and many backcrossed
429		individuals with a low level of introgression that often resemble the
430		parental species in phenotype. Can be unimodal (if backcrossing is into
431		just one parental species) or bimodal (backcrossing into both parental
432		species) [22]
433	٠	Credible interval: the range of possible values surrounding a point
434		estimate, representing the uncertainty in the estimate
435	٠	Diagnostic markers: markers with fixed allele differences across
436		populations
437	•	\mathbf{D}_{xy} : an absolute measure of genetic differentiation, calculated as the
438		proportion of nucleotides that differ between two homologous sequences
439		within the same or different population.
440	٠	Efficiency: Proportion of correctly identified individuals in each group
441		[33]. If the null hypothesis is that an individual is from the parental
442		species rather than a hybrid individual, then low efficiency suggests a

443		high rate of type II errors, in which hybrid individuals are incorrectly
444		assigned as parental species.
445	•	\mathbf{F}_{ST} – A measure of genetic differentiation between populations based on
446		the difference in allele frequencies within and between populations [64]
447	•	Hybridization: mating of individuals from diverged populations
448	•	Hybrid: an individual that has an intermediate genotype between two
449		diverged, parental populations, as the result of interbreeding between
450		these populations
451	٠	Hybrid swarm: a hybridizing population that includes F1 hybrids and
452		various backcrosses, due to a total breakdown of assortative mating. Also
453		known as a unimodal hybrid zone [22].
454	٠	Introgression: the movement of alleles between genetically
455		differentiated forms (including populations, species, etc), mediated by
456		backcrossing [65]
457	•	Secondary contact: Occurs when two (or more) species that have been in
458		allopatry come back into sympatry
459	•	Sympatry: species in overlapping ranges
460		

461 Figure 1: Anthropogenic hybridization falls into three main categories. These are 1) systems with inviable or infertile hybrids, 2) bimodal hybrid zones in which 462 463 there is either mating preference for parental species phenotypes or the relative 464 abundance of parental species means most matings are backcrosses and 3) 465 hybrid swarms in which there is random mating and many hybrid individuals. In 466 this schematic figure we illustrate for each type of anthropogenic hybridization 467 system how many individuals of each admixture (Q) score might be found and 468 typical distributions of mating success across Q scores according to whether 469 there is a high likelihood of hybrid individuals mating with the parental species 470 phenotypes present. While we represent hybrid swarms and bimodal hybrid 471 zones as categorically different, these are probably ends of a continuum and 472 some systems may be intermediate between them. Note that we have 473 represented (2) as a bimodal hybrid zone due to backcrossing into both parental 474 species. Alternatively there can be a single (i.e. unimodal) hybrid zone due to 475 unidirectional backcrossing.

476 Box 1 - How many markers do I need to discover backcrossed individuals 477 in my system?

478

479 Substantial power is needed to detect individuals that are the result of repeated 480 generations of backcrossing. General rules have been suggested, including that 481 for each additional generation twice as many markers are needed [44], and that 482 at least 48 markers would be needed to consistently detect first generation 483 backcrossing in hybrids with parental species that have an $F_{st} = 0.21$ [33]. 484 However, we are now in the age of genomics, when the cost of increasing marker 485 density is dramatically decreasing [58], and thus marker numbers should be less of a barrier than previously. So, how many markers does a study need to reliably 486 487 detect backcrossed individuals? 488 489 To maximize detection of backcrossed individuals, researchers can increase their

490 power in three ways; through increased divergence, the use of diagnostic

491 markers, or with increased numbers of markers. Studies with high divergence

492 between hybridizing species have high power [33]. However, as many

493 conservation biologists choose their study system based on conservation

494 concerns and not to maximize power, this advice is not helpful. Diagnostic

495 markers have fixed allelic differences between parental species and are the most

496 powerful for backcross detection [25]. Ancestry informative markers, those with

497 strong allele frequency divergence between species, are also very powerful [63].

498 Loci with weak allele frequency divergence between species are least useful.

499 Diagnostic and ancestry informative markers can be determined based on

500 genotyping and contrasting known parental species individuals, although this is

not always feasible (e.g. [55]). Additionally, the diagnostic properties of markers

are a function of the populations and individuals that have been sampled; more

503 extensive sampling sometimes demonstrates that selected markers are not

504 diagnostic for all populations [66]. Generally speaking, the more markers used,

the higher the chance of detection of admixture in an individual [33, 44].

506

507 Assuming diagnostic markers, it is ideal to know the number of elapsed508 generations since the initial hybridization, as, for every further generation of

backcrossing, the proportion of introgressed genome halves [44]. The number of

510 generations since hybridization should be interpreted with an eye to policy.

511 After some number of generations of uni-directional backcrossing, policy will

512 dictate that we consider an individual to be parental species (again) [67]. It's

513 best to make this decision prior to marker selection, as it is impossible to apply

514 policy decisions regarding the acceptability of backcrossed individuals without

515 sufficient detection power.

516

517 If we are interested in all generations of backcrossing, then we can extend the 518 deterministic model developed by Boecklen and Howard ([44]; Equation 2) for 519 the genomics era. We made the same assumptions, specifically that backcrossing 520 is unidirectional, loci are independent and Mendelian, all markers are diagnostic, 521 all backcrossing is between the previous generation of backcrosses and parental 522 species, and all genotypes are equally fecund [44]. We asked what proportion of 523 backcrossed individuals are undetectable because they are homozygous for all 524 diagnostic markers. We modeled 10 generations of backcrossing, and each of 10, 525 100 and 1000 diagnostic markers (Figure I). When using 10 diagnostic markers, 526 52% of 4th generation backcrosses are homozygous for one parental species at 527 all loci, and thus undetectable as backcrosses. In contrast, 1000 diagnostic markers allow for powerful (85%) detection of 9th generation backcrosses. 528 529 530 531 532 Figure I: An extension of the deterministic model presented by Boecklen and 533 Howard [44]. The proportion of hybrid individuals that are homozygous at all

the (diagnostic) markers, and are hence indistinguishable from the parental

535 species that is being introgressed, increases with each generation of

536 backcrossing, but decreases with increased marker density. This demonstrates

that more markers than are typically used in studies of anthropogenic

538 hybridization are needed to detect advanced backcrosses.

539

540 **Box 2 – Reporting Error**

541 Credible (or confidence) intervals (CIs) are a powerful, intuitive way to assess 542 confidence in the estimates being presented [68, 69]. Measures of uncertainty 543 are not always presented in estimates of anthropogenic hybridization (although 544 see [53, 70-73] for exceptions), perhaps because the uncertainty is so high where 545 estimated. Credible intervals can be calculated using STRUCTURE [25] and 546 standard errors can be calculated using ADMIXTURE [29], so reporting of error 547 estimates is easily implemented in a routine workflow.

548

549 There are practical implications of the reporting of credible intervals,

550 particularly for individuals with very low or very high admixture values (Q). Cut-

off thresholds have been used to determine if individuals are members of the

parental populations or are admixed, but these thresholds are usually based on

the detection power of a study (see main text). Since these are hard cut-offs,

individuals with very similar levels of admixture can be assigned to very

different populations. For example, with a Q cut-off of 0.80, if individual 1 is

assessed as Q=0.79, it is determined to be admixed and, depending on the

557 management of the system, may be culled. In contrast, if individual 2 is estimated

to have Q=0.81, it would be considered a parental species individual and be

retained for breeding. There may be no substantive difference between these

560 individuals, although this is impossible to tell using only point estimates.

561

562 We recommend that credible intervals should also be included in visual

563 depictions of admixture. Typically, the key figure from a paper on anthropogenic

564 hybridization is the characteristic "STRUCTURE Bar Plot" [25], that uses stacked

565 colours to denote genetic contributions from different source populations. These

566 plots show the point estimates for each individual, and allow the author to

567 determine thresholds for inclusion in each group. While such figures are

568 compelling and easily interpreted, they do not convey the uncertainty around

569 individual point estimates.

570

571 Allendorf and colleagues [11] noted that it is very difficult to make policy

572 decisions when comparing different low point estimates of admixture. We

- 573 recommend that researchers should focus on the uncertainty around Q estimates
- 574 when making decisions about the genetic group each individual belongs to. It has
- 575 been pointed out that the use of credible intervals demonstrates the high levels
- 576 of uncertainty researchers are facing [70]. As they should! This problem will of
- 577 course be substantially alleviated by using more markers (see Box 1).
- 578

579 Box 3: Lessons from Natural Systems

580

581 Naturally occurring hybrid zones have long been used as 'natural laboratories' to 582 study the speciation process [74]. The field of speciation genomics works to 583 understand how genomic differences build up to cause eventual reproductive 584 isolation [75-78]. Recently, population geneticists have used genome wide 585 markers to ask questions regarding the genomic architecture of reproductive 586 isolation and speciation, and how the genomes of diverged populations change in 587 the face of on-going gene flow [43, 78, 79]. Further, many studies of natural 588 hybrid zones have focused on isolating signals from historical vs. contemporary 589 hybridization (main text 2.1.1, [78]). These questions that speciation biologists 590 ask using hybrid zones could equally be asked in anthropogenic hybrid zones, 591 particularly in studies that used whole genome sequence data. Indeed, studies of 592 anthropogenic hybrid zones may even have more power than those with 593 naturally occurring secondary contact as in some cases of introduced or escaped 594 heterospecifics, phenotypic divergence is more extreme, meaning that fewer 595 individuals would need to be sampled for, for example, admixture mapping [78]. 596

597 Use of genomic data allows speciation geneticists to examine heterogeneity in 598 divergence across the genome. Indeed, the questions we noted above are most 599 interesting when heterogeneity is found. Genome scans look for regions of high 600 divergence between species (F_{st} or d_{xy}) which may indicate regions that resist 601 introgression, also known as 'speciation islands' [80], or 'islands of 602 differentiation' [79]. While such signals are not without controversy [81], and in 603 some cases may represent phylogenetically derived regions of low 604 recombination, rather than reproductive isolation [82], they represent 605 interesting candidate regions for fixed differences between hybridizing species, 606 and thus could be used diagnostically by conservation biologists. For example, 607 golden-winged (Vermivora chrysoptera) and blue-winged warblers (V. 608 *cyanoptera*), which hybridize in eastern North America are phenotypically 609 distinct but undistinguishable when using low density, microsatellite marker 610 panels [83]. Only with the use of whole genome sequencing were six small 611 divergent regions of the genome discovered, four of which are associated with

- 612 either pigmentation or feather development genes and explain more than 90% of
- 613 the variation in plumage [39]. This demonstrates that a focus on the use of high
- 614 density markers to explore heterogeneity across the genome allows for higher
- 615 power to both distinguish between closely related, hybridizing species
- 616 genetically, and to associate genomic regions with diverged phenotypes, two
- 617 possible goals of conservation biologists working on anthropogenic hybrid
- cones. We echo the call of [1] that conservation biologists can take a cue from
- 619 speciation biologists that have, in many cases, developed methods that use
- 620 genomics to ask interesting questions of hybrid zones.
- 621

622 References

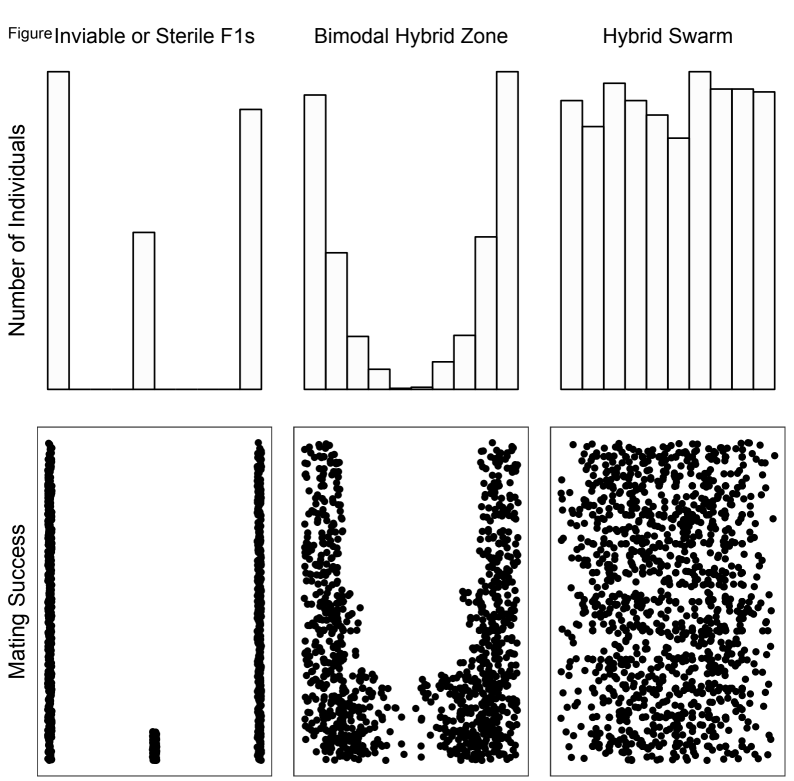
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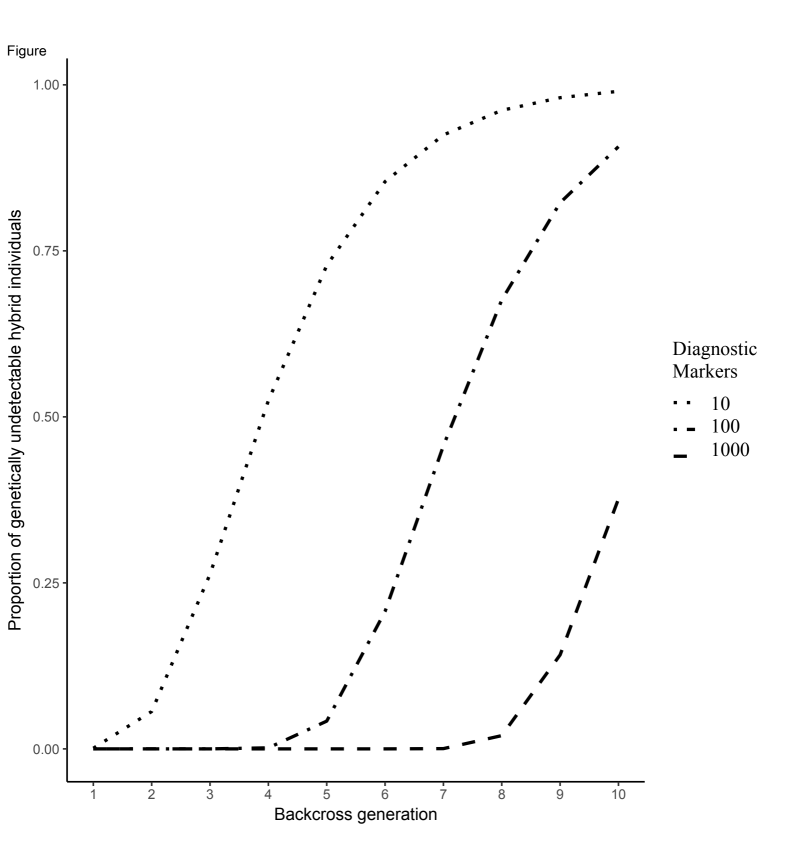
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Admixture (Q) scores from 0 - 1



Outstanding questions:

1) Do replicate anthropogenic hybrid zones show similar patterns of introgression?

There are big evolutionary questions that could be answered by the sorts of data that conservation biologists working on anthropogenic hybridization could answer. For example, there are multiple replicate hybrid zones occurring in the wolf/dog, wild cat/domestic cat, red deer/sika deer, westslope cut throat trout/rainbow trout systems. But in many cases, there is limited communication and collaboration between researchers, or different markers are used across studies [60]. Clearly this isn't a problem unique to this field, but it is the case that collaboration between researchers would be made easier with standardized genome wide data aligned to a common genome. Genomic data make cross study comparisons easier, and would allow for easier comparison between studies.

2) Once there has been a breakdown of reproductive isolation characterized as hybridization, how common is maintenance of within parental species assortative mating? Is the strength of assortative mating stronger when species are more diverged, or perhaps between closely related species that have recently evolved reproductive isolation?

3) What is the relative frequency of hybrid swarms vs bimodal hybrid zones? We expect that the prevalence of bimodal hybrid zones has been underestimated because of the difficulty of detecting highly introgressed backcrosses. Increased use of high-density markers will make these cases easier to detect and would enhance our understanding of the systems that are bimodal hybrid zones.

Highlights:

Anthropogenic hybridization is increasingly common and likely to result in a breakdown of reproductive isolation between 'good' species.

Backcrossed individuals that have only a small proportion of one parental genome are difficult to differentiate from parental individuals using the most common current technologies.

Bimodal hybrid zones are characterized by introgression and backcrossing. The majority of hybrid individuals in these systems have low levels of introgression. The problems posed by bimodal hybrid zones have been largely overlooked in the literature.

Genome wide sampling of genetic markers at high densities allow for increased precision in the estimate of admixture proportions, which makes it feasible to detect multi-generation backcrosses, and will thus make it easier to differentiate bimodal hybrid zones from hybrid swarms or systems without introgression.